

Package ‘misha’

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Type Package

Title Toolkit for Analysis of Genomic Data

Version 5.6.6

Description A toolkit for analysis of genomic data. The 'misha' package implements an efficient data structure for storing genomic data, and provides a set of functions for data extraction, manipulation and analysis. Some of the 2D genome algorithms were described in Yaffe and Tanay (2011) <[doi:10.1038/ng.947](https://doi.org/10.1038/ng.947)>.

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URL <https://tanaylab.github.io/misha/>,
<https://github.com/tanaylab/misha>

BugReports <https://github.com/tanaylab/misha/issues>

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misha-package *Toolkit for analysis of genomic data*

Description

'misha' package is intended to help users to efficiently analyze genomic data achieved from various experiments.

Details

For a complete list of help resources, use `library(help = "misha")`.

The following options are available for the package. Use 'options' function to alter the value of the options.

| NAME | DEFAULT | DESCRIPTION |
|---------------------|----------|---|
| gmax.data.size | AUTO | Auto-configured based on system RAM and processes. Formula: $\min((\text{RAM} * 0.7) / \text{gmax.processes}, 10\text{GB})$. Controls max in-memory result buffer size for streaming/sampling. Operations like gextract, gscreen use this as upper bound. |
| gbig.intervals.size | 1000000 | Threshold for converting interval sets to disk-based "big" format. When interval count exceeds this, intervals are stored per-chromosome instead of all in memory. Note: Independent of gmax.data.size (different purposes). |
| gmax.mem.usage | 10000000 | Maximal memory consumption of all child processes in KB before the limiting algorithm is invoked. |
| gmax.processes | AUTO | Auto-configured to 70% of CPU cores. Maximal number of processes for multitasking. |
| gmax.processes2core | 2 | Maximal number of processes per CPU core for multitasking |

| | | |
|-------------------------------|--------|--|
| gmin.scope4process | 10000 | Minimal scope range (for 2D: surface) assigned to a process in multitasking mode. |
| gbuf.size | 1000 | Size of track expression values buffer. |
| gmultitask.max.records.factor | 64 | Multiplier to inflate multitask max_records estimates to avoid under-allocation. |
| gtrack.chunk.size | 100000 | Chunk size in bytes of a 2D track. If '0' chunk size is unlimited. |
| gtrack.num.chunks | 0 | Maximal number of 2D track chunks simultaneously stored in memory. |
| gmultitasking | TRUE | Enable/disable automatic parallelization. Some functions may choose single-threaded mode for very small workloads. |

More information about the options can be found in 'User manual' of the package.

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See Also

Useful links:

- <https://tanaylab.github.io/misha/>
- <https://github.com/tanaylab/misha>
- Report bugs at <https://github.com/tanaylab/misha/issues>

gbins.quantiles

Calculates quantiles of a track expression for bins

Description

Calculates quantiles of a track expression for bins.

Usage

```
gbins.quantiles(
  ...,
  expr = NULL,
  percentiles = 0.5,
  intervals = get("ALLGENOME", envir = .misha),
  include.lowest = FALSE,
  iterator = NULL,
  band = NULL
)
```

Arguments

| | |
|----------------|---|
| ... | pairs of track expressions ('bin_expr') that determines the bins and breaks that define the bins. See gdist . |
| expr | track expression for which quantiles are calculated |
| percentiles | an array of percentiles of quantiles in [0, 1] range |
| intervals | genomic scope for which the function is applied. |
| include.lowest | if 'TRUE', the lowest value of the range determined by breaks is included |
| iterator | track expression iterator. If 'NULL' iterator is determined implicitly based on track expressions. |
| band | track expression band. If 'NULL' no band is used. |

Details

This function is a binned version of 'gquantiles'. For each iterator interval the value of 'bin_expr' is calculated and assigned to the corresponding bin determined by 'breaks'. The quantiles of 'expr' are calculated then separately for each bin.

The bins can be multi-dimensional depending on the number of 'bin_expr'-'breaks' pairs.

The range of bins is determined by 'breaks' argument. For example: 'breaks=c(x1, x2, x3, x4)' represents three different intervals (bins): (x1, x2], (x2, x3], (x3, x4].

If 'include.lowest' is 'TRUE' the the lowest value will be included in the first interval, i.e. in [x1, x2].

Value

Multi-dimensional array representing quantiles for each percentile and bin.

See Also

[gquantiles](#), [gintervals.quantiles](#), [gdist](#)

Examples

```
gdb.init_examples()
gbins.quantiles("dense_track", c(0, 0.2, 0.4, 2), "sparse_track",
  percentiles = c(0.2, 0.5),
  intervals = gintervals(1),
  iterator = "dense_track"
)
```

| | |
|----------------------------|---|
| <code>gbins.summary</code> | <i>Calculates summary statistics of a track expression for bins</i> |
|----------------------------|---|

Description

Calculates summary statistics of a track expression for bins.

Usage

```
gbins.summary(
  ...,
  expr = NULL,
  intervals = get("ALLGENOME", envir = .misha),
  include.lowest = FALSE,
  iterator = NULL,
  band = NULL
)
```

Arguments

| | |
|-----------------------------|---|
| <code>...</code> | pairs of track expressions ('bin_expr') that determines the bins and breaks that define the bins. See gdist . |
| <code>expr</code> | track expression for which summary statistics is calculated |
| <code>intervals</code> | genomic scope for which the function is applied |
| <code>include.lowest</code> | if 'TRUE', the lowest value of the range determined by breaks is included |
| <code>iterator</code> | track expression iterator. If 'NULL' iterator is determined implicitly based on track expressions. |
| <code>band</code> | track expression band. If 'NULL' no band is used. |

Details

This function is a binned version of 'gsummary'. For each iterator interval the value of 'bin_expr' is calculated and assigned to the corresponding bin determined by 'breaks'. The summary statistics of 'expr' are calculated then separately for each bin.

The bins can be multi-dimensional depending on the number of 'bin_expr'-'breaks' pairs.

The range of bins is determined by 'breaks' argument. For example: 'breaks=c(x1, x2, x3, x4)' represents three different intervals (bins): (x1, x2], (x2, x3], (x3, x4].

If 'include.lowest' is 'TRUE' the the lowest value will be included in the first interval, i.e. in [x1, x2].

Value

Multi-dimensional array representing summary statistics for each bin.

See Also

[gsummary](#), [gintervals.summary](#), [gdist](#)

Examples

```
gdb.init_examples()
gbins.summary("dense_track", c(0, 0.2, 0.4, 2), "sparse_track",
  intervals = gintervals(1), iterator = "dense_track"
)
```

gcis_decay

Calculates distribution of contact distances

Description

Calculates distribution of contact distances.

Usage

```
gcis_decay(
  expr = NULL,
  breaks = NULL,
  src = NULL,
  domain = NULL,
  intervals = NULL,
  include.lowest = FALSE,
  iterator = NULL,
  band = NULL
)
```

Arguments

| | |
|--------|-------------------------------|
| expr | track expression |
| breaks | breaks that determine the bin |
| src | source intervals |

| | |
|----------------|---|
| domain | domain intervals |
| intervals | genomic scope for which the function is applied |
| include.lowest | if 'TRUE', the lowest value of the range determined by breaks is included |
| iterator | 2D track expression iterator. If 'NULL' iterator is determined implicitly based on track expressions. |
| band | track expression band. If 'NULL' no band is used. |

Details

A 2D iterator interval '(chrom1, start1, end1, chrom2, start2, end2)' is said to represent a contact between two 1D intervals I1 and I2: '(chrom1, start1, end1)' and '(chrom2, start2, end2)'.

For contacts where 'chrom1' equals to 'chrom2' and I1 is within source intervals the function calculates the distribution of distances between I1 and I2. The distribution is calculated separately for intra-domain and inter-domain contacts.

An interval is within source intervals if the unification of all source intervals fully overlaps it. 'src' intervals are allowed to contain overlapping intervals.

Two intervals I1 and I2 are within the same domain (intra-domain contact) if among the domain intervals exists an interval that fully overlaps both I1 and I2. Otherwise the contact is considered to be inter-domain. 'domain' must contain only non-overlapping intervals.

The distance between I1 and I2 is the absolute distance between the centers of these intervals, i.e.: $|(start1 + end1 - start2 - end2) / 2|$.

The range of distances for which the distribution is calculated is defined by 'breaks' argument. For example: 'breaks=c(x1, x2, x3, x4)' represents three different intervals (bins): (x1, x2], (x2, x3], (x3, x4].

If 'include.lowest' is 'TRUE' the the lowest value will be included in the first interval, i.e. in [x1, x2]

Value

2-dimensional vector representing the distribution of contact distances for inter and intra domains.

See Also

[gdist](#), [gtrack.2d.import_contacts](#)

Examples

```
gdb.init_examples()

src <- rbind(
  gintervals(1, 10, 100),
  gintervals(1, 200, 300),
  gintervals(1, 400, 500),
  gintervals(1, 600, 700),
  gintervals(1, 7000, 9100),
  gintervals(1, 9000, 18000),
```

```

    gintervals(1, 30000, 31000),
    gintervals(2, 1130, 15000)
  )

  domain <- rbind(
    gintervals(1, 0, 483000),
    gintervals(2, 0, 300000)
  )

  gcis_decay("rects_track", 50000 * (1:10), src, domain)

```

gcluster.run

Runs R commands on a cluster

Description

Runs R commands on a cluster that supports SGE.

Usage

```

gcluster.run(
  ...,
  opt.flags = "",
  max.jobs = 400,
  debug = FALSE,
  R = "R",
  control_dir = NULL
)

```

Arguments

| | |
|-------------|---|
| ... | R commands |
| opt.flags | optional flags for qsub command |
| max.jobs | maximal number of simultaneously submitted jobs |
| debug | if 'TRUE', additional reports are printed |
| R | command that launches R |
| control_dir | directory where the control files are stored. Note that this directory should be accessible from all nodes. If 'NULL', a temporary directory would be created under the current misha database. |

Details

This function runs R commands on a cluster by distributing them among cluster nodes. It must run on a machine that supports Sun Grid Engine (SGE). The order in which the commands are executed can not be guaranteed, therefore the commands must be inter-independent.

Optional flags to 'qsub' command can be passed through 'opt.flags' parameter. Users are strongly recommended to use only '-l' flag as other flags might interfere with those that are already used (-terse, -S, -o, -e, -V). For additional information please refer to the manual of 'qsub'.

The maximal number of simultaneously submitted jobs is controlled by 'max.jobs'.

Set 'debug' argument to 'TRUE' to allow additional report prints.

'gcluster.run' launches R on the cluster nodes to execute the commands. 'R' argument specifies how R executable should be invoked.

Value

Return value ('retv') is a list, such that 'retv[[i]]' represents the result of the run of command number 'i'. Each result consists of 4 fields that can be accessed by 'retv[[i]]\$FIELDNAME':

| <i>FIELDNAME</i> | <i>DESCRIPTION</i> |
|------------------|---|
| exit.status | Exit status of the command. Possible values: 'success', 'failure' or 'interrupted'. |
| retv | Return value of the command. |
| stdout | Standard output of the command. |
| stderr | Standard error of the command. |

Examples

```
gdb.init_examples()
# Run only on systems with Sun Grid Engine (SGE)
if (FALSE) {
  v <- 17
  gcluster.run(
    gsummary("dense_track + v"),
    {
      inters <- gscreen("dense_track > 0.1", gintervals(1, 2))
      gsummary("sparse_track", inters)
    },
    gsummary("rects_track")
  )
}
```

gcompute_strands_autocorr

Computes auto-correlation between the strands for a file of mapped sequences

Description

Calculates auto-correlation between plus and minus strands for the given chromosome in a file of mapped sequences.

Usage

```
gcompute_strands_autocorr(
  file = NULL,
  chrom = NULL,
  binsize = NULL,
  maxread = 400,
  cols.order = c(9, 11, 13, 14),
  min.coord = 0,
  max.coord = 3e+08
)
```

Arguments

| | |
|------------|---|
| file | the name of the file containing mapped sequences |
| chrom | chromosome for which the auto-correlation is computed |
| binsize | calculate the auto-correlation for bins in the range of [-maxread, maxread] |
| maxread | maximal length of the sequence used for statistics |
| cols.order | order of sequence, chromosome, coordinate and strand columns in file |
| min.coord | minimal coordinate used for statistics |
| max.coord | maximal coordinate used for statistics |

Details

This function calculates auto-correlation between plus and minus strands for the given chromosome in a file of mapped sequences. Each line in the file describes one read. Each column is separated by a TAB character.

The following columns must be presented in the file: sequence, chromosome, coordinate and strand. The position of these columns are controlled by 'cols.order' argument accordingly. The default value of 'cols.order' is a vector (9,11,13,14) meaning that sequence is expected to be found at column number 9, chromosome - at column 11, coordinate - at column 13 and strand - at column 14. The first column should be referenced by 1 and not by 0.

Coordinates that are not in [min.coord, max.coord] range are ignored.

gcompute_strands_autocorr outputs the total statistics and the auto-correlation given by bins. The size of the bin is indicated by 'binsize' parameter. Statistics is calculated for bins in the range of [-maxread, maxread].

Value

Statistics for each strand and auto-correlation by given bins.

Examples

```
gdb.init_examples()
gcompute_strands_autocorr(paste(.misha$GROOT, "reads", sep = "/"),
  "chr1", 50,
```

```

    maxread = 300
)

```

gcor

Calculates correlation between track expressions

Description

Calculates correlation between track expressions over iterator bins inside the supplied genomic scope. Expressions are processed in pairs: (expr1, expr2), (expr3, expr4), etc. Only bins where both expressions are not NaN are used.

Usage

```

gcor(
  expr1 = NULL,
  expr2 = NULL,
  ...,
  intervals = NULL,
  iterator = NULL,
  band = NULL,
  method = c("pearson", "spearman", "spearman.exact"),
  details = FALSE,
  names = NULL
)

```

Arguments

| | |
|-----------|--|
| expr1 | first track expression |
| expr2 | second track expression |
| ... | additional track expressions, supplied as pairs (expr3, expr4, ...) |
| intervals | genomic scope for which the function is applied |
| iterator | track expression iterator. If 'NULL' iterator is determined implicitly based on track expression. |
| band | track expression band. If 'NULL' no band is used. |
| method | correlation method to use. One of 'pearson' (default), 'spearman' (approximate, memory-efficient), or 'spearman.exact' (exact, requires O(n) memory where n is number of non-NaN pairs). |
| details | if 'TRUE' returns summary statistics for each pair, otherwise returns correlations only. For Pearson, includes n, n.na, mean1, mean2, sd1, sd2, cov, cor. For Spearman methods, includes n, n.na, cor. |
| names | optional names for the pairs. If supplied, length must match the number of pairs. |

Value

If 'details' is 'FALSE', a numeric vector of correlations. If 'details' is 'TRUE', a data frame with summary statistics for each pair.

See Also

[gextract](#), [gscreen](#), [gsummary](#)

Examples

```
gdb.init_examples()
gcor("dense_track", "sparse_track", intervals = gintervals(1, 0, 10000), iterator = 1000)

# Spearman correlation (approximate, memory-efficient)
gcor("dense_track", "sparse_track",
     intervals = gintervals(1, 0, 10000),
     iterator = 1000, method = "spearman"
)

# Exact Spearman correlation
gcor("dense_track", "sparse_track",
     intervals = gintervals(1, 0, 10000),
     iterator = 1000, method = "spearman.exact"
)
```

`gdataset.example_path` *Create an example dataset on the fly*

Description

Creates a small dataset in a temporary directory using the built-in example database. This function has side effects: it calls [gdb.init_examples](#) which resets the working database, and it creates then deletes temporary tracks ('example_dataset_track') and intervals ('example_dataset_intervals') in that database.

Usage

```
gdataset.example_path()
```

Details

This function performs the following steps:

1. Calls `gdb.init_examples()` to set the working database
2. Removes any existing 'example_dataset_track' and 'example_dataset_intervals'

3. Creates temporary track and intervals in the example database
4. Saves them to a new dataset in a temporary directory
5. Removes the temporary track and intervals from the example database

This is primarily intended for use in examples and tests. Users should be aware that calling this function will change their current working database.

Value

Path to the created dataset directory (in a temporary location)

See Also

[gdataset.save](#), [gdataset.load](#), [gdb.init_examples](#)

Examples

```
dataset_path <- gdataset.example_path()
gdataset.load(dataset_path)
gdataset.unload(dataset_path)
```

gdataset.info

Get dataset information

Description

Returns metadata and contents of a dataset.

Usage

```
gdataset.info(path)
```

Arguments

path Path to any dataset (loaded or not)

Value

List with dataset information

See Also

[gdataset.ls](#), [gdataset.load](#)

Examples

```
dataset_path <- gdataset.example_path()
gdataset.info(dataset_path)
```

| | |
|---------------|--|
| gdataset.load | <i>Load a dataset into the namespace</i> |
|---------------|--|

Description

Loads tracks and intervals from a dataset directory, making them available for analysis alongside the working database.

Usage

```
gdataset.load(path, force = FALSE, verbose = FALSE)
```

Arguments

| | |
|---------|--|
| path | Path to a dataset or misha database directory |
| force | If TRUE, ignore name collisions (working db wins; for dataset-to-dataset, later-loaded wins) |
| verbose | If TRUE, print loaded track/interval names and summary counts |

Value

Invisibly returns a list with:

| | |
|--------------------|--|
| tracks | Number of visible tracks loaded |
| intervals | Number of visible intervals loaded |
| shadowed_tracks | Number of tracks shadowed by collisions |
| shadowed_intervals | Number of intervals shadowed by collisions |

See Also

[gdataset.unload](#), [gdataset.save](#), [gdataset.ls](#)

Examples

```
dataset_path <- gdataset.example_path()
gdataset.load(dataset_path)
gdataset.unload(dataset_path)
```

`gdataset.ls` *List working database and loaded datasets*

Description

Returns a list of the working database and all loaded datasets.

Usage

```
gdataset.ls(dataframe = FALSE)
```

Arguments

`dataframe` If FALSE, return character vector; if TRUE, return data frame

Value

Character vector of paths or data frame with detailed information

See Also

[gdataset.load](#), [gdataset.info](#)

Examples

```
dataset_path <- gdataset.example_path()
gdataset.load(dataset_path)
gdataset.ls()
gdataset.unload(dataset_path)
```

`gdataset.save` *Save a dataset*

Description

Creates a new dataset directory containing selected tracks and/or intervals from the working database.

Usage

```
gdataset.save(
  path,
  description,
  tracks = NULL,
  intervals = NULL,
  symlinks = FALSE,
  copy_seq = FALSE
)
```

Arguments

| | |
|-------------|---|
| path | Destination directory (must not exist) |
| description | Required description for metadata |
| tracks | Character vector of track names to include |
| intervals | Character vector of interval set names to include |
| symlinks | If TRUE, create symlinks to tracks/intervals instead of copying |
| copy_seq | If TRUE, copy seq/ directory instead of symlinking |

Value

Invisible path

See Also

[gdataset.load](#), [gdataset.info](#)

Examples

```
gdb.init_examples()
example_intervs <- gintervals(1, 0, 10000)
gintervals.save("example_dataset_intervals", example_intervs)
gtrack.create(
  "example_dataset_track",
  "Example dataset track",
  "dense_track",
  iterator = "example_dataset_intervals"
)
dataset_path <- tempfile("misha_dataset_")
gdataset.save(
  path = dataset_path,
  description = "Example dataset",
  tracks = "example_dataset_track",
  intervals = "example_dataset_intervals"
)
gtrack.rm("example_dataset_track", force = TRUE)
gintervals.rm("example_dataset_intervals", force = TRUE)
```

gdataset.unload

Unload a dataset from the namespace

Description

Removes all tracks and intervals from a previously loaded dataset. If a track was shadowing another, the shadowed track becomes visible again.

Usage

```
gdataset.unload(path, validate = FALSE)
```

Arguments

| | |
|----------|--|
| path | Path to a previously loaded dataset |
| validate | If TRUE, error if path is not currently loaded; otherwise silently no-op |

Value

Invisible NULL

See Also

[gdataset.load](#), [gdataset.ls](#)

Examples

```
dataset_path <- gdataset.example_path()
gdataset.load(dataset_path)
gdataset.unload(dataset_path, validate = TRUE)
```

`gdb.convert_to_indexed`

Change Database to Indexed Genome Format

Description

Converts a per-chromosome database to indexed genome format with a single consolidated genome.seq file and genome.idx index. Optionally also converts tracks and interval sets to indexed format.

Usage

```
gdb.convert_to_indexed(
  groot = NULL,
  remove_old_files = FALSE,
  force = FALSE,
  validate = TRUE,
  convert_tracks = FALSE,
  convert_intervals = FALSE,
  verbose = FALSE,
  chunk_size = 104857600
)
```

Arguments

| | |
|-------------------|---|
| groot | Root directory of the database to change to indexed format. If NULL, uses the currently active database. |
| remove_old_files | Logical. If TRUE, removes old per-chromosome files after successful conversion. Default: FALSE. |
| force | Logical. If TRUE, forces the conversion without confirmation. Default: FALSE. |
| validate | Logical. If TRUE, validates the conversion by comparing sequences. Default: TRUE. |
| convert_tracks | Logical. If TRUE, also converts all eligible tracks to indexed format. Default: FALSE. |
| convert_intervals | Logical. If TRUE, also converts all eligible interval sets to indexed format. Default: FALSE. |
| verbose | Logical. If TRUE, prints verbose messages. Default: FALSE. |
| chunk_size | Integer. The size of the chunk to read from the sequence files. Default: 104857600 (100MB). Reduce if you are running into memory issues. |

Details

This function converts a per-chromosome database (with separate .seq files per contig) to indexed format (single genome.seq + genome.idx). The indexed format provides better performance and scalability, especially for genomes with many contigs.

Important: Preserving Chromosome Order

For exact conversion that produces bit-for-bit identical results before and after conversion, you should load the source database first using `gsetroot()` or `gdb.init()`:

- If database is loaded: Uses chromosome order from ALLGENOME (exact preservation)
- If database is not loaded: Uses order from chrom_sizes.txt (may differ from ALLGENOME)

This ensures that the converted database has the exact same chromosome ordering, which affects iteration order, interval IDs, and other operations that depend on chromosome order.

The conversion process:

1. Checks if database is already in indexed format
2. Gets chromosome order from ALLGENOME (if loaded) or chrom_sizes.txt
3. Consolidates all per-chromosome .seq files into genome.seq
4. Creates genome.idx with CRC64 checksum
5. Optionally validates the conversion
6. Optionally removes old .seq files
7. If convert_tracks=TRUE, converts all eligible tracks (1D: dense, sparse, array; 2D: rectangles, points)
8. If convert_intervals=TRUE, converts all eligible interval sets (1D and 2D)

Tracks and intervals that cannot be converted (and are skipped):

- Tracks: virtual tracks, single-file tracks, already converted tracks
- Intervals: Single-file interval sets, already converted interval sets

Value

Invisible NULL

See Also

[gdb.create](#), [gdb.init](#), [gtrack.convert_to_indexed](#), [gintervals.convert_to_indexed](#), [gintervals.2d.convert_to_indexed](#)

Examples

```
## Not run:
# Recommended: Load database first for exact conversion
gsetroot("/path/to/database")
gdb.convert_to_indexed(
  convert_tracks = TRUE,
  convert_intervals = TRUE,
  remove_old_files = TRUE,
  verbose = TRUE
)

# Convert current database to indexed format (genome only)
gdb.convert_to_indexed()

# Convert specific database without loading it first
# Note: chromosome order may differ from ALLGENOME
gdb.convert_to_indexed(groot = "/path/to/database")

# Convert genome and all tracks to indexed format
gdb.convert_to_indexed(convert_tracks = TRUE)

# Full conversion with validation and cleanup
gsetroot("/path/to/database") # Load first for exact order preservation
gdb.convert_to_indexed(
  convert_tracks = TRUE,
  convert_intervals = TRUE,
  remove_old_files = TRUE,
  validate = TRUE,
  verbose = TRUE
)

## End(Not run)
```

`gdb.create`

Creates a new Genomic Database

Description

Creates a new Genomic Database.

Usage

```
gdb.create(
  groot = NULL,
  fasta = NULL,
  genes.file = NULL,
  annots.file = NULL,
  annots.names = NULL,
  format = NULL,
  verbose = FALSE
)
```

Arguments

| | |
|---------------------------|--|
| <code>groot</code> | path to newly created database |
| <code>fasta</code> | an array of names or URLs of FASTA files. Can contain wildcards for multiple files |
| <code>genes.file</code> | name or URL of file that contains genes. If 'NULL' no genes are imported |
| <code>annots.file</code> | name of URL file that contains annotations. If 'NULL' no annotations are imported |
| <code>annots.names</code> | annotations names |
| <code>format</code> | database format: "indexed" (default, single genome.seq + genome.idx) or "per-chromosome" (separate .seq file per contig). If NULL, uses the value from <code>getOption("gmulticontig.indexed_format", TRUE)</code> |
| <code>verbose</code> | if TRUE, prints verbose messages |

Details

This function creates a new Genomic Database at the location specified by 'groot'. FASTA files are converted to 'Seq' format and appropriate 'chrom_sizes.txt' file is generated (see "User Manual" for more details).

Two database formats are supported:

- **indexed**: Single genome.seq + genome.idx (default). Recommended for genomes with many contigs. Provides better performance and scalability.
- **per-chromosome**: Separate .seq file per contig.

If 'genes.file' is not 'NULL' four sets of intervals are created in the database: tss, exons, utr3 and utr5. See [gintervals.import_genes](#) for more details about importing genes intervals.

'fasta', 'genes.file' and 'annots.file' can be either a file path or URL in a form of 'ftp://[address]/[file]'. 'fasta' can also contain wildcards to indicate multiple files. Files that these arguments point to can be zipped or unzipped.

See the 'Genomes' vignette for details on how to create a database from common genome sources.

Value

None.

See Also

[gdb.init](#), [gdb.reload](#), [gintervals.import_genes](#)

Examples

```
# ftp <- "ftp://hgdownload.soe.ucsc.edu/goldenPath/mm10"
# mm10_dir <- file.path(tempdir(), "mm10")
# # only a single chromosome is loaded in this example
# # see "Genomes" vignette how to download all of them and how
# # to download other genomes
# gdb.create(
#   mm10_dir,
#   paste(ftp, "chromosomes", paste0(
#     "chr", c("X"),
#     ".fa.gz"
#   ), sep = "/"),
#   paste(ftp, "database/knownGene.txt.gz", sep = "/"),
#   paste(ftp, "database/kgXref.txt.gz", sep = "/"),
#   c(
#     "kgID", "mRNA", "spID", "spDisplayID", "geneSymbol",
#     "refseq", "protAcc", "description", "rfamAcc",
#     "tRnaName"
#   )
# )
# gdb.init(mm10_dir)
# gintervals.ls()
# gintervals.all()
```

gdb.create_genome

Create and Load a Genome Database

Description

This function downloads, extracts, and loads a misha genome database for the specified genome.

Usage

```
gdb.create_genome(genome, path = getwd(), tmpdir = tempdir())
```

Arguments

| | |
|--------|---|
| genome | A character string specifying the genome to download. Supported genomes are "mm9", "mm10", "mm39", "hg19", and "hg38". |
| path | A character string specifying the directory where the genome will be extracted. Defaults to genome name (e.g. "mm10") in the current working directory. |
| tmpdir | A character string specifying the directory for storing temporary files. This is used for storing the downloaded genome file. |

Details

The function checks if the specified genome is available. If tmpdir, it constructs the download URL, downloads the genome file, extracts it to the specified directory, and loads the genome database using gsetroot. The function also calls gdb.reload to reload the genome database.

Value

None.

Examples

```
## Not run:
mm10_dir <- tempdir()
gdb.create_genome("mm10", path = mm10_dir)
list.files(file.path(mm10_dir, "mm10"))
gsetroot(file.path(mm10_dir, "mm10"))
gintervals.ls()

## End(Not run)
```

| | |
|-------------------|--|
| gdb.create_linked | <i>Create a linked database with symlinks to a parent database</i> |
|-------------------|--|

Description

Creates a new database directory structure with symbolic links to the parent database's seq/ directory and chrom_sizes.txt file.

Usage

```
gdb.create_linked(path, parent)
```

Arguments

| | |
|--------|--|
| path | Path for the new linked database |
| parent | Path to the parent database (with seq and chrom_sizes.txt) |

Details

This is useful for creating a writable database that shares sequence data with a read-only main database. The new database can be set as the working database via gsetroot(), and then the parent database can be loaded as a dataset via gdataset.load().

Value

Invisible TRUE on success

See Also

[gsetroot](#), [gdb.create](#), [gdataset.load](#), [gdataset.ls](#)

Examples

```
## Not run:
# Create linked database sharing sequence data with main database
gdb.create_linked("~/my_tracks", parent = "/shared/genomics/hg38")

# Set linked database as working database and load parent as dataset
gsetroot("~/my_tracks")
gdataset.load("/shared/genomics/hg38")

## End(Not run)
```

`gdb.export_fasta` *Export a database genome as FASTA*

Description

Writes all contigs from a misha database to a multi-FASTA file.

Usage

```
gdb.export_fasta(
  file = NULL,
  groot = NULL,
  line_width = 80L,
  chunk_size = 1000000L,
  overwrite = FALSE,
  verbose = FALSE
)
```

Arguments

| | |
|-------------------------|---|
| <code>file</code> | Output FASTA file path |
| <code>groot</code> | Optional database root path. If NULL, uses current database. |
| <code>line_width</code> | Number of bases per FASTA line. Default: 80. |
| <code>chunk_size</code> | Number of bases to extract per chunk while writing. Default: 1000000. |
| <code>overwrite</code> | Logical. If TRUE, overwrite existing output file. Default: FALSE. |
| <code>verbose</code> | Logical. If TRUE, prints progress messages. Default: FALSE. |

Details

By default, the currently active database is used. You can also provide `groot` to export another database without changing the caller's active database.

Value

Invisibly returns file.

See Also

[gdb.init](#), [gseq.extract](#)

Examples

```
## Not run:
gdb.init_examples()
out <- tempfile(fileext = ".fa")
gdb.export_fasta(out)
head(readLines(out))

## End(Not run)
```

`gdb.get_readonly_attrs`

Returns a list of read-only track attributes

Description

Returns a list of read-only track attributes.

Usage

```
gdb.get_readonly_attrs()
```

Details

This function returns a list of read-only track attributes. These attributes are not allowed to be modified or deleted.

If no attributes are marked as read-only a 'NULL' is returned.

Value

A list of read-only track attributes.

See Also

[gdb.set_readonly_attrs](#), [gtrack.attr.get](#), [gtrack.attr.set](#)

`gdb.info`*Get Database Information*

Description

Returns information about a misha genome database including format, number of chromosomes, total genome size, and whether it uses the indexed format.

Usage

```
gdb.info(groot = NULL)
```

Arguments

`groot` Root directory of the database. If NULL, uses the currently active database.

Value

A list with database information:

- `path` - Full path to the database
- `is_db` - TRUE if this is a valid misha database
- `format` - "indexed" or "per-chromosome"
- `num_chromosomes` - Number of chromosomes/contigs
- `genome_size` - Total length of genome in bases
- `chromosomes` - Data frame with chromosome names and sizes

Examples

```
## Not run:  
# Get info about currently active database  
info <- gdb.info()  
cat("Database format:", info$format, "\n")  
cat("Genome size:", info$genome_size / 1e6, "Mb\n")  
  
# Get info about specific database  
info <- gdb.info("/path/to/database")  
  
## End(Not run)
```

| | |
|----------|---|
| gdb.init | <i>Initializes connection with Genomic Database</i> |
|----------|---|

Description

Initializes connection with Genomic Database: loads the list of tracks, intervals, etc.

Usage

```
gdb.init(groot = NULL, dir = NULL, rescan = FALSE)
```

```
gdb.init_examples()
```

```
gsetroot(groot = NULL, dir = NULL, rescan = FALSE)
```

Arguments

| | |
|--------|---|
| groot | the root directory of the Genomic Database |
| dir | the current working directory inside the Genomic Database |
| rescan | indicates whether the file structure should be rescanned |

Details

'gdb.init' initializes the connection with the Genomic Database. It is typically called first prior to any other function. When the package is attached it internally calls to 'gdb.init.examples' which opens the connection with the database located at 'PKGDIR/trackdb/test' directory, where 'PKGDIR' is the directory where the package is installed.

The current working directory inside the Genomic Database is set to 'dir'. If 'dir' is 'NULL', the current working directory is set to 'GROOT/tracks'.

If 'rescan' is 'TRUE', the list of tracks and intervals is achieved by rescanning directory structure under the current current working directory. Otherwise 'gdb.init' attempts to use the cached list that resides in 'groot/db.cache' file.

Upon completion the connection is established with the database. If auto-completion mode is switched on (see 'gset_input_method') the list of tracks and intervals sets is loaded and added as variables to the global environment allowing auto-completion of object names with <TAB> key. Also a few variables are defined at an environment called .misha, and can be accessed using .misha\$variable, e.g. .misha\$ALLGENOME. These variables should not be modified by user.

| | |
|---------------------|---|
| GROOT | Root directory of Genomic Database |
| GWD | Current working directory inside Genomic Database |
| GTRACKS | List of all available tracks |
| GINTERVS | List of all available intervals |
| GVTRACKS | List of all available virtual tracks |
| ALLGENOME | List of all chromosomes and their sizes |
| GITERATOR.INTERVALS | A set of iterator intervals for which the track expression is evaluated |

When option 'gmulticontig.indexed_format' is set to TRUE, the function loads a database with "indexed" track format.

Value

None.

See Also

[gdb.reload](#), [gdb.create](#), [gdir.cd](#), [gtrack.ls](#), [gintervals.ls](#), [gvtrack.ls](#)

`gdb.mark_cache_dirty` *Mark cached track list as dirty*

Description

When tracks or interval sets are modified outside of misha (e.g. files copied manually), the cached inventory may become out of date. Calling this helper marks the cache as dirty so the next `gsetroot()` forces a rescan.

Usage

```
gdb.mark_cache_dirty()
```

Value

Invisible TRUE if the dirty flag was written, FALSE otherwise.

See Also

[gdb.reload](#), [gsetroot](#)

| | |
|------------|---------------------------------------|
| gdb.reload | <i>Reloads database from the disk</i> |
|------------|---------------------------------------|

Description

Reloads database from disk: list of tracks, intervals, etc.

Usage

```
gdb.reload(rescan = TRUE)
```

Arguments

rescan indicates whether the file structure should be rescanned

Details

Reloads Genomic Database from disk: list of tracks, intervals, etc. Use this function if you manually add tracks or if for any reason the database becomes corrupted. If 'rescan' is 'TRUE', the list of tracks and intervals is achieved by rescanning directory structure under the current current working directory. Otherwise 'gdb.reload' attempts to use the cached list that resides in 'GROOT/.db.cache' file.

Value

No return value, called for side effects.

See Also

[gdb.init](#), [gdb.create](#), [gdir.cd](#),

| | |
|------------------------|--|
| gdb.set_readonly_attrs | <i>Sets read-only track attributes</i> |
|------------------------|--|

Description

Sets read-only track attributes.

Usage

```
gdb.set_readonly_attrs(attrs)
```

Arguments

attrs a vector of read-only attributes names or 'NULL'

Details

This function sets the list of read-only track attributes. The specified attributes may or may not already exist in the tracks.

If 'attrs' is 'NULL' the list of read-only attributes is emptied.

Value

None.

See Also

[gdb.get_readonly_attrs](#), [gtrack.attr.get](#), [gtrack.attr.set](#)

gdir.cd

Changes current working directory in Genomic Database

Description

Changes current working directory in Genomic Database.

Usage

```
gdir.cd(dir = NULL)
```

Arguments

dir directory path

Details

This function changes the current working directory in Genomic Database (not to be confused with shell's current working directory). The list of database objects - tracks, intervals, track variables - is rescanned recursively under 'dir'. Object names are updated with the respect to the new current working directory. Example: a track named 'subdir.dense' will be referred as 'dense' once current working directory is set to 'subdir'. All virtual tracks are removed.

Value

None.

See Also

[gdb.init](#), [gdir.cwd](#), [gdir.create](#), [gdir.rm](#)

Examples

```
gdb.init_examples()
gdir.cd("subdir")
gtrack.ls()
gdir.cd("../")
gtrack.ls()
```

| | |
|-------------|--|
| gdir.create | <i>Creates a new directory in Genomic Database</i> |
|-------------|--|

Description

Creates a new directory in Genomic Database.

Usage

```
gdir.create(dir = NULL, showWarnings = TRUE, mode = "0777")
```

Arguments

| | |
|--------------|------------------|
| dir | directory path |
| showWarnings | see 'dir.create' |
| mode | see 'dir.create' |

Details

This function creates a new directory in Genomic Database. Creates only the last element in the specified path.

Value

None.

Note

A new directory cannot be created within an existing track directory.

See Also

[dir.create](#), [gdb.init](#), [gdir.cwd](#), [gdir.rm](#)

| | |
|----------|---|
| gdir.cwd | Returns the current working directory in Genomic Database |
|----------|---|

Description

Returns the absolute path of the current working directory in Genomic Database.

Usage

```
gdir.cwd()
```

Details

This function returns the absolute path of the current working directory in Genomic Database (not to be confused with shell's current working directory).

Value

A character string of the path.

See Also

[gdb.init](#), [gdir.cd](#), [gdir.create](#), [gdir.rm](#)

| | |
|---------|---|
| gdir.rm | Deletes a directory from Genomic Database |
|---------|---|

Description

Deletes a directory from Genomic Database.

Usage

```
gdir.rm(dir = NULL, recursive = FALSE, force = FALSE)
```

Arguments

| | |
|-----------|---|
| dir | directory path |
| recursive | if 'TRUE', the directory is deleted recursively |
| force | if 'TRUE', suppresses user confirmation of tracks/intervals removal |

Details

This function deletes a directory from Genomic Database. If 'recursive' is 'TRUE', the directory is deleted with all the files/directories it contains. If the directory contains tracks or intervals, the user is prompted to confirm the deletion. Set 'force' to 'TRUE' to suppress the prompt.

Value

None.

See Also[gdb.init](#), [gdir.create](#), [gdir.cd](#), [gdir.cwd](#)

gdist*Calculates distribution of track expressions*

Description

Calculates distribution of track expressions' values over the given set of bins.

Usage

```

gdist(
  ...,
  intervals = NULL,
  include.lowest = FALSE,
  iterator = NULL,
  band = NULL,
  dataframe = FALSE,
  names = NULL
)

```

Arguments

| | |
|----------------|--|
| ... | pairs of 'expr', 'breaks' where 'expr' is a track expression and the breaks determine the bin |
| intervals | genomic scope for which the function is applied |
| include.lowest | if 'TRUE', the lowest value of the range determined by breaks is included |
| iterator | track expression iterator. If 'NULL' iterator is determined implicitly based on track expressions. |
| band | track expression band. If 'NULL' no band is used. |
| dataframe | return a data frame instead of an N-dimensional vector. |
| names | names for track expressions in the returned dataframe (only relevant when dataframe == TRUE) |

Details

This function calculates the distribution of values of the numeric track expressions over the given set of bins.

The range of bins is determined by 'breaks' argument. For example: 'breaks=c(x1, x2, x3, x4)' represents three different intervals (bins): (x1, x2], (x2, x3], (x3, x4].

If 'include.lowest' is 'TRUE' the the lowest value will be included in the first interval, i.e. in [x1, x2]

'gdist' can work with any number of dimensions. If more than one 'expr'-'breaks' pair is passed, the result is a multidimensional vector, and an individual value can be accessed by [i1,i2,...,iN] notation, where 'i1' is the first track and 'iN' is the last track expression.

Value

N-dimensional vector where N is the number of 'expr'-'breaks' pairs. If dataframe == TRUE - a data frame with a column for each track expression and an additional column 'n' with counts.

See Also

[gextract](#)

Examples

```
gdb.init_examples()

## calculate the distribution of dense_track for bins:
## (0, 0.2], (0.2, 0.5] and (0.5, 1]
gdist("dense_track", c(0, 0.2, 0.5, 1))

## calculate two-dimensional distribution:
## dense_track vs. sparse_track
gdist("dense_track", seq(0, 1, by = 0.1), "sparse_track",
      seq(0, 2, by = 0.2),
      iterator = 100
    )
```

gextract

Returns evaluated track expression

Description

Returns the result of track expressions evaluation for each of the iterator intervals.

Usage

```

gextract(
  ...,
  intervals = NULL,
  colnames = NULL,
  iterator = NULL,
  band = NULL,
  file = NULL,
  intervals.set.out = NULL
)

```

Arguments

| | |
|-------------------|--|
| ... | track expression |
| intervals | genomic scope for which the function is applied |
| colnames | sets the columns names in the returned value. If 'NULL' names are set to track expression. |
| iterator | track expression iterator. If 'NULL' iterator is determined implicitly based on track expressions. |
| band | track expression band. If 'NULL' no band is used. |
| file | file name where the function result is optionally outputted in tab-delimited format |
| intervals.set.out | intervals set name where the function result is optionally outputted |

Details

This function returns the result of track expressions evaluation for each of the iterator intervals. The returned value is a set of intervals with an additional column for each of the track expressions. This value can be used as an input for any other function that accepts intervals. If the intervals inside 'intervals' argument overlap gextract returns the overlapped coordinate more than once.

The order inside the result might not be the same as the order of intervals. An additional column 'intervalID' is added to the return value. Use this column to refer to the index of the original interval from the supplied 'intervals'.

If 'file' parameter is not 'NULL' the result is outputted to a tab-delimited text file (without 'intervalID' column) rather than returned to the user. This can be especially useful when the result is too big to fit into the physical memory. The resulted file can be used as an input for 'gtrack.import' or 'gtrack.array.import' functions.

If 'intervals.set.out' is not 'NULL' the result is saved as an intervals set. Similarly to 'file' parameter 'intervals.set.out' can be useful to overcome the limits of the physical memory.

'colnames' parameter controls the names of the columns that contain the evaluated expressions. By default the column names match the track expressions.

Value

If 'file' and 'intervals.set.out' are 'NULL' a set of intervals with an additional column for each of the track expressions and 'columnID' column.

See Also

[gtrack.array.extract](#), [gsample](#), [gtrack.import](#), [gtrack.array.import](#), [glookup](#), [gpartition](#), [gdist](#)

Examples

```
gdb.init_examples()

## get values of 'dense_track' for [0, 400), chrom 1
gextract("dense_track", gintervals(1, 0, 400))

## get values of 'rects_track' (a 2D track) for a 2D interval
gextract(
  "rects_track",
  gintervals.2d("chr1", 0, 4000, "chr2", 2000, 5000)
)
```

| | |
|------------|--------------------------------------|
| gintervals | <i>Creates a set of 1D intervals</i> |
|------------|--------------------------------------|

Description

Creates a set of 1D intervals.

Usage

```
gintervals(chroms = NULL, starts = 0, ends = -1, strands = NULL)
```

Arguments

| | |
|---------|---|
| chroms | chromosomes - an array of strings with or without "chr" prefixes or an array of integers (like: '1' for "chr1") |
| starts | an array of start coordinates |
| ends | an array of end coordinates. If '-1' chromosome size is assumed. |
| strands | 'NULL' or an array consisting of '-1', '0' or '1' values |

Details

This function returns a set of one-dimensional intervals. The returned value can be used in all functions that accept 'intervals' argument.

One-dimensional intervals is a data frame whose first three columns are 'chrom', 'start' and 'end'. Each row of the data frame represents a genomic interval of the specified chromosome in the range of [start, end). Additional columns can be presented in 1D intervals object yet these columns must be added after the three obligatory ones.

If 'strands' argument is not 'NULL' an additional column "strand" is added to the intervals. The possible values of a strand can be '1' (plus strand), '-1' (minus strand) or '0' (unknown).

Value

A data frame representing the intervals.

See Also

[gintervals.2d](#), [gintervals.force_range](#)

Examples

```
gdb.init_examples()

## the following 3 calls produce identical results
gintervals(1)
gintervals("1")
gintervals("chrX")

gintervals(1, 1000)
gintervals(c("chr2", "chrX"), 10, c(3000, 5000))
```

| | |
|---------------|--------------------------------------|
| gintervals.2d | <i>Creates a set of 2D intervals</i> |
|---------------|--------------------------------------|

Description

Creates a set of 2D intervals.

Usage

```
gintervals.2d(
  chroms1 = NULL,
  starts1 = 0,
  ends1 = -1,
  chroms2 = NULL,
  starts2 = 0,
  ends2 = -1
)
```

Arguments

| | |
|---------|---|
| chroms1 | chromosomes1 - an array of strings with or without "chr" prefixes or an array of integers (like: '1' for "chr1") |
| starts1 | an array of start1 coordinates |
| ends1 | an array of end1 coordinates. If '-1' chromosome size is assumed. |
| chroms2 | chromosomes2 - an array of strings with or without "chr" prefixes or an array of integers (like: '1' for "chr1"). If 'NULL', 'chroms2' is assumed to be equal to 'chroms1'. |

starts2 an array of start2 coordinates
 ends2 an array of end2 coordinates. If '-1' chromosome size is assumed.

Details

This function returns a set of two-dimensional intervals. The returned value can be used in all functions that accept 'intervals' argument.

Two-dimensional intervals is a data frame whose first six columns are 'chrom1', 'start1', 'end1', 'chrom2', 'start2' and 'end2'. Each row of the data frame represents two genomic intervals from two chromosomes in the range of [start, end). Additional columns can be presented in 2D intervals object yet these columns must be added after the six obligatory ones.

Value

A data frame representing the intervals.

See Also

[gintervals](#), [gintervals.force_range](#)

Examples

```
gdb.init_examples()

## the following 3 calls produce identical results
gintervals.2d(1)
gintervals.2d("1")
gintervals.2d("chrX")

gintervals.2d(1, 1000, 2000, "chrX", 400, 800)
gintervals.2d(c("chr2", "chrX"), 10, c(3000, 5000), 1)
```

gintervals.2d.all *Returns 2D intervals that cover the whole genome*

Description

Returns 2D intervals that cover the whole genome.

Usage

```
gintervals.2d.all()
```

Details

This function returns a set of two-dimensional intervals that cover the whole genome as it is defined by 'chrom_sizes.txt' file.

Value

A data frame representing the intervals.

See Also

[gintervals.2d](#)

gintervals.2d.band_intersect

Intersects two-dimensional intervals with a band

Description

Intersects two-dimensional intervals with a band.

Usage

```
gintervals.2d.band_intersect(  
  intervals = NULL,  
  band = NULL,  
  intervals.set.out = NULL  
)
```

Arguments

| | |
|-------------------|--|
| intervals | two-dimensional intervals |
| band | track expression band. If 'NULL' no band is used. |
| intervals.set.out | intervals set name where the function result is optionally outputted |

Details

This function intersects each two-dimensional interval from 'intervals' with 'band'. If the intersection is not empty, the interval is shrunk to the minimal rectangle that contains the band and added to the return value.

If 'intervals.set.out' is not 'NULL' the result is saved as an intervals set. Use this parameter if the result size exceeds the limits of the physical memory.

Value

If 'intervals.set.out' is 'NULL' a data frame representing the intervals.

See Also

[gintervals.2d](#), [gintervals.intersect](#)

Examples

```
gdb.init_examples()
gintervals.2d.band_intersect(gintervals.2d(1), c(10000, 20000))
```

```
gintervals.2d.convert_to_indexed
```

Convert 2D interval set to indexed format

Description

Converts a per-chromosome interval set to indexed format (intervals2d.dat + intervals2d.idx) which reduces file descriptor usage.

Usage

```
gintervals.2d.convert_to_indexed(  
    set.name = NULL,  
    remove.old = FALSE,  
    force = FALSE  
)
```

Arguments

| | |
|------------|---|
| set.name | name of 2D interval set to convert |
| remove.old | if TRUE, removes old per-chromosome files after successful conversion |
| force | if TRUE, re-converts even if already in indexed format |

Details

The indexed format stores all chromosome pairs in a single intervals2d.dat file with an intervals2d.idx index file. This dramatically reduces file descriptor usage, especially for genomes with many chromosomes ($N*(N-1)/2$ files to just 2).

Only non-empty pairs are stored in the index, avoiding $O(N^2)$ space overhead.

The conversion process:

1. Scans directory for existing per-pair files
2. Creates temporary intervals2d.dat.tmp and intervals2d.idx.tmp files
3. Concatenates all per-pair files into intervals2d.dat.tmp
4. Builds index with pair offsets and checksums
5. Atomically renames temporary files to final names
6. Optionally removes old per-pair files

The indexed format is 100

Value

invisible NULL

Examples

```
## Not run:  
# Convert a 2D interval set  
gintervals.2d.convert_to_indexed("my_2d_intervals")  
  
# Convert and remove old files  
gintervals.2d.convert_to_indexed("my_2d_intervals", remove.old = TRUE)  
  
# Force re-conversion  
gintervals.2d.convert_to_indexed("my_2d_intervals", force = TRUE)  
  
## End(Not run)
```

| | |
|----------------|---|
| gintervals.all | <i>Returns 1D intervals that cover the whole genome</i> |
|----------------|---|

Description

Returns 1D intervals that cover the whole genome.

Usage

```
gintervals.all()
```

Details

This function returns a set of one-dimensional intervals that cover the whole genome as it is defined by 'chrom_sizes.txt' file.

Value

A data frame representing the intervals.

See Also

[gintervals](#)

`gintervals.annotate` *Annotates 1D intervals using nearest neighbors*

Description

Annotates one-dimensional intervals by finding nearest neighbors in another set of intervals and adding selected columns from the neighbors to the original intervals.

Usage

```
gintervals.annotate(
  intervals,
  annotation_intervals,
  annotation_columns = NULL,
  column_names = NULL,
  dist_column = "dist",
  max_dist = Inf,
  na_value = NA,
  maxneighbors = 1,
  tie_method = c("first", "min.start", "min.end"),
  overwrite = FALSE,
  keep_order = TRUE,
  intervals.set.out = NULL,
  ...
)
```

Arguments

| | |
|-----------------------------------|---|
| <code>intervals</code> | Intervals to annotate (1D). |
| <code>annotation_intervals</code> | Source intervals containing annotation data (1D). |
| <code>annotation_columns</code> | Character vector of column names to copy from <code>annotation_intervals</code> . If NULL (default), all non-basic columns are used, i.e. everything beyond the coordinate/strand columns among: <code>chrom</code> , <code>start</code> , <code>end</code> , <code>chrom1</code> , <code>start1</code> , <code>end1</code> , <code>chrom2</code> , <code>start2</code> , <code>end2</code> , <code>strand</code> . |
| <code>column_names</code> | Optional custom names for the annotation columns. If provided, must have the same length as <code>annotation_columns</code> . Defaults to using the original names. |
| <code>dist_column</code> | Name of the distance column to include. Use NULL to omit the distance column. Defaults to "dist". |
| <code>max_dist</code> | Maximum absolute distance. When finite, neighbors with $ dist > max_dist$ result in annotation columns being set to <code>na_value</code> for those rows, while the row itself is retained. |
| <code>na_value</code> | Value(s) to use for annotations when beyond <code>max_dist</code> or when no neighbor is found. Can be a single scalar recycled for all columns, or a named list/vector supplying per-column values matching <code>column_names</code> . |

| | |
|-------------------|---|
| maxneighbors | Maximum number of neighbors per interval (duplicates intervals as needed). Defaults to 1. |
| tie_method | Tie-breaking when distances are equal: one of "first" (arbitrary but stable), "min.start" (smaller neighbor start first), or "min.end" (smaller neighbor end first). Applies when maxneighbors > 1. |
| overwrite | When FALSE (default), errors if selected annotation columns would overwrite existing columns in intervals. When TRUE, conflicting base columns are replaced by the annotation columns. |
| keep_order | If TRUE (default), preserves the original order of intervals rows in the output. |
| intervals.set.out | intervals set name where the function result is optionally outputted |
| ... | Additional arguments forwarded to gintervals.neighbors (e.g., mindist, maxdist). |

Details

The function wraps and extends `gintervals.neighbors` to provide convenient column selection/renaming, optional distance inclusion, distance thresholding with custom NA values, multiple neighbors per interval, and deterministic tie-breaking. Currently supports 1D intervals only.

- When `annotation_columns = NULL`, all non-basic columns present in `annotation_intervals` are included. - Setting `dist_column = NULL` omits the distance column. - If no neighbor is found for an interval, annotation columns are filled with `na_value` and the distance (when present) is `NA_real_`. - Column name collisions are handled as follows: when `overwrite=FALSE` a clear error is emitted; when `overwrite=TRUE`, base columns with the same names are replaced by annotation columns.

Value

A data frame containing the original intervals plus the requested annotation columns (and optional distance column). If `maxneighbors > 1`, rows may be duplicated per input interval to accommodate multiple neighbors.

Examples

```
# Prepare toy data
intervs <- gintervals(1, c(1000, 5000), c(1100, 5050))
ann <- gintervals(1, c(900, 5400), c(950, 5500))
ann$remark <- c("a", "b")
ann$score <- c(10, 20)

# Basic usage with default columns (all non-basic columns)
gintervals.annotate(intervs, ann)

# Select specific columns, with custom names and distance column name
gintervals.annotate(
  intervs, ann,
  annotation_columns = c("remark"),
  column_names = c("ann_remark"),
  dist_column = "ann_dist"
)
```

```

# Distance threshold with scalar NA replacement
gintervals.annotate(
  intervals, ann,
  annotation_columns = c("remark"),
  max_dist = 200,
  na_value = "no_ann"
)

# Multiple neighbors with deterministic tie-breaking
nbrs <- gintervals.annotate(
  gintervals(1, 1000, 1100),
  {
    x <- gintervals(1, c(800, 1200), c(900, 1300))
    x$label <- c("left", "right")
    x
  },
  annotation_columns = "label",
  maxneighbors = 2,
  tie_method = "min.start"
)
nbrs

# Overwrite existing columns in the base intervals
intervals2 <- intervals
intervals2$remark <- c("orig1", "orig2")
gintervals.annotate(intervals2, ann, annotation_columns = "remark", overwrite = TRUE)

```

`gintervals.as_chain` *Transforms existing intervals to a chain format*

Description

Transforms existing intervals to a chain format by validating required columns and adding chain attributes.

Usage

```

gintervals.as_chain(
  intervals = NULL,
  src_overlap_policy = "error",
  tgt_overlap_policy = "auto",
  min_score = NULL
)

```

Arguments

`intervals` a data frame with chain columns: `chrom`, `start`, `end`, `strand`, `chromsrc`, `startsrc`, `endsrc`, `strandsrc`, `chain_id`, `score`

```

src_overlap_policy      source overlap policy: "error", "keep", or "discard"
tgt_overlap_policy      target overlap policy: "error", "auto", "auto_first", "auto_longer", "auto_score",
                        "discard", "keep", or "agg"
min_score               optional minimum alignment score threshold

```

Details

This function checks that the input intervals data frame has all the required columns for a chain format and adds the necessary attributes. A chain format requires both target coordinates (chrom, start, end, strand) and source coordinates (chromsrc, startsrc, endsrc, strandsrc), as well as chain_id and score columns.

Value

A data frame in chain format with chain attributes set

See Also

[gintervals.load_chain](#), [gintervals.liftover](#)

Examples

```

gdb.init_examples()

# Create a chain from existing intervals
chain_data <- data.frame(
  chrom = "chr1",
  start = 1000,
  end = 2000,
  strand = 0,
  chromsrc = "chr1",
  startsrc = 5000,
  endsrc = 6000,
  strandsrc = 0,
  chain_id = 1L,
  score = 1000.0
)
chain <- gintervals.as_chain(chain_data)

```

```
gintervals.attr.export
```

Returns interval set attributes values

Description

Returns interval set attributes values.

Usage

```
gintervals.attr.export(intervals.set = NULL, attrs = NULL)
```

Arguments

`intervals.set` a vector of interval set names or 'NULL'
`attrs` a vector of attribute names or 'NULL'

Details

This function returns a data frame that contains interval set attribute values. Column names of the data frame consist of the attribute names, row names contain the interval set names.

The list of required interval sets is specified by 'intervals.set' argument. If 'intervals.set' is 'NULL' the attribute values of all existing interval sets are returned.

Likewise the list of required attributes is controlled by 'attrs' argument. If 'attrs' is 'NULL' all attribute values of the specified interval sets are returned. The columns are sorted then by "popularity" of an attribute, i.e. the number of interval sets containing this attribute. This sorting is not applied if 'attrs' is not 'NULL'.

Empty character string in a table cell marks a non-existing attribute.

Value

A data frame containing interval set attribute values.

See Also

[gintervals.attr.import](#), [gintervals.attr.get](#), [gintervals.attr.set](#)

Examples

```
gdb.init_examples()
gintervals.attr.set("annotations", "test_attr", "value1")
gintervals.attr.export()
gintervals.attr.export(intervals.set = "annotations")
gintervals.attr.export(attrs = "test_attr")
gintervals.attr.set("annotations", "test_attr", "")
```

gintervals.attr.get *Returns value of an interval set attribute*

Description

Returns value of an interval set attribute.

Usage

```
gintervals.attr.get(intervals.set = NULL, attr = NULL)
```

Arguments

| | |
|---------------|-------------------|
| intervals.set | interval set name |
| attr | attribute name |

Details

This function returns the value of an interval set attribute. If the attribute does not exist an empty string is returned.

Value

Interval set attribute value (character string).

See Also

[gintervals.attr.set](#), [gintervals.attr.export](#), [gintervals.attr.import](#)

Examples

```
gdb.init_examples()
gintervals.attr.set("annotations", "test_attr", "value")
gintervals.attr.get("annotations", "test_attr")
gintervals.attr.set("annotations", "test_attr", "")
```

`gintervals.attr.import`*Imports interval set attributes values*

Description

Imports interval set attributes values.

Usage

```
gintervals.attr.import(table = NULL, remove.others = FALSE)
```

Arguments

`table` a data frame containing attribute values

`remove.others` specifies what to do with the attributes that are not in the table

Details

This function imports attribute values contained in a data frame 'table'. The format of a table is similar to the one returned by 'gintervals.attr.export'. The values of the table must be character strings. Column names of the table should specify the attribute names, while row names should contain the interval set names.

The specified attributes of the specified interval sets are modified. If an attribute value is an empty string this attribute is removed from the interval set.

If 'remove.others' is 'TRUE' all attributes that do not appear in the table are removed, otherwise they are preserved unchanged.

Value

None.

See Also

[gintervals.attr.export](#), [gintervals.attr.set](#), [gintervals.attr.get](#)

Examples

```
gdb.init_examples()
t <- data.frame(
  myattr = "val1", row.names = "annotations",
  stringsAsFactors = FALSE
)
gintervals.attr.import(t)
gintervals.attr.export(attrs = "myattr")

# roll-back the changes
```

```
t$myattr <- ""  
gintervals.attr.import(t)
```

gintervals.attr.set *Assigns value to an interval set attribute*

Description

Assigns value to an interval set attribute.

Usage

```
gintervals.attr.set(intervals.set = NULL, attr = NULL, value = NULL)
```

Arguments

| | |
|---------------|--------------------------|
| intervals.set | interval set name |
| attr | attribute name |
| value | value (character string) |

Details

This function creates an interval set attribute and assigns 'value' to it. If the attribute already exists its value is overwritten.

If 'value' is an empty string the attribute is removed.

Value

None.

See Also

[gintervals.attr.get](#), [gintervals.attr.export](#), [gintervals.attr.import](#)

Examples

```
gdb.init_examples()  
gintervals.attr.set("annotations", "test_attr", "value")  
gintervals.attr.get("annotations", "test_attr")  
gintervals.attr.set("annotations", "test_attr", "")
```

gintervals.canonic *Converts intervals to canonic form*

Description

Converts intervals to canonic form.

Usage

```
gintervals.canonic(intervals = NULL, unify_touching_intervals = TRUE)
```

Arguments

intervals intervals to be converted
unify_touching_intervals
 if 'TRUE', touching one-dimensional intervals are unified

Details

This function converts 'intervals' into a "canonic" form: properly sorted with no overlaps. The result can be used later in the functions that require the intervals to be in canonic form. Use 'unify_touching_intervals' to control whether the intervals that touch each other (i.e. the end coordinate of one equals to the start coordinate of the other) are unified. 'unify_touching_intervals' is ignored if two-dimensional intervals are used.

Since 'gintervals.canonic' unifies overlapping or touching intervals, the number of the returned intervals might be less than the number of the original intervals. To allow the user to find the origin of the new interval 'mapping' attribute is attached to the result. It maps between the original intervals and the resulted intervals. Use 'attr(retv_of_gintervals.canonic, "mapping")' to retrieve the map.

Value

A data frame representing the canonic intervals and an attribute 'mapping' that maps the original intervals to the resulted ones.

See Also

[gintervals](#), [gintervals.2d](#)

Examples

```
gdb.init_examples()

## Create intervals manually by using 'data.frame'.
## Note that we add an additional column 'data'.
## Return value:
##  chrom start  end data
```

```
## 1 chr1 11000 12000 10
## 2 chr1 100 200 20
## 3 chr1 10000 13000 30
## 4 chr1 10500 10600 40
intervs <- data.frame(
  chrom = "chr1",
  start = c(11000, 100, 10000, 10500),
  end = c(12000, 200, 13000, 10600),
  data = c(10, 20, 30, 40)
)

## Convert the intervals into the canonic form.
## The function discards any columns besides chrom, start and end.
## Return value:
## chrom start end
## 1 chr1 100 200
## 2 chr1 10000 13000
res <- gintervals.canonic(intervs)

## By inspecting mapping attribute we can see how the new
## intervals were created: "2 1 2 2" means that the first
## interval in the result was created from the second interval in
## the original set (we look for the indices in mapping where "1"
## appears). Likewise the second interval in the result was
## created from 3 intervals in the original set. Their indices are
## 1, 3 and 4 (once again we look for the indices in mapping where
## "2" appears).
## Return value:
## 2 1 2 2
attr(res, "mapping")

## Finally (and that is the most useful part of 'mapping'
## attribute): we add a new column 'data' to our result which is
## the mean value of the original data column. The trick is done
## using 'tapply' on par with 'mapping' attribute. For example,
## 20.00000 equals is a result of 'mean(intervs[2,]$data' while
## 26.66667 is a result of 'mean(intervs[c(1,3,4),]$data)'.
## 'res' after the following call:
## chrom start end data
## 1 chr1 100 200 20.00000
## 2 chr1 10000 13000 26.66667
res$data <- tapply(intervs$data, attr(res, "mapping"), mean)
```

gintervals.chrom_sizes

Returns number of intervals per chromosome

Description

Returns number of intervals per chromosome (or chromosome pair).

Usage

```
gintervals.chrom_sizes(intervals = NULL)
```

Arguments

```
intervals      intervals set
```

Details

This function returns number of intervals per chromosome (for 1D intervals) or chromosome pair (for 2D intervals).

Value

Data frame representing number of intervals per chromosome (for 1D intervals) or chromosome pair (for 2D intervals).

See Also

[gintervals.load](#), [gintervals.save](#), [gintervals.exists](#), [gintervals.ls](#), [gintervals](#), [gintervals.2d](#)

Examples

```
gdb.init_examples()  
gintervals.chrom_sizes("annotations")
```

```
gintervals.convert_to_indexed
```

Convert 1D interval set to indexed format

Description

Converts a per-chromosome interval set to indexed format (intervals.dat + intervals.idx) which reduces file descriptor usage.

Usage

```
gintervals.convert_to_indexed(  
  set.name = NULL,  
  remove.old = FALSE,  
  force = FALSE  
)
```

Arguments

| | |
|-------------------------|---|
| <code>set.name</code> | name of interval set to convert |
| <code>remove.old</code> | if TRUE, removes old per-chromosome files after successful conversion |
| <code>force</code> | if TRUE, re-converts even if already in indexed format |

Details

The indexed format stores all chromosomes in a single `intervals.dat` file with an `intervals.idx` index file. This reduces file descriptor usage from N files (one per chromosome) to just 2 files.

The conversion process:

1. Creates temporary `intervals.dat.tmp` and `intervals.idx.tmp` files
2. Concatenates all per-chromosome files into `intervals.dat.tmp`
3. Builds index with offsets and checksums
4. Atomically renames temporary files to final names
5. Optionally removes old per-chromosome files

The indexed format is 100

Value

invisible NULL

See Also

[gintervals.save](#), [gintervals.load](#)

Examples

```
## Not run:  
# Convert an interval set  
gintervals.convert_to_indexed("my_intervals")  
  
# Convert and remove old files  
gintervals.convert_to_indexed("my_intervals", remove.old = TRUE)  
  
# Force re-conversion  
gintervals.convert_to_indexed("my_intervals", force = TRUE)  
  
## End(Not run)
```

`gintervals.coverage_fraction`*Calculate fraction of genomic space covered by intervals*

Description

Returns the fraction of a genomic space that is covered by a set of intervals.

Usage

```
gintervals.coverage_fraction(intervals1 = NULL, intervals2 = NULL)
```

Arguments

| | |
|-------------------------|---|
| <code>intervals1</code> | set of one-dimensional intervals (the covering set) |
| <code>intervals2</code> | set of one-dimensional intervals to be covered (default: <code>NULL</code> , meaning the entire genome) |

Details

This function calculates what fraction of 'intervals2' is covered by 'intervals1'. If 'intervals2' is `NULL`, it calculates the fraction of the entire genome that is covered by 'intervals1'. Overlapping intervals in either set are automatically unified before calculation.

Value

A single numeric value between 0 and 1 representing the fraction of 'intervals2' (or the genome) covered by 'intervals1'.

See Also

[gintervals](#), [gintervals.intersect](#), [gintervals.covered_bp](#), [gintervals.all](#)

Examples

```
gdb.init_examples()

# Create some intervals
intervs1 <- gscreen("dense_track > 0.15")
intervs2 <- gintervals(c("chr1", "chr2"), 0, c(100000, 100000))

# Calculate fraction of interv2 covered by interv1
gintervals.coverage_fraction(intervs1, interv2)

# Calculate fraction of entire genome covered by interv1
gintervals.coverage_fraction(intervs1)
```

`gintervals.covered_bp` *Calculate total base pairs covered by intervals*

Description

Returns the total number of base pairs covered by a set of intervals.

Usage

```
gintervals.covered_bp(intervals = NULL)
```

Arguments

`intervals` set of one-dimensional intervals

Details

This function first canonicalizes the intervals to remove overlaps and touching intervals, then sums up the lengths of all resulting intervals. Overlapping intervals are counted only once.

Value

A single numeric value representing the total number of base pairs covered by the intervals.

See Also

[gintervals](#), [gintervals.canonic](#), [gintervals.coverage_fraction](#)

Examples

```
gdb.init_examples()

# Create some intervals
intervs <- gintervals(
  c("chr1", "chr1", "chr2"),
  c(100, 150, 1000),
  c(200, 250, 2000)
)

# Calculate total bp covered
# Note: intervals [100,200) and [150,250) overlap,
# so total is (200-100) + (250-150) + (2000-1000) = 100 + 100 + 1000 = 1200
# But after canonicalization: [100,250) + [1000,2000) = 150 + 1000 = 1150
gintervals.covered_bp(intervs)
```

`gintervals.dataset` *Returns the database/dataset path for interval sets*

Description

Returns the path of the database or dataset containing an interval set.

Usage

```
gintervals.dataset(intervals = NULL)
```

Arguments

`intervals` interval set name or a vector of interval set names

Details

When datasets are loaded, interval sets can come from either the working database or from loaded datasets. This function returns the source path for each interval set.

Value

A character vector containing the database paths for each interval set. Returns NA for interval sets that don't exist in any connected database.

See Also

[gintervals.dbs](#), [gintervals.exists](#), [gintervals.ls](#), [gdataset.ls](#)

Examples

```
gdb.init_examples()  
gintervals.dataset("annotations1")
```

`gintervals.dbs` *Returns all database paths containing an interval set*

Description

Returns all database paths that contain a version of an interval set.

Usage

```
gintervals.dbs(intervals = NULL, dataframe = FALSE)
```

Arguments

| | |
|-----------|---|
| intervals | interval set name |
| dataframe | return a data frame with columns intervals and db |

Details

When datasets are loaded, an interval set may exist in multiple locations. This function computes on-demand and returns all such paths.

Value

A named character vector of database paths. If dataframe is TRUE, returns a data frame with columns intervals and db.

See Also

[gintervals.dataset](#), [gintervals.ls](#), [gdataset.ls](#)

Examples

```
gdb.init_examples()
gintervals.dbs("annotations1")
```

| | |
|-----------------|--|
| gintervals.diff | <i>Calculates difference of two intervals sets</i> |
|-----------------|--|

Description

Returns difference of two sets of intervals.

Usage

```
gintervals.diff(intervals1 = NULL, intervals2 = NULL, intervals.set.out = NULL)
```

Arguments

| | |
|------------------------|--|
| intervals1, intervals2 | set of one-dimensional intervals |
| intervals.set.out | intervals set name where the function result is optionally outputted |

Details

This function returns a genomic space that is covered by 'intervals1' but not covered by 'intervals2'. If 'intervals.set.out' is not 'NULL' the result is saved as an intervals set. Use this parameter if the result size exceeds the limits of the physical memory.

Value

If 'intervals.set.out' is 'NULL' a data frame representing the intervals.

See Also

[gintervals](#), [gintervals.intersect](#), [gintervals.union](#)

Examples

```
gdb.init_examples()

intervs1 <- gscreen("dense_track > 0.15")
intervs2 <- gscreen("dense_track < 0.2")

## 'res3' equals to 'res4'
res3 <- gintervals.diff(intervs1, intervs2)
res4 <- gscreen("dense_track >= 0.2")
```

| | |
|-------------------|--|
| gintervals.exists | <i>Tests for a named intervals set existence</i> |
|-------------------|--|

Description

Tests for a named intervals set existence.

Usage

```
gintervals.exists(intervals.set = NULL)
```

Arguments

intervals.set name of an intervals set

Details

This function returns 'TRUE' if a named intervals set exists in Genomic Database.

Value

'TRUE' if a named intervals set exists. Otherwise 'FALSE'.

See Also

[gintervals.ls](#), [gintervals.load](#), [gintervals.rm](#), [gintervals.save](#), [gintervals](#), [gintervals.2d](#)

Examples

```
gdb.init_examples()  
gintervals.exists("annotations")
```

`gintervals.force_range`

Limits intervals to chromosomal range

Description

Limits intervals to chromosomal range.

Usage

```
gintervals.force_range(intervals = NULL, intervals.set.out = NULL)
```

Arguments

| | |
|--------------------------------|--|
| <code>intervals</code> | <code>intervals</code> |
| <code>intervals.set.out</code> | intervals set name where the function result is optionally outputted |

Details

This function enforces the intervals to be within the chromosomal range [0, chrom length) by altering the intervals' boundaries. Intervals that lay entirely outside of the chromosomal range are eliminated. The new intervals are returned.

If 'intervals.set.out' is not 'NULL' the result is saved as an intervals set. Use this parameter if the result size exceeds the limits of the physical memory.

Value

If 'intervals.set.out' is 'NULL' a data frame representing the intervals.

See Also

[gintervals](#), [gintervals.2d](#), [gintervals.canonic](#)

Examples

```
gdb.init_examples()
intervs <- data.frame(
  chrom = "chr1",
  start = c(11000, -100, 10000, 10500),
  end = c(12000, 200, 13000000, 10600)
)
gintervals.force_range(intervs)
```

```
gintervals.import_genes
```

Imports genes and annotations from files

Description

Imports genes and annotations from files.

Usage

```
gintervals.import_genes(
  genes.file = NULL,
  annots.file = NULL,
  annots.names = NULL
)
```

Arguments

| | |
|--------------|---|
| genes.file | name or URL of file that contains genes |
| annots.file | name of URL file that contains annotations. If 'NULL' no annotations are imported |
| annots.names | annotations names |

Details

This function reads a definition of genes from 'genes.file' and returns four sets of intervals: TSS, exons, 3utr and 5utr. In addition to the regular intervals columns 'strand' column is added. It contains '1' values for '+' strands and '-1' values for '-' strands.

If annotation file 'annots.file' is given then annotations are attached too to the intervals. The names of the annotations as they would appear in the return value must be specified in 'annots.names' argument.

Both 'genes.file' and 'annots.file' can be either a file path or URL in a form of 'ftp://[address]/[file]'. Files that these arguments point to can be zipped or unzipped.

Examples of 'genes.file' and 'annots.file' can be found here:

`ftp://hgdownload.soe.ucsc.edu/goldenPath/hg19/database/knownGene.txt.gz` `ftp://hgdownload.soe.ucsc.edu`

If a few intervals overlap (for example: two TSS regions) they are all unified to an interval that covers the whole overlapping region. 'strand' value is set to '0' if two or more of the overlapping intervals have different strands. The annotations of the overlapping intervals are concatenated to a single character string separated by semicolons. Identical values of overlapping intervals' annotation are eliminated.

Value

A list of four intervals sets named 'tss', 'exons', 'utr3' and 'utr5'. 'strand' column and annotations are attached to the intervals.

See Also

[gintervals](#), [gdb.create](#)

gintervals.intersect *Calculates an intersection of two sets of intervals*

Description

Calculates an intersection of two sets of intervals.

Usage

```
gintervals.intersect(  
  intervals1 = NULL,  
  intervals2 = NULL,  
  intervals.set.out = NULL  
)
```

Arguments

```
intervals1, intervals2  
                set of intervals  
intervals.set.out  
                intervals set name where the function result is optionally outputted
```

Details

This function returns intervals that represent a genomic space which is achieved by intersection of 'intervals1' and 'intervals2'.

If 'intervals.set.out' is not 'NULL' the result is saved as an intervals set. Use this parameter if the result size exceeds the limits of the physical memory.

Value

If 'intervals.set.out' is 'NULL' a data frame representing the intersection of intervals.

See Also

[gintervals.2d.band_intersect](#), [gintervals.diff](#), [gintervals.union](#), [gintervals](#), [gintervals.2d](#)

Examples

```
gdb.init_examples()

intervs1 <- gscreen("dense_track > 0.15")
intervs2 <- gscreen("dense_track < 0.2")

## 'intervs3' and 'intervs4' are identical
intervs3 <- gintervals.intersect(intervs1, intervs2)
intervs4 <- gscreen("dense_track > 0.15 & dense_track < 0.2")
```

`gintervals.is.bigset` *Tests for big intervals set*

Description

Tests for big intervals set.

Usage

```
gintervals.is.bigset(intervals.set = NULL)
```

Arguments

`intervals.set` name of an intervals set

Details

This function tests whether 'intervals.set' is a big intervals set. Intervals set is big if it is stored in big intervals set format and given the current limits it cannot be fully loaded into memory.

Memory limit is controlled by 'gmax.data.size' option (see: 'getOption("gmax.data.size")').

Value

'TRUE' if intervals set is big, otherwise 'FALSE'.

See Also

[gintervals.load](#), [gintervals.save](#), [gintervals.exists](#), [gintervals.ls](#)

Examples

```
gdb.init_examples()
gintervals.is.bigset("annotations")
```

| | |
|---------------------|---|
| gintervals.liftover | <i>Converts intervals from another assembly</i> |
|---------------------|---|

Description

Converts intervals from another assembly to the current one.

Usage

```
gintervals.liftover(
  intervals = NULL,
  chain = NULL,
  src_overlap_policy = "error",
  tgt_overlap_policy = "auto",
  min_score = NULL,
  include_metadata = FALSE,
  canonic = FALSE,
  value_col = NULL,
  multi_target_agg = c("mean", "median", "sum", "min", "max", "count", "first", "last",
    "nth", "max.coverage_len", "min.coverage_len", "max.coverage_frac",
    "min.coverage_frac"),
  params = NULL,
  na.rm = TRUE,
  min_n = NULL
)
```

Arguments

| | |
|--------------------|--|
| intervals | intervals from another assembly |
| chain | name of chain file or data frame as returned by 'gintervals.load_chain' |
| src_overlap_policy | policy for handling source overlaps: "error" (default), "keep", or "discard". "keep" allows one source interval to map to multiple target intervals, "discard" discards all source intervals that have overlaps and "error" throws an error if source overlaps are detected. |
| tgt_overlap_policy | policy for handling target overlaps. One of: |

| Policy | Description |
|--------|--|
| error | Throws an error if any target overlaps are detected. |

| | |
|----------------------------------|---|
| <code>auto</code> | Default. Alias for "auto_score". |
| <code>auto_score</code> | Resolves overlaps by segmenting the target region and selecting the best chain for each segment based on score. |
| <code>auto_longer</code> | Resolves overlaps by segmenting and selecting the chain with the longest span for each segment. Tie-breaker is score. |
| <code>auto_first</code> | Resolves overlaps by segmenting and selecting the chain with the lowest chain_id for each segment. |
| <code>keep</code> | Preserves all overlapping intervals. |
| <code>discard</code> | Discards any chain interval that has a target overlap with another chain interval. |
| <code>agg</code> | Segments overlaps into smaller disjoint regions where each region contains all contributing chains, allowing for aggregation. |
| <code>best_source_cluster</code> | Best source cluster strategy based on source overlap. When multiple chains map a source interval, cluster the best one. |

| | |
|-------------------------------|---|
| <code>min_score</code> | optional minimum alignment score threshold. Chains with scores below this value are filtered out. Useful for excluding low-quality alignments. |
| <code>include_metadata</code> | logical; if TRUE, adds 'score' column to the output indicating the alignment score of the chain used for each mapping. Only applicable with "auto_score" or "auto" policy. |
| <code>canonic</code> | logical; if TRUE, merges adjacent target intervals that originated from the same source interval (same intervalID) and same chain (same chain_id). This is useful when a source interval maps to multiple adjacent target blocks due to chain gaps. |
| <code>value_col</code> | optional character string specifying the name of a numeric column in the intervals data frame to track through the liftover. When specified, this column's values are preserved in the output with the same column name. Use with <code>multi_target_agg</code> to aggregate values when multiple source intervals map to overlapping target regions. |
| <code>multi_target_agg</code> | aggregation method to use when <code>value_col</code> is specified. One of: "mean", "median", "sum", "min", "max", "count", "first", "last", "nth", "max.coverage_len", "min.coverage_len", "max.coverage_frac", "min.coverage_frac". Default: "mean". Ignored when <code>value_col</code> is NULL. |
| <code>params</code> | additional parameters for specific aggregation methods. Currently only used for "nth" aggregation, where it specifies which element to select (e.g., <code>params = 2</code> for second element, or <code>params = list(n = 2)</code>). |
| <code>na.rm</code> | logical; if TRUE (default), NA values are removed before aggregation. If FALSE, any NA in the values will cause the result to be NA. Only used when <code>value_col</code> is specified. |
| <code>min_n</code> | optional minimum number of non-NA observations required for aggregation. If fewer observations are available, the result is NA. NULL (default) means no minimum. Only used when <code>value_col</code> is specified. |

Details

This function converts 'intervals' from another assembly to the current one. Chain file instructs how the conversion of coordinates should be done. It can be either a name of a chain file or a data frame in the same format as returned by `'gintervals.load_chain'` function.

The converted intervals are returned. An additional column named 'intervalID' is added to the resulted data frame. For each interval in the resulted intervals it indicates the index of the original interval.

Note: When passing a pre-loaded chain (data frame), overlap policies cannot be specified - they are taken from the chain's attributes that were set during loading. When passing a chain file path, policies can be specified and will be used for loading.

Value

A data frame representing the converted intervals. For 1D intervals, always includes 'intervalID' (index of original interval) and 'chain_id' (identifier of the chain that produced the mapping) columns. The chain_id column is essential for distinguishing results when a source interval maps to multiple target regions via different chains (duplications). When include_metadata=TRUE, also includes 'score' column. When value_col is specified, includes the value column with its original name.

See Also

[gintervals.load_chain](#), [gtrack.liftover](#), [gintervals](#)

Examples

```
gdb.init_examples()
chainfile <- paste(.misha$GROOT, "data/test.chain", sep = "/")
intervs <- data.frame(
  chrom = "chr25", start = c(0, 7000),
  end = c(6000, 20000)
)
# Liftover with default policies
gintervals.liftover(intervs, chainfile)

# Liftover keeping source overlaps (one source interval may map to multiple targets)
# gintervals.liftover(intervs, chainfile, src_overlap_policy = "keep")
```

| | |
|-----------------|------------------------------------|
| gintervals.load | <i>Loads a named intervals set</i> |
|-----------------|------------------------------------|

Description

Loads a named intervals set.

Usage

```
gintervals.load(
  intervals.set = NULL,
  chrom = NULL,
  chrom1 = NULL,
  chrom2 = NULL
)
```

Arguments

| | |
|----------------------------|--|
| <code>intervals.set</code> | name of an intervals set |
| <code>chrom</code> | chromosome for 1D intervals set |
| <code>chrom1</code> | first chromosome for 2D intervals set |
| <code>chrom2</code> | second chromosome for 2D intervals set |

Details

This function loads and returns intervals stored in a named intervals set.

If intervals set contains 1D intervals and 'chrom' is not 'NULL' only the intervals of the given chromosome are returned.

Likewise if intervals set contains 2D intervals and 'chrom1', 'chrom2' are not 'NULL' only the intervals of the given pair of chromosomes are returned.

For big intervals sets 'chrom' parameter (1D case) / 'chrom1', 'chrom2' parameters (2D case) must be specified. In other words: big intervals sets can be loaded only by chromosome or chromosome pair.

Value

A data frame representing the intervals.

See Also

[gintervals.save](#), [gintervals.is.bigset](#), [gintervals.exists](#), [gintervals.ls](#), [gintervals](#), [gintervals.2d](#)

Examples

```
gdb.init_examples()  
gintervals.load("annotations")
```

`gintervals.load_chain` *Loads assembly conversion table from a chain file*

Description

Loads assembly conversion table from a chain file.

Usage

```
gintervals.load_chain(
    file = NULL,
    src_overlap_policy = "error",
    tgt_overlap_policy = "auto",
    src_groot = NULL,
    min_score = NULL
)
```

Arguments

`file` name of chain file

`src_overlap_policy` policy for handling source overlaps: "error" (default), "keep", or "discard". "keep" allows one source interval to map to multiple target intervals, "discard" discards all source intervals that have overlaps and "error" throws an error if source overlaps are detected.

`tgt_overlap_policy` policy for handling target overlaps. One of:

| Policy | Description |
|------------------------|--|
| error | Throws an error if any target overlaps are detected. |
| auto | Default. Alias for "auto_score". |
| auto_score | Resolves overlaps by segmenting the target region and selecting the best chain for each segment based on score. |
| auto_longer | Resolves overlaps by segmenting and selecting the chain with the longest span for each segment. Tie-breaker is score. |
| auto_first | Resolves overlaps by segmenting and selecting the chain with the lowest chain_id for each segment. |
| keep | Preserves all overlapping intervals. |
| discard | Discards any chain interval that has a target overlap with another chain interval. |
| agg | Segments overlaps into smaller disjoint regions where each region contains all contributing chains, allowing for a single chain to represent the entire region. |
| best_source_cluster | Best source cluster strategy based on source overlap. When multiple chains map a source interval, cluster the best one. |
| <code>src_groot</code> | optional path to source genome database for validating source chromosomes and coordinates. If provided, the function temporarily switches to this database to verify that all source chromosomes exist and coordinates are within bounds, then restores the original database. |
| <code>min_score</code> | optional minimum alignment score threshold. Chains with scores below this value are filtered out. Useful for excluding low-quality alignments. |

Details

This function reads a file in 'chain' format and returns assembly conversion table that can be used in 'gtrack.liftover' and 'gintervals.liftover'.

Source overlaps occur when the same source genome position maps to multiple target genome positions. Target overlaps occur when multiple source positions map to overlapping regions in the target genome.

The 'src_overlap_policy' controls how source overlaps are handled:

- "error" (default): Throw an error if source overlaps are detected
- "keep": Keep all mappings, allowing one source to map to multiple targets
- "discard": Remove all chain intervals involved in source overlaps

The 'tgt_overlap_policy' controls how target overlaps are handled:

- "error": Throw an error if target overlaps are detected
- "auto" (default) / "auto_first": Keep the first overlapping chain (original file order) by trimming or discarding later overlaps while keeping source/target lengths consistent
- "auto_longer": Keep the longer overlapping chain and trim/drop the shorter ones
- "discard": Remove all chain intervals involved in target overlaps
- "keep": Allow target overlaps to remain untouched (liftover must be able to handle them)

Value

A data frame representing assembly conversion table with columns: chrom, start, end, strand, chromsrc, startsrc, endsrc, strandsrc, chain_id, score.

See Also

[gintervals.liftover](#), [gtrack.liftover](#)

Examples

```
gdb.init_examples()
chainfile <- paste(.misha$GROOT, "data/test.chain", sep = "/")
# Load chain file with default policies
gintervals.load_chain(chainfile)
```

gintervals.ls

Returns a list of named intervals sets

Description

Returns a list of named intervals sets in Genomic Database.

Usage

```
gintervals.ls(
  pattern = "",
  db = NULL,
  ignore.case = FALSE,
  perl = FALSE,
  fixed = FALSE,
  useBytes = FALSE
)
```

Arguments

pattern, ignore.case, perl, fixed, useBytes
 see 'grep'
 db optional database path to filter intervals. If specified, only interval sets from that database are returned.

Details

This function returns a list of named intervals sets that match the pattern (see 'grep'). If called without any arguments all named intervals sets are returned.

When multiple databases are connected, the 'db' parameter can be used to filter intervals to only those from a specific database.

Value

An array that contains the names of intervals sets.

See Also

[grep](#), [gintervals.exists](#), [gintervals.load](#), [gintervals.save](#), [gintervals.rm](#), [gintervals](#), [gintervals.2d](#), [gintervals.dataset](#)

Examples

```
gdb.init_examples()
gintervals.ls()
gintervals.ls(pattern = "annot*")
```

`gintervals.mapply` *Applies a function to values of track expressions*

Description

Applies a function to values of track expressions for each interval.

Usage

```
gintervals.mapply(
  FUN = NULL,
  ...,
  intervals = NULL,
  enable.gapply.intervals = FALSE,
  iterator = NULL,
  band = NULL,
  intervals.set.out = NULL,
  colnames = "value"
)
```

Arguments

| | |
|--------------------------------------|---|
| <code>FUN</code> | function to apply, found via <code>'match.fun'</code> |
| <code>...</code> | track expressions whose values are used as arguments for <code>'FUN'</code> |
| <code>intervals</code> | intervals for which track expressions are calculated |
| <code>enable.gapply.intervals</code> | if <code>'TRUE'</code> , then a variable <code>'GAPPLY.INTERVALS'</code> is available |
| <code>iterator</code> | track expression iterator. If <code>'NULL'</code> iterator is determined implicitly based on track expressions. |
| <code>band</code> | track expression band. If <code>'NULL'</code> no band is used. |
| <code>intervals.set.out</code> | intervals set name where the function result is optionally outputted |
| <code>colnames</code> | name of the column that contains the return values of <code>'FUN'</code> . Default is <code>"value"</code> . |

Details

This function evaluates track expressions for each interval from `'intervals'`. The resulted vectors are passed then as arguments to `'FUN'`.

If the intervals are one-dimensional and have an additional column named `'strand'` whose value is `'-1'`, the values of the track expression are placed to the vector in reverse order.

The current interval index (1-based) is stored in `'GAPPLY.INTERVID'` variable that is available during the execution of `'gintervals.mapply'`. There is no guarantee about the order in which the intervals are processed. Do not rely on any specific order and use `'GITERATOR.INTERVID'` variable to detect the current interval id.

If `'enable.gapply.intervals'` is `'TRUE'`, an additional variable `'GAPPLY.INTERVALS'` is defined during the execution of `'gintervals.mapply'`. This variable stores the current iterator intervals prior to track expression evaluation. Please note that setting `'enable.gapply.intervals'` to `'TRUE'` might severely affect the run-time of the function.

Note: all the changes made in R environment by `'FUN'` will be void if multitasking mode is switched on. One should also refrain from performing any other operations in `'FUN'` that might be not "thread-safe" such as updating files, etc. Please switch off multitasking (`'options(gmultitasking = FALSE)'`) if you wish to perform such operations.

If `'intervals.set.out'` is not `'NULL'` the result is saved as an intervals set. Use this parameter if the result size exceeds the limits of the physical memory.

Value

If `'intervals.set.out'` is `'NULL'` a data frame representing intervals with an additional column that contains the return values of `'FUN'`. The name of this additional column is specified by the `'colnames'` parameter.

See Also

[mapply](#)

Examples

```

gdb.init_examples()
gintervals.mapply(
  max, "dense_track",
  gintervals(c(1, 2), 0, 10000)
)
gintervals.mapply(
  function(x, y) {
    max(x + y)
  }, "dense_track",
  "sparse_track", gintervals(c(1, 2), 0, 10000),
  iterator = "sparse_track"
)
# Using custom column name
gintervals.mapply(
  max, "dense_track",
  gintervals(c(1, 2), 0, 10000),
  colnames = "max_value"
)

```

```
gintervals.mark_overlaps
```

Mark overlapping intervals with a group ID

Description

Mark overlapping intervals with a group ID

Usage

```

gintervals.mark_overlaps(
  intervals,
  group_col = "overlap_group",
  unify_touching_intervals = TRUE
)

```

Arguments

| | |
|--------------------------|--|
| intervals | intervals set |
| group_col | name of the column to store the overlap group IDs (default: "overlap_group") |
| unify_touching_intervals | if 'TRUE', touching one-dimensional intervals are unified |

Value

The intervals set with an additional column containing group IDs from `gintervals.canonic` mapping. All overlapping intervals will have the same group ID.

Examples

```

gdb.init_examples()
# Create sample overlapping intervals
intervs <- data.frame(
  chrom = "chr1",
  start = c(11000, 100, 10000, 10500),
  end = c(12000, 200, 13000, 10600),
  data = c(10, 20, 30, 40)
)

# Mark overlapping intervals
intervs_marked <- gintervals.mark_overlaps(intervs)

# Use custom column name
intervs_marked <- gintervals.mark_overlaps(intervs, group_col = "my_groups")

```

`gintervals.neighbors` *Finds neighbors between two sets of intervals*

Description

For each interval in 'intervals1', finds the closest intervals from 'intervals2'. Distance directionality can be determined by either the strand of the target intervals (intervals2, default) or the query intervals (intervals1). When no strand column is present, all intervals are treated as positive strand (strand = 1).

Usage

```

gintervals.neighbors(
  intervals1 = NULL,
  intervals2 = NULL,
  maxneighbors = 1,
  mindist = -1e+09,
  maxdist = 1e+09,
  mindist1 = -1e+09,
  maxdist1 = 1e+09,
  mindist2 = -1e+09,
  maxdist2 = 1e+09,
  na.if.notfound = FALSE,
  use_intervals1_strand = FALSE,
  warn.ignored.strand = TRUE,
  intervals.set.out = NULL
)

```


The names of the returned columns are made unique using `make.unique(colnames(df), sep = "")`, assuming 'df' is the result.

If 'intervals.set.out' is not 'NULL' the result is saved as an intervals set. Use this parameter if the result size exceeds the limits of the physical memory.

Value

If 'intervals.set.out' is 'NULL' a data frame containing the pairs of intervals from 'intervals1', intervals from 'intervals2' and an additional column named 'dist' ('dist1' and 'dist2' for 2D intervals) representing the distance between the corresponding intervals. The intervals from intervals2 would be changed to 'chrom1', 'start1', and 'end1' and for 2D intervals chrom11, start11, end11 and chrom22, start22, end22. If 'na.if.notfound' is 'TRUE', the data frame contains all the intervals from 'intervals1' including those for which no matching neighbor was found. For the latter intervals an 'NA' neighboring interval is stated. If 'na.if.notfound' is 'FALSE', the data frame contains only intervals from 'intervals1' for which matching neighbor(s) was found.

See Also

[gintervals](#), [gintervals.neighbors.upstream](#), [gintervals.neighbors.downstream](#)

Examples

```
gdb.init_examples()

# Basic intervals
intervs1 <- giterator.intervals("dense_track",
  gintervals(1, 0, 4000),
  iterator = 233
)
intervs2 <- giterator.intervals(
  "sparse_track",
  gintervals(1, 0, 2000)
)

# Original behavior - no strand considerations
gintervals.neighbors(intervs1, intervs2, 10,
  mindist = -300,
  maxdist = 500
)

# Add strand to intervals2 - affects distance directionality (original behavior)
intervs2$strand <- c(1, 1, -1, 1)
gintervals.neighbors(intervs1, intervs2, 10,
  mindist = -300,
  maxdist = 500
)

# TSS analysis example - use intervals1 (TSS) strand for directionality
tss <- data.frame(
  chrom = c("chr1", "chr1", "chr1"),
```

```
    start = c(1000, 2000, 3000),
    end = c(1001, 2001, 3001),
    strand = c(1, -1, 1), # +, -, +
    gene = c("GeneA", "GeneB", "GeneC")
  )

features <- data.frame(
  chrom = "chr1",
  start = c(500, 800, 1200, 1800, 2200, 2800, 3200),
  end = c(600, 900, 1300, 1900, 2300, 2900, 3300),
  feature_id = paste0("F", 1:7)
)

# Use TSS strand for distance directionality
result <- gintervals.neighbors(tss, features,
  maxneighbors = 2,
  mindist = -1000, maxdist = 1000,
  use_intervals1_strand = TRUE
)

# Convenience functions for common TSS analysis
# Find upstream neighbors (negative distances for + strand genes)
upstream <- gintervals.neighbors.upstream(tss, features,
  maxneighbors = 2, maxdist = 1000
)

# Find downstream neighbors (positive distances for + strand genes)
downstream <- gintervals.neighbors.downstream(tss, features,
  maxneighbors = 2, maxdist = 1000
)

# Find both directions
both_directions <- gintervals.neighbors.directional(tss, features,
  maxneighbors_upstream = 1,
  maxneighbors_downstream = 1,
  maxdist = 1000
)
```

gintervals.neighbors.upstream

Directional neighbor finding functions

Description

These functions find neighbors using query strand directionality, where upstream/downstream directionality is determined by the strand of the query intervals rather than the target intervals. This is particularly useful for TSS analysis where you want distances relative to gene direction.

Usage

```

gintervals.neighbors.upstream(
    query_intervals,
    target_intervals,
    maxneighbors = 1,
    maxdist = 1e+09,
    ...
)

gintervals.neighbors.downstream(
    query_intervals,
    target_intervals,
    maxneighbors = 1,
    maxdist = 1e+09,
    ...
)

gintervals.neighbors.directional(
    query_intervals,
    target_intervals,
    maxneighbors_upstream = 1,
    maxneighbors_downstream = 1,
    maxdist = 1e+09,
    ...
)

```

Arguments

| | |
|--------------------------------------|--|
| <code>query_intervals</code> | intervals with strand information (query intervals) |
| <code>target_intervals</code> | intervals to search for neighbors |
| <code>maxneighbors</code> | maximum number of neighbors per query interval (default: 1) |
| <code>maxdist</code> | maximum distance to search (default: 1e+09) |
| <code>...</code> | additional arguments passed to <code>gintervals.neighbors</code> |
| <code>maxneighbors_upstream</code> | maximum upstream neighbors per query interval (default: 1) |
| <code>maxneighbors_downstream</code> | maximum downstream neighbors per query interval (default: 1) |

Details

****Distance interpretation:****

- ****Positive strand queries:**** upstream distances < 0, downstream distances > 0
- ****Negative strand queries:**** upstream distances > 0, downstream distances < 0

If no strand column is present, all intervals are treated as positive strand.

Value

gintervals.neighbors.upstream data frame of upstream neighbors

gintervals.neighbors.downstream data frame of downstream neighbors

gintervals.neighbors.directional list with 'upstream' and 'downstream' components

See Also

[gintervals.neighbors](#)

Examples

```
gdb.init_examples()

# Create TSS intervals with strand information
tss <- data.frame(
  chrom = c("chr1", "chr1", "chr1"),
  start = c(1000, 2000, 3000),
  end = c(1001, 2001, 3001),
  strand = c(1, -1, 1), # +, -, +
  gene = c("GeneA", "GeneB", "GeneC")
)

# Create regulatory features
features <- data.frame(
  chrom = "chr1",
  start = c(500, 800, 1200, 1800, 2200, 2800, 3200),
  end = c(600, 900, 1300, 1900, 2300, 2900, 3300),
  feature_id = paste0("F", 1:7)
)

# Find upstream neighbors (promoter analysis)
upstream <- gintervals.neighbors.upstream(tss, features,
  maxneighbors = 2, maxdist = 1000
)
print(upstream)

# Find downstream neighbors (gene body analysis)
downstream <- gintervals.neighbors.downstream(tss, features,
  maxneighbors = 2, maxdist = 5000
)
print(downstream)

# Find both directions in one call
both <- gintervals.neighbors.directional(tss, features,
  maxneighbors_upstream = 1,
  maxneighbors_downstream = 1,
  maxdist = 1000
)
print(both$upstream)
print(both$downstream)
```

gintervals.normalize *Normalize intervals to fixed or variable sizes*

Description

This function normalizes intervals by computing their centers and then expanding them to fixed or variable sizes, while ensuring they don't cross chromosome boundaries.

Usage

```
gintervals.normalize(intervals = NULL, size = NULL, intervals.set.out = NULL)
```

Arguments

`intervals` intervals set

`size` target size(s) for normalized intervals. Can be either:

- A single positive integer: all intervals normalized to this size
- A numeric vector: each interval normalized to its corresponding size. Vector length must exactly match the number of intervals, OR a single interval can be provided with multiple sizes to create multiple output intervals (one-to-many expansion).

`intervals.set.out` intervals set name where the function result is saved. If NULL, the result is returned to the user.

Value

Normalized intervals set with fixed or variable sizes, or NULL if result is saved to `intervals.set.out`

See Also

[gintervals.force_range](#)

Examples

```
gdb.init_examples()

# Single size (all intervals normalized to 500bp)
intervs <- gintervals(1, c(1000, 5000), c(2000, 6000))
gintervals.normalize(intervs, 500)

# Vector of sizes (each interval gets its own size)
intervs <- gintervals(1, c(1000, 3000, 5000), c(2000, 4000, 6000))
gintervals.normalize(intervs, c(500, 1000, 750))
```

```
# One-to-many: single interval with multiple sizes
interv <- gintervals(1, 1000, 2000)
gintervals.normalize(interv, c(500, 1000, 1500))
```

| | |
|-----------------|--|
| gintervals.path | <i>Returns the path on disk of an interval set</i> |
|-----------------|--|

Description

Returns the path on disk of an interval set.

Usage

```
gintervals.path(intervals.set = NULL)
```

Arguments

`intervals.set` name of an interval set or a vector of interval set names

Details

This function returns the actual file system path where an interval set is stored. The function works with a single interval set name or a vector of names.

Value

A character vector containing the full paths to the interval sets on disk.

See Also

[gintervals.exists](#), [gintervals.ls](#), [gtrack.path](#)

Examples

```
gdb.init_examples()
gintervals.path("annotations")
gintervals.path(c("annotations", "coding"))
```

gintervals.quantiles *Calculates quantiles of a track expression for intervals*

Description

Calculates quantiles of a track expression for intervals.

Usage

```
gintervals.quantiles(  
    expr = NULL,  
    percentiles = 0.5,  
    intervals = NULL,  
    iterator = NULL,  
    band = NULL,  
    intervals.set.out = NULL  
)
```

Arguments

| | |
|-------------------|--|
| expr | track expression for which quantiles are calculated |
| percentiles | an array of percentiles of quantiles in [0, 1] range |
| intervals | set of intervals |
| iterator | track expression iterator. If 'NULL' iterator is determined implicitly based on track expressions. |
| band | track expression band. If 'NULL' no band is used. |
| intervals.set.out | intervals set name where the function result is optionally outputted |

Details

This function calculates quantiles of 'expr' for each interval in 'intervals'.

If 'intervals.set.out' is not 'NULL' the result is saved as an intervals set. Use this parameter if the result size exceeds the limits of the physical memory.

Value

If 'intervals.set.out' is 'NULL' a set of intervals with additional columns representing quantiles for each percentile.

See Also

[gquantiles](#), [gbins.quantiles](#)

Examples

```
gdb.init_examples()
intervs <- gintervals(c(1, 2), 0, 5000)
gintervals.quantiles("dense_track",
  percentiles = c(0.5, 0.3, 0.9), intervs
)
```

| | |
|-------------------|---|
| gintervals.random | <i>Generate random genome intervals</i> |
|-------------------|---|

Description

Generate random genome intervals with a specified number of regions of a specified size. This function samples intervals uniformly across the genome, weighted by chromosome length.

Usage

```
gintervals.random(
  size,
  n,
  dist_from_edge = 3000000,
  chromosomes = NULL,
  filter = NULL
)
```

Arguments

| | |
|----------------|---|
| size | The size of the intervals to generate (in base pairs) |
| n | The number of intervals to generate |
| dist_from_edge | The minimum distance from the edge of the chromosome for a region to start or end (default: 3e6) |
| chromosomes | The chromosomes to sample from (default: all chromosomes). Can be a character vector of chromosome names. |
| filter | A set of intervals to exclude from sampling (default: NULL). Generated intervals will not overlap with these regions. |

Details

The function samples intervals randomly across the genome, with chromosomes weighted by their length. Each interval is guaranteed to:

- Be of the specified size
- Start and end at least `dist_from_edge` bases away from chromosome boundaries
- Fall entirely within a single chromosome

- Not overlap with any intervals in the filter (if provided)

When a filter is provided, the function pre-computes valid genome segments (regions not in the filter) and samples from these segments. Note that this can be slow when the filter contains many intervals.

The function uses R's random number generator, so `set.seed()` can be used for reproducibility.

This function is implemented in C++ for high performance and can generate millions of intervals quickly.

Value

A data.frame with columns chrom, start, and end representing genomic intervals

Examples

```
## Not run:
gdb.init_examples()

# Generate 1000 random intervals of 100bp
intervals <- gintervals.random(100, 1000)
head(intervals)

# Generate intervals only on chr1 and chr2
intervals <- gintervals.random(100, 1000, chromosomes = c("chr1", "chr2"))

# Generate intervals avoiding specific regions
filter_regions <- gintervals(c("chr1", "chr2"), c(1000, 5000), c(2000, 6000))
intervals <- gintervals.random(100, 1000, filter = filter_regions)

# Verify no overlaps with filter
overlaps <- gintervals.intersect(intervals, filter_regions)
nrow(overlaps) # Should be 0

# For reproducibility
set.seed(123)
intervals1 <- gintervals.random(100, 100)
set.seed(123)
intervals2 <- gintervals.random(100, 100)
identical(intervals1, intervals2) # TRUE

## End(Not run)
```

gintervals.rbind

Combines several sets of intervals

Description

Combines several sets of intervals into one set.

Usage

```
gintervals.rbind(..., intervals.set.out = NULL)
```

Arguments

```
...           intervals sets to combine
intervals.set.out
               intervals set name where the function result is optionally outputted
intervals     intervals set
```

Details

This function combines several intervals sets into one set. It works in a similar manner as 'rbind' yet it is faster. Also it supports intervals sets that are stored in files including the big intervals sets.

If 'intervals.set.out' is not 'NULL' the result is saved as an intervals set. If the format of the output intervals is set to be "big" (determined implicitly based on the result size and options), the order of the resulted intervals is altered as they are sorted by chromosome (or chromosomes pair - for 2D).

Value

If 'intervals.set.out' is 'NULL' a data frame combining intervals sets.

See Also

[gintervals](#), [gintervals.2d](#), [gintervals.canonic](#)

Examples

```
gdb.init_examples()

intervs1 <- gextract("sparse_track", gintervals(c(1, 2), 1000, 4000))
intervs2 <- gextract("sparse_track", gintervals(c(2, "X"), 2000, 5000))
gintervals.save("testintervs", interv2)
gintervals.rbind(intervs1, "testintervs")
gintervals.rm("testintervs", force = TRUE)
```

gintervals.rm

Deletes a named intervals set

Description

Deletes a named intervals set.

Usage

```
gintervals.rm(intervals.set = NULL, force = FALSE, db = NULL)
```

Arguments

intervals.set name of an intervals set
 force if 'TRUE', suppresses user confirmation of a named intervals set removal
 db optional database path. When multiple databases are connected, this specifies which database to delete the intervals set from. If NULL (the default), the intervals set is deleted from the working database (GROOT).

Details

This function deletes a named intervals set from the Genomic Database. By default 'gintervals.rm' requires the user to interactively confirm the deletion. Set 'force' to 'TRUE' to suppress the user prompt.

Value

None.

See Also

[gintervals.save](#), [gintervals.exists](#), [gintervals.ls](#), [gintervals](#), [gintervals.2d](#), [gtrack.rm](#)

Examples

```
gdb.init_examples()
intervs <- gintervals(c(1, 2))
gintervals.save("testintervs", intervs)
gintervals.ls()
gintervals.rm("testintervs", force = TRUE)
gintervals.ls()
```

gintervals.save *Creates a named intervals set*

Description

Saves intervals to a named intervals set.

Usage

```
gintervals.save(intervals.set.out = NULL, intervals = NULL)
```

Arguments

intervals.set.out name of the new intervals set
 intervals intervals to save

Details

This function saves 'intervals' as a named intervals set.

Value

None.

See Also

[gintervals.rm](#), [gintervals.load](#), [gintervals.exists](#), [gintervals.ls](#), [gintervals](#), [gintervals.2d](#)

Examples

```
gdb.init_examples()
intervs <- gintervals(c(1, 2))
gintervals.save("testintervs", intervs)
gintervals.ls()
gintervals.rm("testintervs", force = TRUE)
```

| | |
|--------------------|--|
| gintervals.summary | <i>Calculates summary statistics of track expression for intervals</i> |
|--------------------|--|

Description

Calculates summary statistics of track expression for intervals.

Usage

```
gintervals.summary(
  expr = NULL,
  intervals = NULL,
  iterator = NULL,
  band = NULL,
  intervals.set.out = NULL
)
```

Arguments

| | |
|-------------------|---|
| expr | track expression |
| intervals | set of intervals |
| iterator | track expression iterator. If 'NULL' iterator is determined implicitly based on track expression. |
| band | track expression band. If 'NULL' no band is used. |
| intervals.set.out | intervals set name where the function result is optionally outputted |

Details

This function returns summary statistics of a track expression for each interval 'intervals': total number of bins, total number of bins whose value is NaN, min, max, sum, mean and standard deviation of the values.

If 'intervals.set.out' is not 'NULL' the result is saved as an intervals set. Use this parameter if the result size exceeds the limits of the physical memory.

Value

If 'intervals.set.out' is 'NULL' a set of intervals with additional columns representing summary statistics for each percentile and interval.

See Also

[gsummary](#), [gbins.summary](#)

Examples

```
gdb.init_examples()
intervals <- gintervals(c(1, 2), 0, 5000)
gintervals.summary("dense_track", intervals)
```

`gintervals.union`

Calculates a union of two sets of intervals

Description

Calculates a union of two sets of intervals.

Usage

```
gintervals.union(
  intervals1 = NULL,
  intervals2 = NULL,
  intervals.set.out = NULL
)
```

Arguments

`intervals1`, `intervals2`
 set of one-dimensional intervals

`intervals.set.out`
 intervals set name where the function result is optionally outputted

Details

This function returns intervals that represent a genomic space covered by either 'intervals1' or 'intervals2'.

If 'intervals.set.out' is not 'NULL' the result is saved as an intervals set. Use this parameter if the result size exceeds the limits of the physical memory.

Value

If 'intervals.set.out' is 'NULL' a data frame representing the union of intervals.

See Also

[gintervals.intersect](#), [gintervals.diff](#), [gintervals](#), [gintervals.2d](#)

Examples

```
gdb.init_examples()

intervs1 <- gscreen("dense_track > 0.15 & dense_track < 0.18")
intervs2 <- gscreen("dense_track >= 0.18 & dense_track < 0.2")

## 'intervs3' and 'intervs4' are identical
intervs3 <- gintervals.union(intervs1, intervs2)
intervs4 <- gscreen("dense_track > 0.15 & dense_track < 0.2")
```

| | |
|-------------------|--------------------------------------|
| gintervals.update | <i>Updates a named intervals set</i> |
|-------------------|--------------------------------------|

Description

Updates a named intervals set.

Usage

```
gintervals.update(
  intervals.set = NULL,
  intervals = "",
  chrom = NULL,
  chrom1 = NULL,
  chrom2 = NULL
)
```

Arguments

| | |
|----------------------------|--|
| <code>intervals.set</code> | name of an intervals set |
| <code>intervals</code> | intervals or 'NULL' |
| <code>chrom</code> | chromosome for 1D intervals set |
| <code>chrom1</code> | first chromosome for 2D intervals set |
| <code>chrom2</code> | second chromosome for 2D intervals set |

Details

This function replaces all intervals of given chromosome (or chromosome pair) within 'intervals.set' with 'intervals'. Chromosome is specified by 'chrom' for 1D intervals set or 'chrom1', 'chrom2' for 2D intervals set.

If 'intervals' is 'NULL' all intervals of given chromosome are removed from 'intervals.set'.

Value

None.

See Also

[gintervals.save](#), [gintervals.load](#), [gintervals.exists](#), [gintervals.ls](#)

Examples

```
gdb.init_examples()
intervs <- gscreen(
  "sparse_track > 0.2",
  gintervals(c(1, 2), 0, 10000)
)
gintervals.save("testintervs", intervs)
gintervals.load("testintervs")
gintervals.update("testintervs", intervs[intervs$chrom == "chr2", ][1:5, ], chrom = 2)
gintervals.load("testintervs")
gintervals.update("testintervs", NULL, chrom = 2)
gintervals.load("testintervs")
gintervals.rm("testintervs", force = TRUE)
```

giterator.cartesian_grid

Creates a cartesian-grid iterator

Description

Creates a cartesian grid two-dimensional iterator that can be used by any function that accepts an iterator argument.

Usage

```
giterator.cartesian_grid(
  intervals1 = NULL,
  expansion1 = NULL,
  intervals2 = NULL,
  expansion2 = NULL,
  min.band.idx = NULL,
  max.band.idx = NULL
)
```

Arguments

| | |
|---|--|
| <code>intervals1</code> | one-dimensional intervals |
| <code>expansion1</code> | an array of integers that define expansion around <code>intervals1</code> centers |
| <code>intervals2</code> | one-dimensional intervals. If 'NULL' then 'intervals2' is considered to be equal to 'intervals1' |
| <code>expansion2</code> | an array of integers that define expansion around <code>intervals2</code> centers. If 'NULL' then 'expansion2' is considered to be equal to 'expansion1' |
| <code>min.band.idx, max.band.idx</code> | integers that limit iterator intervals to band |

Details

This function creates and returns a cartesian grid two-dimensional iterator that can be used by any function that accepts an iterator argument.

Assume 'centers1' and 'centers2' to be the central points of each interval from 'intervals1' and 'intervals2', and 'C1', 'C2' to be two points from 'centers1', 'centers2' accordingly. Assume also that the values in 'expansion1' and 'expansion2' are unique and sorted.

'giterator.cartesian_grid' creates a set of all possible unique and non-overlapping two-dimensional intervals of form: '(chrom1, start1, end1, chrom2, start2, end2)'. Each '(chrom1, start1, end1)' is created by taking a point 'C1' - '(chrom1, coord1)' and converting it to 'start1' and 'end1' such that 'start1 == coord1+E1[i]', 'end1 == coord1+E1[i+1]', where 'E1[i]' is one of the sorted 'expansion1' values. Overlaps between rectangles or expansion beyond the limits of chromosome are avoided.

'min.band.idx' and 'max.band.idx' parameters control whether a pair of 'C1' and 'C2' is skipped or not. If both of these parameters are not 'NULL' AND if both 'C1' and 'C2' share the same chromosome AND the delta of indices of 'C1' and 'C2' ('C1 index - C2 index') lays within '[min.band.idx, max.band.idx]' range - only then the pair will be used to create the intervals. Otherwise 'C1-C2' pair is filtered out. Note: if 'min.band.idx' and 'max.band.idx' are not 'NULL', i.e. band indices filtering is applied, then 'intervals2' parameter must be set to 'NULL'.

Value

A list containing the definition of cartesian iterator.

See Also[giterator.intervals](#)**Examples**

```
gdb.init_examples()

intervs1 <- gintervals(
  c(1, 1, 2), c(100, 300, 200),
  c(300, 500, 300)
)
intervs2 <- gintervals(
  c(1, 2, 2), c(400, 1000, 3000),
  c(800, 2000, 4000)
)
itr <- giterator.cartesian_grid(
  intervs1, c(-20, 100), intervs2,
  c(-40, -10, 50)
)
giterator.intervals(iterator = itr)

itr <- giterator.cartesian_grid(intervs1, c(-20, 50, 100))
giterator.intervals(iterator = itr)

itr <- giterator.cartesian_grid(intervs1, c(-20, 50, 100),
  min.band.idx = -1,
  max.band.idx = 0
)
giterator.intervals(iterator = itr)
```

`giterator.intervals` *Returns iterator intervals*

Description

Returns iterator intervals given track expression, scope, iterator and band.

Usage

```
giterator.intervals(
  expr = NULL,
  intervals = .misha$ALLGENOME,
  iterator = NULL,
  band = NULL,
  intervals.set.out = NULL,
  interval_relative = FALSE,
  partial_bins = c("clip", "exact", "drop")
)
```

Arguments

| | |
|--------------------------------|---|
| <code>expr</code> | track expression |
| <code>intervals</code> | genomic scope |
| <code>iterator</code> | track expression iterator. If 'NULL' iterator is determined implicitly based on track expression. |
| <code>band</code> | track expression band. If 'NULL' no band is used. |
| <code>intervals.set.out</code> | intervals set name where the function result is optionally outputted |
| <code>interval_relative</code> | if TRUE, and iterator is numeric, bins start at each interval's start position instead of chromosome position 0. Returns <code>intervalID</code> column. Default: FALSE. |
| <code>partial_bins</code> | how to handle partial bins at interval boundaries when <code>interval_relative</code> is TRUE. One of "clip" (default, truncate last bin to interval boundary), "exact" or "drop" (only output full-size bins). |

Details

This function returns a set of intervals used by the iterator `intervals` for the given track expression, genomic scope, iterator and band. Some functions accept an iterator without accepting a track expression (like `gtrack.create_pwm_energy`). These functions generate the values for each iterator interval by themselves. Use set `expr` to 'NULL' to simulate the work of these functions.

If `intervals.set.out` is not 'NULL' the result is saved as an intervals set. Use this parameter if the result size exceeds the limits of the physical memory.

When `interval_relative` is TRUE, bins are aligned to each input interval's start position rather than chromosome position 0. This mode requires a numeric iterator (binsize) and returns an additional `intervalID` column indicating which input interval spawned each bin.

Value

If `intervals.set.out` is 'NULL' a data frame representing iterator intervals. When `interval_relative` is TRUE, includes an `intervalID` column.

See Also

[giterator.cartesian_grid](#)

Examples

```
gdb.init_examples()

## iterator is set implicitly to bin size of 'dense' track
giterator.intervals("dense_track", gintervals(1, 0, 200))

## iterator = 30
giterator.intervals("dense_track", gintervals(1, 0, 200), 30)
```

```

## iterator is an intervals set named 'annotations'
giterator.intervals("dense_track", .misha$ALLGENOME, "annotations")

## iterator is set implicitly to intervals of 'array_track' track
giterator.intervals("array_track", gintervals(1, 0, 200))

## iterator is a rectangle 100000 by 50000
giterator.intervals(
  "rects_track",
  gintervals.2d(chroms1 = 1, chroms2 = "chrX"),
  c(100000, 50000)
)

## interval_relative mode: bins aligned to each interval's start
intervs <- gintervals(1, c(100, 500), c(300, 700))
giterator.intervals(NULL, intervs, iterator = 50, interval_relative = TRUE)

```

glookup

Returns values from a lookup table based on track expression

Description

Evaluates track expression and translates the values into bin indices that are used in turn to retrieve and return values from a lookup table.

Usage

```

glookup(
  lookup_table = NULL,
  ...,
  intervals = NULL,
  include.lowest = FALSE,
  force.binning = TRUE,
  iterator = NULL,
  band = NULL,
  intervals.set.out = NULL
)

```

Arguments

`lookup_table` a multi-dimensional array containing the values that are returned by the function

`...` pairs of 'expr', 'breaks' where 'expr' is a track expression and the breaks determine the bin

`intervals` genomic scope for which the function is applied

`include.lowest` if 'TRUE', the lowest value of the range determined by breaks is included

| | |
|--------------------------------|---|
| <code>force.binning</code> | if 'TRUE', the values smaller than the minimal break will be translated to index 1, and the values that exceed the maximal break will be translated to index N-1 where N is the number of breaks. If 'FALSE' the out-of-range values will produce NaN values. |
| <code>iterator</code> | track expression iterator. If 'NULL' iterator is determined implicitly based on track expressions. |
| <code>band</code> | track expression band. If 'NULL' no band is used. |
| <code>intervals.set.out</code> | intervals set name where the function result is optionally outputted |

Details

This function evaluates the track expression for all iterator intervals and translates this value into an index based on the breaks. This index is then used to address the lookup table and return the according value. More than one 'expr'-'breaks' pair can be used. In that case 'lookup_table' is addressed in a multidimensional manner, i.e. 'lookup_table[i1, i2, ...]'.

The range of bins is determined by 'breaks' argument. For example: 'breaks = c(x1, x2, x3, x4)' represents three different intervals (bins): (x1, x2], (x2, x3], (x3, x4].

If 'include.lowest' is 'TRUE' then the lowest value is included in the first interval, i.e. in [x1, x2].

'force.binning' parameter controls what should be done when the value of 'expr' exceeds the range determined by 'breaks'. If 'force.binning' is 'TRUE' then values smaller than the minimal break will be translated to index 1, and the values exceeding the maximal break will be translated to index 'M-1' where 'M' is the number of breaks. If 'force.binning' is 'FALSE' the out-of-range values will produce 'NaN' values.

Regardless of 'force.binning' value if the value of 'expr' is 'NaN' then result is 'NaN' too.

The order inside the result might not be the same as the order of intervals. Use 'intervalID' column to refer to the index of the original interval from the supplied 'intervals'.

If 'intervals.set.out' is not 'NULL' the result (without 'columnID' column) is saved as an intervals set. Use this parameter if the result size exceeds the limits of the physical memory.

Value

If 'intervals.set.out' is 'NULL' a set of intervals with additional 'value' and 'columnID' columns.

See Also

[gtrack.lookup](#), [gextract](#), [gpartition](#), [gdist](#)

Examples

```
gdb.init_examples()

## one-dimensional lookup table
breaks1 <- seq(0.1, 0.2, length.out = 6)
glookup(1:5, "dense_track", breaks1, gintervals(1, 0, 200))
```

```
## two-dimensional lookup table
t <- array(1:15, dim = c(5, 3))
breaks2 <- seq(0.31, 0.37, length.out = 4)
glookup(
  t, "dense_track", breaks1, "2 * dense_track", breaks2,
  gintervals(1, 0, 200)
)
```

gpartition

Partitions the values of track expression

Description

Converts the values of track expression to intervals that match corresponding bin.

Usage

```
gpartition(
  expr = NULL,
  breaks = NULL,
  intervals = NULL,
  include.lowest = FALSE,
  iterator = NULL,
  band = NULL,
  intervals.set.out = NULL
)
```

Arguments

| | |
|-------------------|---|
| expr | track expression |
| breaks | breaks that determine the bin |
| intervals | genomic scope for which the function is applied |
| include.lowest | if 'TRUE', the lowest value of the range determined by breaks is included |
| iterator | track expression iterator. If 'NULL' iterator is determined implicitly based on track expression. |
| band | track expression band. If 'NULL' no band is used. |
| intervals.set.out | intervals set name where the function result is optionally outputted |

Details

This function converts first the values of track expression into 1-based bin's index according 'breaks' argument. It returns then the intervals with the corresponding bin's index.

The range of bins is determined by 'breaks' argument. For example: 'breaks=c(x1, x2, x3, x4)' represents three different intervals (bins): (x1, x2], (x2, x3], (x3, x4].

If 'include.lowest' is 'TRUE' the the lowest value will be included in the first interval, i.e. in [x1, x2].

If 'intervals.set.out' is not 'NULL' the result is saved as an intervals set. Use this parameter if the result size exceeds the limits of the physical memory.

Value

If 'intervals.set.out' is 'NULL' a set of intervals with an additional column that indicates the corresponding bin index.

See Also

[gscreen](#), [gextract](#), [glookup](#), [gdist](#)

Examples

```
gdb.init_examples()
breaks <- seq(0, 0.2, by = 0.05)
gpartition("dense_track", breaks, gintervals(1, 0, 5000))
```

gquantiles

Calculates quantiles of a track expression

Description

Calculates the quantiles of a track expression for the given percentiles.

Usage

```
gquantiles(
  expr = NULL,
  percentiles = 0.5,
  intervals = get("ALLGENOME", envir = .misha),
  iterator = NULL,
  band = NULL
)
```

Arguments

| | |
|-------------|---|
| expr | track expression |
| percentiles | an array of percentiles of quantiles in [0, 1] range |
| intervals | genomic scope for which the function is applied |
| iterator | track expression iterator. If 'NULL' iterator is determined implicitly based on track expression. |
| band | track expression band. If 'NULL' no band is used. |

Details

This function calculates the quantiles for the given percentiles.

If data size exceeds the limit (see: 'getOption(gmax.data.size)'), the data is randomly sampled to fit the limit. A warning message is generated. The seed of the pseudo-random generator can be controlled through 'grnd.seed' option.

Note: this function is capable to run in multitasking mode. Sampling may vary according to the extent of multitasking. Since multitasking depends on the number of available CPU cores, running the function on two different machines might give different results. Please switch off multitasking if you want to achieve identical results on any machine. For more information regarding multitasking please refer "User Manual".

Value

An array that represent quantiles.

See Also

[gbins.quantiles](#), [gintervals.quantiles](#), [gdist](#)

Examples

```
gdb.init_examples()
gquantiles("dense_track", c(0.1, 0.6, 0.8), gintervals(c(1, 2)))
```

grevcomp

Get reverse complement of DNA sequence

Description

Takes a DNA sequence string and returns its reverse complement.

Usage

```
grevcomp(seq)
```

Arguments

seq A character vector containing DNA sequences (using A,C,G,T). Ignores other characters and NA values.

Value

A character vector of the same length as the input, containing the reverse complement sequences

Examples

```
grevcomp("ACTG") # Returns "CAGT"  
grevcomp(c("ACTG", "GGCC")) # Returns c("CAGT", "GGCC")  
grevcomp(c("ACTG", NA, "GGCC")) # Returns c("CAGT", NA, "GGCC")
```

| | |
|---------|--|
| gsample | <i>Returns samples from the values of track expression</i> |
|---------|--|

Description

Returns a sample of the specified size from the values of track expression.

Usage

```
gsample(expr = NULL, n = NULL, intervals = NULL, iterator = NULL, band = NULL)
```

Arguments

| | |
|-----------|---|
| expr | track expression |
| n | a number of items to choose |
| intervals | genomic scope for which the function is applied |
| iterator | track expression iterator. If 'NULL' iterator is determined implicitly based on track expression. |
| band | track expression band. If 'NULL' no band is used. |

Details

This function returns a sample of the specified size from the values of track expression. If 'n' is less than the total number of values, the data is randomly sampled. The seed of the pseudo-random generator can be controlled through 'grnd.seed' option.

If 'n' is higher than the total number of values, all values are returned (yet reshuffled).

Value

An array that represent quantiles.

See Also

[gextract](#)

Examples

```
gdb.init_examples()  
gsample("sparse_track", 10)
```

gscreen

Finds intervals that match track expression

Description

Finds all intervals where track expression is 'TRUE'.

Usage

```
gscreen(  
  expr = NULL,  
  intervals = NULL,  
  iterator = NULL,  
  band = NULL,  
  intervals.set.out = NULL  
)
```

Arguments

| | |
|-------------------|---|
| expr | logical track expression |
| intervals | genomic scope for which the function is applied |
| iterator | track expression iterator. If 'NULL' iterator is determined implicitly based on track expression. |
| band | track expression band. If 'NULL' no band is used. |
| intervals.set.out | intervals set name where the function result is optionally outputted |

Details

This function finds all intervals where track expression's value is 'TRUE'.

If 'intervals.set.out' is not 'NULL' the result is saved as an intervals set. Use this parameter if the result size exceeds the limits of the physical memory.

Value

If 'intervals.set.out' is 'NULL' a set of intervals that match track expression.

See Also

[gsegment](#), [gextract](#)

Examples

```
gdb.init_examples()
gscreen("dense_track > 0.2 & sparse_track < 0.4",
        iterator = "dense_track"
)
```

gsegment

Divides track expression into segments

Description

Divides the values of track expression into segments by using Wilcoxon test.

Usage

```
gsegment(
  expr = NULL,
  minsegment = NULL,
  maxpval = 0.05,
  onetailed = TRUE,
  intervals = NULL,
  iterator = NULL,
  intervals.set.out = NULL
)
```

Arguments

| | |
|-------------------|---|
| expr | track expression |
| minsegment | minimal segment size |
| maxpval | maximal P-value that separates two adjacent segments |
| onetailed | if 'TRUE', Wilcoxon test is performed one tailed, otherwise two tailed |
| intervals | genomic scope for which the function is applied |
| iterator | track expression iterator of "fixed bin" type. If 'NULL' iterator is determined implicitly based on track expression. |
| intervals.set.out | intervals set name where the function result is optionally outputted |

Details

This function divides the values of track expression into segments, where each segment size is at least of 'minsegment' size and the P-value of comparing the segment with the first 'minsegment' values from the next segment is at most 'maxpval'. Comparison is done using Wilcoxon (also known as Mann-Whitney) test.

If 'intervals.set.out' is not 'NULL' the result is saved as an intervals set. Use this parameter if the result size exceeds the limits of the physical memory.

Value

If 'intervals.set.out' is 'NULL' a set of intervals where each interval represents a segment.

See Also

[gscreen](#), [gwilcox](#)

Examples

```
gdb.init_examples()
gsegment("dense_track", 5000, 0.0001)
```

gseq.comp

Complement DNA sequence

Description

Takes a DNA sequence string and returns its complement (without reversing).

Usage

```
gseq.comp(seq)
```

Arguments

| | |
|-----|--|
| seq | A character vector containing DNA sequences (using A,C,G,T). Preserves case and handles NA values. |
|-----|--|

Value

A character vector of the same length as the input, containing the complemented sequences

See Also

[gseq.revcomp](#), [gseq.rev](#)

Examples

```
gseq.comp("ACTG") # Returns "TGAC"
gseq.comp(c("ACTG", "GGCC")) # Returns c("TGAC", "CCGG")
gseq.comp(c("ACTG", NA, "GGCC")) # Returns c("TGAC", NA, "CCGG")
```

| | |
|--------------|-----------------------|
| gseq.extract | Returns DNA sequences |
|--------------|-----------------------|

Description

Returns DNA sequences for given intervals

Usage

```
gseq.extract(intervals = NULL)
```

Arguments

intervals intervals for which DNA sequence is returned

Details

This function returns an array of sequence strings for each interval from 'intervals'. If intervals contain an additional 'strand' column and its value is '-1', the reverse-complementary sequence is returned.

Value

An array of character strings representing DNA sequence.

See Also

[gextract](#)

Examples

```
gdb.init_examples()
intervs <- gintervals(c(1, 2), 10000, 10020)
gseq.extract(intervs)
```

gseq.kmer

*Score DNA sequences with a k-mer over a region of interest***Description**

Counts exact matches of a k-mer in DNA sequences over a specified region of interest (ROI). The ROI is defined by `start_pos` and `end_pos` (1-based, inclusive), with optional extension controlled by `extend`.

Usage

```
gseq.kmer(
  seqs,
  kmer,
  mode = c("count", "frac"),
  strand = 0L,
  start_pos = NULL,
  end_pos = NULL,
  extend = FALSE,
  skip_gaps = TRUE,
  gap_chars = c("-", ".")
)
```

Arguments

| | |
|------------------------|---|
| <code>seqs</code> | character vector of DNA sequences (A/C/G/T/N; case-insensitive) |
| <code>kmer</code> | single character string containing the k-mer to search for (A/C/G/T only) |
| <code>mode</code> | character; one of "count" or "frac" |
| <code>strand</code> | integer; 1=forward, -1=reverse, 0=both strands (default: 0) |
| <code>start_pos</code> | integer or NULL; 1-based inclusive start of ROI (default: 1) |
| <code>end_pos</code> | integer or NULL; 1-based inclusive end of ROI (default: sequence length) |
| <code>extend</code> | logical or integer; extension of allowed window starts (default: FALSE) |
| <code>skip_gaps</code> | logical; if TRUE, treat gap characters as holes and skip them while scanning. Windows are k consecutive non-gap bases (default: TRUE) |
| <code>gap_chars</code> | character vector; which characters count as gaps (default: c("-", ".")) |

Details

This function counts k-mer occurrences in DNA sequences directly without requiring a genomics database. For detailed documentation on k-mer counting parameters, see [gvtrack.create](#) (functions "kmer.count" and "kmer.frac").

The ROI (region of interest) is defined by `start_pos` and `end_pos`. The `extend` parameter controls whether k-mer matches can extend beyond the ROI boundaries. For palindromic k-mers, use `strand=1` or `-1` to avoid double counting.

When `skip_gaps=TRUE`, characters specified in `gap_chars` are treated as gaps. Windows are defined as `k` consecutive non-gap bases. The `frac` denominator counts the number of possible logical starts (non-gap windows) in the region. `start_pos` and `end_pos` are interpreted as physical coordinates on the full sequence.

Value

Numeric vector with counts (for "count" mode) or fractions (for "frac" mode). Returns 0 when sequence is too short or ROI is invalid.

See Also

[gvtrack.create](#) for detailed k-mer parameter documentation

Examples

```
## Not run:
# Example sequences
seqs <- c("CGCGCGCGCG", "ATATATATAT", "ACGTACGTACGT")

# Count CG dinucleotides on both strands
gseq.kmer(seqs, "CG", mode = "count", strand = 0)

# Count on forward strand only
gseq.kmer(seqs, "CG", mode = "count", strand = 1)

# Get CG fraction
gseq.kmer(seqs, "CG", mode = "frac", strand = 0)

# Count in a specific region
gseq.kmer(seqs, "CG", mode = "count", start_pos = 2, end_pos = 8)

# Allow k-mer to extend beyond ROI boundaries
gseq.kmer(seqs, "CG", mode = "count", start_pos = 2, end_pos = 8, extend = TRUE)

# Calculate GC content by summing G and C fractions
g_frac <- gseq.kmer(seqs, "G", mode = "frac", strand = 1)
c_frac <- gseq.kmer(seqs, "C", mode = "frac", strand = 1)
gc_content <- g_frac + c_frac
gc_content

# Compare AT counts on different strands
at_forward <- gseq.kmer(seqs, "AT", mode = "count", strand = 1)
at_reverse <- gseq.kmer(seqs, "AT", mode = "count", strand = -1)
at_both <- gseq.kmer(seqs, "AT", mode = "count", strand = 0)
data.frame(forward = at_forward, reverse = at_reverse, both = at_both)

## End(Not run)
```

| | |
|----------------|--|
| gseq.kmer.dist | <i>Compute k-mer distribution in genomic intervals</i> |
|----------------|--|

Description

Counts the occurrence of all k-mers (of size k) within the specified genomic intervals, optionally excluding masked regions.

Usage

```
gseq.kmer.dist(intervals, k = 6L, mask = NULL)
```

Arguments

| | |
|-----------|---|
| intervals | Genomic intervals to analyze |
| k | Integer k-mer size (1-10). Default is 6. |
| mask | Optional intervals to exclude from counting. Positions within the mask will not contribute to k-mer counts. |

Value

A data frame with columns:

kmer Character string representing the k-mer sequence

count Number of occurrences of this k-mer

Only k-mers with count > 0 are included. K-mers containing N bases are not counted.

See Also

[gseq.extract](#), [gseq.kmer](#)

Examples

```
gdb.init_examples()

# Count all 6-mers in first 10kb of chr1
intervals <- data.frame(chrom = "chr1", start = 0, end = 10000)
kmer_dist <- gseq.kmer.dist(intervals, k = 6)
head(kmer_dist)

# Count dinucleotides
dinucs <- gseq.kmer.dist(intervals, k = 2)
dinucs

# Count with mask
mask <- data.frame(chrom = "chr1", start = 5000, end = 6000)
kmer_dist_masked <- gseq.kmer.dist(intervals, k = 6, mask = mask)
```

gseq.pwm

*Score DNA sequences with a PWM over a region of interest***Description**

Scores full DNA sequences using a Position Weight Matrix (PWM) over a specified region of interest (ROI). The ROI is defined by `start_pos` and `end_pos` (1-based, inclusive), with optional extension controlled by `extend`. All reported positions are on the full input sequence.

Usage

```
gseq.pwm(
  seqs,
  pssm,
  mode = c("lse", "max", "pos", "count"),
  bidirect = TRUE,
  strand = 0L,
  score.thresh = 0,
  start_pos = NULL,
  end_pos = NULL,
  extend = FALSE,
  spat.factor = NULL,
  spat.bin = 1L,
  spat.min = NULL,
  spat.max = NULL,
  return_strand = FALSE,
  skip_gaps = TRUE,
  gap_chars = c("-", "."),
  neutral_chars = c("N", "n", "*"),
  neutral_chars_policy = c("average", "log_quarter", "na"),
  prior = 0.01
)
```

Arguments

| | |
|---------------------------|---|
| <code>seqs</code> | character vector of DNA sequences (A/C/G/T/N; case-insensitive) |
| <code>pssm</code> | numeric matrix or data frame with columns named A, C, G, T (additional columns are allowed and will be ignored) |
| <code>mode</code> | character; one of "lse", "max", "pos", or "count" |
| <code>bidirect</code> | logical; if TRUE, scans both strands (default: TRUE) |
| <code>strand</code> | integer; 1=forward, -1=reverse, 0=both strands (default: 0) |
| <code>score.thresh</code> | numeric; score threshold for mode="count" (default: 0) |
| <code>start_pos</code> | integer or NULL; 1-based inclusive start of ROI (default: 1) |
| <code>end_pos</code> | integer or NULL; 1-based inclusive end of ROI (default: sequence length) |
| <code>extend</code> | logical or integer; extension of allowed window starts (default: FALSE) |

| | |
|-----------------------------------|--|
| <code>spat.factor</code> | numeric vector; spatial weighting factors (optional) |
| <code>spat.bin</code> | integer; bin size for spatial weighting |
| <code>spat.min</code> | numeric; start of scanning window |
| <code>spat.max</code> | numeric; end of scanning window |
| <code>return_strand</code> | logical; if TRUE and mode="pos", returns data.frame with pos and strand columns |
| <code>skip_gaps</code> | logical; if TRUE, treat gap characters as holes and skip them while scanning. Windows are w consecutive non-gap bases (default: TRUE) |
| <code>gap_chars</code> | character vector; which characters count as gaps (default: c("-", ".")) |
| <code>neutral_chars</code> | character vector; bases treated as unknown and scored with the average log probability per position (default: c("N", "n", "*")) |
| <code>neutral_chars_policy</code> | character string; how to treat neutral characters. One of "average" (default; use the column's mean log-probability), "log_quarter" (always use log(1/4)), or "na" (return NA when a neutral character is encountered in the scanning window). |
| <code>prior</code> | numeric; pseudocount added to frequencies (default: 0.01). Set to 0 for no pseudocounts. |

Details

This function scores DNA sequences directly without requiring a genomics database. For detailed documentation on PWM scoring modes, parameters, and spatial weighting, see [gytrack.create](#) (functions "pwm", "pwm.max", "pwm.max.pos", "pwm.count").

The ROI (region of interest) is defined by `start_pos` and `end_pos`. The `extend` parameter controls whether motif matches can extend beyond the ROI boundaries.

When `skip_gaps=TRUE`, characters specified in `gap_chars` are treated as gaps. Windows are defined as `w` consecutive non-gap bases. All positions (`pos`) are reported as 1-based indices on the original full sequence (including gaps). `start_pos` and `end_pos` are interpreted as physical coordinates on the full sequence.

Neutral characters (`neutral_chars`, default c("N", "n", "*")) are treated as unknown bases in both orientations. Each neutral contributes the mean log-probability of the corresponding PSSM column, yielding identical penalties on forward and reverse strands without hard-coded background scores. In mode = "max" the reported value is the single best strand score after applying any spatial weights; forward and reverse contributions are not aggregated. This matches the default behavior of the PWM virtual tracks (`pwm.max`, `pwm.max.pos`, etc.).

Value

Numeric vector (for "lse"/"max"/"count" modes), integer vector (for "pos" mode), or data.frame with `pos` and `strand` columns (for "pos" mode with `return_strand=TRUE`). Returns NA when no valid windows exist.

See Also

[gytrack.create](#) for detailed PWM parameter documentation

Examples

```

## Not run:
# Create a PSSM (position-specific scoring matrix) with frequency values
pssm <- matrix(
  c(
    0.7, 0.1, 0.1, 0.1, # Position 1: mostly A
    0.1, 0.7, 0.1, 0.1, # Position 2: mostly C
    0.1, 0.1, 0.7, 0.1, # Position 3: mostly G
    0.1, 0.1, 0.1, 0.7 # Position 4: mostly T
  ),
  ncol = 4, byrow = TRUE
)
colnames(pssm) <- c("A", "C", "G", "T")

# Example sequences
seqs <- c("ACGTACGTACGT", "GGGGACGTCCCC", "TTTTTTTTTTTT")

# Score sequences using log-sum-exp (default mode)
gseq.pwm(seqs, pssm, mode = "lse")

# Get maximum score
gseq.pwm(seqs, pssm, mode = "max")

# Find position of best match
gseq.pwm(seqs, pssm, mode = "pos")

# Find position with strand information
gseq.pwm(seqs, pssm, mode = "pos", bidirect = TRUE, return_strand = TRUE)

# Count matches above threshold
gseq.pwm(seqs, pssm, mode = "count", score.thresh = 0.5)

# Score only a region of interest
gseq.pwm(seqs, pssm, mode = "max", start_pos = 3, end_pos = 10)

# Allow matches to extend beyond ROI boundaries
gseq.pwm(seqs, pssm, mode = "count", start_pos = 5, end_pos = 8, extend = TRUE)

# Spatial weighting example: higher weight in the center
spatial_weights <- c(0.5, 1.0, 2.0, 1.0, 0.5)
gseq.pwm(seqs, pssm,
  mode = "lse",
  spat.factor = spatial_weights,
  spat.bin = 2
)

## End(Not run)

```

Description

Parses a HOMER .motif format file and returns a named list of position probability matrices (PPM). Each matrix has rows corresponding to motif positions and columns A, C, G, T. The returned matrices are directly usable with [gseq.pwm](#).

Usage

```
gseq.read_homer(file)
```

Arguments

file character(1) path to a HOMER motif file (.motif).

Value

A named list of numeric matrices. Each matrix has columns A, C, G, T and one row per motif position. List names are derived from the consensus sequence. Each matrix carries the following attributes:

name Motif name / description from the header

consensus Consensus sequence from the header

log_odds_threshold Detection threshold (numeric)

log_p_value Log p-value (numeric)

w Motif width (integer)

source "homer"

See Also

Other motif functions: [gseq.read_jaspar\(\)](#), [gseq.read_meme\(\)](#)

Examples

```
## Not run:
motifs <- gseq.read_homer("known_motifs.motif")
names(motifs)
m <- motifs[[1]]
head(m)
attr(m, "consensus")

## End(Not run)
```

| | |
|------------------|--|
| gseq.read_jaspar | <i>Read motifs from a JASPAR PFM format file</i> |
|------------------|--|

Description

Parses a JASPAR Position Frequency Matrix (PFM) file and returns a named list of position probability matrices (PPM). Supports both the standard JASPAR header format (>ID NAME followed by labeled rows) and the simple 4-row PFM format. Counts are converted to probabilities by dividing each column by its column sum.

Usage

```
gseq.read_jaspar(file)
```

Arguments

file character(1) path to a JASPAR format file (.jaspar, .pfm, .txt).

Value

A named list of numeric matrices. Each matrix has columns A, C, G, T and one row per motif position. List names are motif identifiers. Each matrix carries the following attributes:

name Motif name from the header line

w Motif width (integer)

nsites Total counts per position (numeric; NA for simple-format files)

format Sub-format detected: "jaspar" or "simple"

See Also

Other motif functions: [gseq.read_homer\(\)](#), [gseq.read_meme\(\)](#)

Examples

```
## Not run:
motifs <- gseq.read_jaspar("JASPAR2024_CORE.jaspar")
names(motifs)
m <- motifs[[1]]
head(m)

## End(Not run)
```

| | |
|----------------|--|
| gseq.read_meme | <i>Read motifs from a MEME minimal motif format file</i> |
|----------------|--|

Description

Parses a MEME minimal motif format file and returns a named list of position probability matrices (PPM). Each matrix has rows corresponding to motif positions and columns A, C, G, T. The returned matrices are directly usable with [gseq.pwm](#).

Usage

```
gseq.read_meme(file)
```

Arguments

file character(1) path to a MEME format file (.meme, .txt).

Value

A named list of numeric matrices. Each matrix has columns A, C, G, T and one row per motif position. List names are motif identifiers. Each matrix carries the following attributes:

name Motif name / alternate ID (second token on the MOTIF line)

length Alphabet length (integer, typically 4)

w Motif width (integer, number of positions)

nsites Number of sites used to build the matrix (numeric; NA if absent)

E E-value (numeric; NA if absent)

url URL string if present, otherwise NA

strand Strand specification from the file header (e.g. "+ -")

background Named numeric vector of background frequencies (c(A=..., C=..., G=..., T=...)), or NULL if absent

See Also

Other motif functions: [gseq.read_homer\(\)](#), [gseq.read_jaspar\(\)](#)

Examples

```
## Not run:
motifs <- gseq.read_meme("JASPAR2024_CORE_vertbrates.meme")
names(motifs)
m <- motifs[[1]]
head(m)
attr(m, "name")
attr(m, "nsites")

## End(Not run)
```

| | |
|----------|-----------------------------|
| gseq.rev | <i>Reverse DNA sequence</i> |
|----------|-----------------------------|

Description

Takes a DNA sequence string and returns its reverse (without complementing).

Usage

```
gseq.rev(seq)
```

Arguments

| | |
|-----|--|
| seq | A character vector containing DNA sequences. Preserves case and handles NA values. |
|-----|--|

Value

A character vector of the same length as the input, containing the reversed sequences

See Also

[gseq.revcomp](#), [gseq.comp](#)

Examples

```
gseq.rev("ACTG") # Returns "GTCA"  
gseq.rev(c("ACTG", "GGCC")) # Returns c("GTCA", "CCGG")  
gseq.rev(c("ACTG", NA, "GGCC")) # Returns c("GTCA", NA, "CCGG")
```

| | |
|--------------|---|
| gseq.revcomp | <i>Get reverse complement of DNA sequence</i> |
|--------------|---|

Description

Alias for [grevcomp](#). Takes a DNA sequence string and returns its reverse complement.

Usage

```
gseq.revcomp(seq)
```

Arguments

| | |
|-----|--|
| seq | A character vector containing DNA sequences (using A,C,G,T). Ignores other characters and NA values. |
|-----|--|

Value

A character vector of the same length as the input, containing the reverse complement sequences

See Also

[grevcomp](#), [gseq.rev](#), [gseq.comp](#)

Examples

```
gseq.revcomp("ACTG") # Returns "CAGT"  
gseq.revcomp(c("ACTG", "GGCC")) # Returns c("CAGT", "GGCC")
```

gsummary

Calculates summary statistics of track expression

Description

Calculates summary statistics of track expression.

Usage

```
gsummary(expr = NULL, intervals = NULL, iterator = NULL, band = NULL)
```

Arguments

| | |
|-----------|---|
| expr | track expression |
| intervals | genomic scope for which the function is applied |
| iterator | track expression iterator. If 'NULL' iterator is determined implicitly based on track expression. |
| band | track expression band. If 'NULL' no band is used. |

Details

This function returns summary statistics of a track expression: total number of bins, total number of bins whose value is NaN, min, max, sum, mean and standard deviation of the values.

Value

An array that represents summary statistics.

See Also

[gintervals.summary](#), [gbins.summary](#)

Examples

```
gdb.init_examples()
gsummary("rects_track")
```

| | |
|----------------|---|
| gsynth.bin_map | <i>Create a bin mapping from value-based merge specifications</i> |
|----------------|---|

Description

Converts value-based bin merge specifications into a `bin_map` named vector that can be used with [gsynth.train](#). This allows you to specify merges using actual track values rather than bin indices.

Usage

```
gsynth.bin_map(breaks, merge_ranges = NULL)
```

Arguments

| | |
|---------------------------|---|
| <code>breaks</code> | Numeric vector of bin boundaries (same as used in gsynth.train) |
| <code>merge_ranges</code> | List of merge specifications. Each specification is a named list with: <ul style="list-style-type: none"> from Numeric vector of length 2 <code>c(min, max)</code> defining the source value range to merge. Use <code>-Inf</code> or <code>Inf</code> for open-ended ranges. Can also be a single number (shorthand for <code>c(value, Inf)</code>). to Numeric vector of length 2 <code>c(min, max)</code> defining the target bin that source bins should map to. Must match an existing bin defined by <code>breaks</code>. |

Value

A named vector (`bin_map`) compatible with `bin_map` parameter in [gsynth.train](#). The names are source bin indices (1-based), and values are target bin indices (1-based).

See Also

[gsynth.train](#)

Examples

```
# Define breaks for GC content [0, 1] in 0.025 increments
breaks <- seq(0, 1, 0.025)

# Merge all GC content above 70% (0.7) into the bin (0.675, 0.7]
bin_map <- gsynth.bin_map(
  breaks = breaks,
  merge_ranges = list(
    list(from = 0.7, to = c(0.675, 0.7))
  )
)
```

```
)  
  
# Multiple merges: merge low GC (< 0.3) and high GC (> 0.7) into middle bins  
bin_map2 <- gsynth.bin_map(  
  breaks = breaks,  
  merge_ranges = list(  
    list(from = c(-Inf, 0.3), to = c(0.4, 0.425)), # low GC -> (0.4, 0.425]  
    list(from = 0.7, to = c(0.675, 0.7)) # high GC -> (0.675, 0.7]  
  )  
)
```

gsynth.convert

Convert a legacy RDS gsynth model to .gsm format

Description

Reads a gsynth.model from a legacy RDS file and saves it in the cross-platform .gsm format.

Usage

```
gsynth.convert(input_file, output_file, compress = FALSE)
```

Arguments

| | |
|-------------|---|
| input_file | Path to the legacy RDS model file |
| output_file | Path for the output .gsm model (directory or zip) |
| compress | Logical. If TRUE, save as a ZIP archive. If FALSE (default), save as a directory. |

Value

Invisibly returns the output file path.

See Also

[gsynth.save](#), [gsynth.load](#)

| | |
|-------------|--------------------------------------|
| gsynth.load | <i>Load a gsynth.model from disk</i> |
|-------------|--------------------------------------|

Description

Loads a previously saved Markov model. Auto-detects the format:

- If file is a directory, reads the .gsm directory format
- If file is a file, tries ZIP .gsm format first, then falls back to legacy RDS format

Usage

```
gsynth.load(file)
```

Arguments

| | |
|------|---|
| file | Path to the saved model (directory, .gsm zip, or legacy .rds) |
|------|---|

Value

A gsynth.model object

See Also

[gsynth.save](#), [gsynth.train](#), [gsynth.convert](#)

| | |
|---------------|---|
| gsynth.random | <i>Generate random genome sequences</i> |
|---------------|---|

Description

Generates random DNA sequences based on nucleotide probabilities without using a trained Markov model. Each nucleotide is sampled independently according to the specified probabilities.

Usage

```
gsynth.random(  
  intervals = NULL,  
  output_path = NULL,  
  output_format = c("misha", "fasta", "vector"),  
  nuc_probs = c(A = 0.25, C = 0.25, G = 0.25, T = 0.25),  
  mask_copy = NULL,  
  seed = NULL,  
  n_samples = 1,  
  iterator = 1  
)
```

Arguments

| | |
|----------------------------|--|
| <code>intervals</code> | Genomic intervals to sample. If NULL, uses all chromosomes. |
| <code>output_path</code> | Path to the output file (ignored when <code>output_format = "vector"</code>) |
| <code>output_format</code> | Output format: <ul style="list-style-type: none"> • "misha": .seq binary format (default) • "fasta": FASTA text format • "vector": Return sequences as a character vector (does not write to file) |
| <code>nuc_probs</code> | Nucleotide probabilities. Can be specified as: <ul style="list-style-type: none"> • A named vector: <code>c(A = 0.3, C = 0.2, G = 0.2, T = 0.3)</code> • An unnamed vector in A, C, G, T order: <code>c(0.3, 0.2, 0.2, 0.3)</code> Probabilities are automatically normalized to sum to 1. Default is uniform (0.25 each). |
| <code>mask_copy</code> | Optional intervals to copy from the original genome instead of random sampling. Use this to preserve specific regions exactly as they appear in the reference. |
| <code>seed</code> | Random seed for reproducibility. If NULL, uses current random state. |
| <code>n_samples</code> | Number of samples to generate per interval. Default is 1. |
| <code>iterator</code> | Iterator for position resolution. Default is 1 (base-pair resolution). Larger values may speed up processing but are typically not needed for random sampling. |

Details

Unlike `gsynth.sample` which uses a trained Markov model to generate sequences that preserve k-mer statistics, `gsynth.random` generates purely random sequences where each nucleotide is sampled independently. This is useful for generating baseline random sequences or sequences with specific GC content.

Nucleotide ordering: When using an unnamed vector for `nuc_probs`, the order is A, C, G, T. Named vectors can be in any order.

Value

When `output_format` is "misha" or "fasta", returns invisible NULL and writes the random sequences to `output_path`. When `output_format` is "vector", returns a character vector of sequences (length = `n_intervals * n_samples`).

See Also

[gsynth.sample](#), [gsynth.train](#)

Examples

```
gdb.init_examples()

# Generate random sequences with uniform nucleotide probabilities
seqs <- gsynth.random(
```

```
    intervals = gintervals(1, 0, 1000),
    output_format = "vector",
    seed = 42
)

# Generate GC-rich sequences (60% GC)
gc_rich <- gsynth.random(
  intervals = gintervals(1, 0, 1000),
  output_format = "vector",
  nuc_probs = c(A = 0.2, C = 0.3, G = 0.3, T = 0.2),
  seed = 42
)

# Generate AT-rich sequences
at_rich <- gsynth.random(
  intervals = gintervals(1, 0, 1000),
  output_format = "vector",
  nuc_probs = c(A = 0.35, C = 0.15, G = 0.15, T = 0.35),
  seed = 42
)
```

gsynth.replace_kmer *Iteratively replace a k-mer in the genome*

Description

Performs an iterative replacement of a target k-mer with a replacement sequence. This is useful for creating synthetic genomes with specific motifs removed (e.g., creating a CpG-null genome by iteratively swapping CG to GC).

Usage

```
gsynth.replace_kmer(
  target,
  replacement,
  output_path = NULL,
  output_format = c("misha", "fasta", "vector"),
  intervals = NULL,
  check_composition = TRUE
)
```

Arguments

| | |
|---------------|--|
| target | The k-mer sequence to remove (e.g., "CG"). |
| replacement | The replacement sequence (e.g., "GC"). |
| output_path | Path to the output file (ignored when output_format = "vector"). |
| output_format | Output format: |

- "misha": .seq binary format (default)
 - "fasta": FASTA text format
 - "vector": Return sequences as a character vector (does not write to file)
- intervals Genomic intervals to process. If NULL, uses all chromosomes.
- check_composition Logical. If TRUE (default), ensures target and replacement have the same nucleotide composition (preserving exact base counts).

Details

Bubble Sort / Iterative Logic: The function scans the sequence and replaces occurrences of target with replacement. If a replacement creates a new instance of target (e.g., removing "CG" with "GC" in the sequence "CCG" -> "CGC"), the new instance is also replaced. This continues until the sequence is free of the target k-mer.

When target and replacement are permutations of each other (e.g., "CG" and "GC"), this acts as a "bubble sort" of nucleotides, moving bases locally without altering the total GC content or base counts of the genome.

Value

When output_format is "misha" or "fasta", returns invisible NULL and writes to output_path. When output_format is "vector", returns a character vector of modified sequences.

Examples

```
## Not run:
# Robust removal of all CpG dinucleotides (preserving GC%)
gsynth.replace_kmer(
  target = "CG",
  replacement = "GC",
  output_path = "genome_no_cpG.seq",
  output_format = "misha"
)

## End(Not run)
```

gsynth.sample

Sample a synthetic genome from a trained Markov model

Description

Generates a synthetic genome by sampling from a trained stratified Markov-5 model. The generated genome preserves the k-mer statistics of the original genome within each stratification bin.

Usage

```
gsynth.sample(
  model,
  output_path = NULL,
  output_format = c("misha", "fasta", "vector"),
  mask_copy = NULL,
  seed = NULL,
  intervals = NULL,
  n_samples = 1,
  bin_merge = NULL
)
```

Arguments

| | |
|---------------|---|
| model | A gsynth.model object from gsynth.train |
| output_path | Path to the output file (ignored when output_format = "vector") |
| output_format | Output format: <ul style="list-style-type: none"> • "misha": .seq binary format (default) • "fasta": FASTA text format • "vector": Return sequences as a character vector (does not write to file) |
| mask_copy | Optional intervals to copy from the original genome instead of sampling. Use this to preserve specific regions (e.g., repeats, regulatory elements) exactly as they appear in the reference. Regions not in mask_copy will be sampled using the Markov model. Note: mask_copy intervals should be non-overlapping and sorted by start position within each chromosome. Overlapping intervals may result in only the first overlapping region being copied, with subsequent overlaps skipped due to cursor advancement during sequential processing. |
| seed | Random seed for reproducibility. If NULL, uses current random state. |
| intervals | Genomic intervals to sample. If NULL, uses all chromosomes. |
| n_samples | Number of samples to generate per interval. Default is 1. When n_samples > 1 and output_format = "fasta", headers include "_sampleN". When output_format = "vector", returns n_samples * n_intervals sequences. |
| bin_merge | Optional list of bin merge specifications to apply during sampling, one per dimension (length must equal model\$n_dims). Each element should be: <ul style="list-style-type: none"> • A list of merge specifications (same format as in gsynth.train: each spec is list(from = ..., to = ...)) • Or NULL to use the bin mapping from training for that dimension This allows merging sparse bins at sampling time without re-training. Example for a 2D model: <pre>bin_merge = list(# Dimension 1: merge bins with values >= 0.8 to bin [0.7, 0.8) list(list(from = c(0.8, Inf), to = c(0.7, 0.8))), # Dimension 2: use training-time bin_map (no override) NULL)</pre> |

Details

N bases during sampling: When the sampler needs to initialize the first 5-mer context and encounters regions with only N bases, it falls back to uniform random base selection until a valid context is established. Similarly, if a bin has no learned statistics (sparse bin with NA CDF), uniform sampling is used for that position.

Sparse bins: If the model has sparse bins (from `min_obs` during training), a warning is issued when sampling regions that fall into these bins. Consider using `bin_merge` to redirect sparse bins to well-populated ones.

Value

When `output_format` is "misha" or "fasta", returns invisible NULL and writes the synthetic genome to `output_path`. When `output_format` is "vector", returns a character vector of sequences (length = `n_intervals * n_samples`).

See Also

[gsynth.train](#), [gsynth.save](#)

Examples

```
gdb.init_examples()

# Create virtual tracks
gvtrack.create("g_frac", NULL, "kmer.frac", kmer = "G")
gvtrack.create("c_frac", NULL, "kmer.frac", kmer = "C")
gvtrack.create("cg_frac", NULL, "kmer.frac", kmer = "CG")
gvtrack.create("masked_frac", NULL, "masked.frac")

# Define repeat mask (regions to preserve from original)
repeats <- gscreen("masked_frac > 0.5",
  intervals = gintervals.all(),
  iterator = 100
)

# Train model (excluding repeats from training)
model <- gsynth.train(
  list(expr = "g_frac + c_frac", breaks = seq(0, 1, 0.025)),
  list(expr = "cg_frac", breaks = c(0, 0.01, 0.02, 0.03, 0.04, 0.2)),
  mask = repeats,
  iterator = 200,
  min_obs = 1000
)

# Sample with mask_copy to preserve repeats from original genome
temp_dir <- tempdir()
synthetic_genome_file <- file.path(temp_dir, "synthetic_genome.fa")
gsynth.sample(model, synthetic_genome_file,
  output_format = "fasta",
  mask_copy = repeats,
  seed = 60427,
```

```

    bin_merge = list(
      list(list(from = 0.7, to = c(0.675, 0.7))),
      list(list(from = 0.04, to = c(0.03, 0.04)))
    )
  )
)

```

 gsynth.save

Save a gsynth.model to disk in .gsm format

Description

Saves a trained Markov model in the cross-platform .gsm format, which consists of a metadata YAML file and raw binary arrays for counts and CDFs. The .gsm format can be stored as a directory (default) or a ZIP archive.

Usage

```
gsynth.save(model, file, compress = FALSE)
```

Arguments

| | |
|----------|---|
| model | A gsynth.model object from gsynth.train |
| file | Path to save the model (directory or .zip file) |
| compress | Logical. If TRUE, save as a ZIP archive. If FALSE (default), save as a directory. |

Value

Invisibly returns the file path.

See Also

[gsynth.load](#), [gsynth.train](#), [gsynth.convert](#)

 gsynth.train

Train a stratified Markov-5 model from genome sequences

Description

Computes a 5th-order Markov model optionally stratified by bins of one or more track expressions (e.g., GC content and CG dinucleotide frequency). This model can be used to generate synthetic genomes that preserve the k-mer statistics of the original genome within each stratification bin. When called with no dimension specifications, trains a single unstratified model.

Usage

```
gsynth.train(
  ...,
  mask = NULL,
  intervals = NULL,
  iterator = NULL,
  pseudocount = 1,
  min_obs = 0
)
```

Arguments

| | |
|-------------|--|
| ... | Zero or more dimension specifications. Each specification is a list containing: expr Track expression for this dimension (required) breaks Numeric vector of bin boundaries for this dimension (required) bin_merge Optional list of merge specifications for merging sparse bins. Each specification is a named list with 'from' and 'to' elements. If no dimensions are provided, trains an unstratified model with a single bin. |
| mask | Optional intervals to exclude from training. Regions in the mask will not contribute to k-mer counts. Can be computed using <code>gscreen()</code> . |
| intervals | Genomic intervals to process. If NULL, uses all chromosomes. |
| iterator | Iterator for track evaluation, determines the resolution at which track values are computed. |
| pseudocount | Pseudocount added to all k-mer counts to avoid zero probabilities. Default is 1. |
| min_obs | Minimum number of observations (6-mers) required per bin. Bins with fewer observations will be marked as NA (not learned) and a warning will be issued. Default is 0 (no minimum). During sampling, NA bins will fall back to uniform sampling unless merged via <code>bin_merge</code> . |

Details

Strand symmetry: The training process counts both the forward strand 6-mer and its reverse complement for each position, ensuring strand-symmetric transition probabilities. This means the reported `total_kmers` is approximately double the number of genomic positions processed.

N bases: Positions where the 6-mer contains any N (unknown) bases are skipped during training and counted in `total_n`. The model only learns from valid A/C/G/T sequences.

Value

A `gsynth.model` object containing:

n_dims Number of stratification dimensions

dim_specs List of dimension specifications (`expr`, `breaks`, `num_bins`, `bin_map`)

dim_sizes Vector of bin counts per dimension

total_bins Total number of bins (product of `dim_sizes`)

total_kmers Total number of valid 6-mers counted
per_bin_kmers Number of 6-mers counted per bin
total_masked Number of positions skipped due to mask
total_n Number of positions skipped due to N bases
model_data Internal model data (counts and CDFs)

See Also

[gsynth.sample](#), [gsynth.save](#), [gsynth.load](#), [gsynth.bin_map](#)

Examples

```
gdb.init_examples()

# Create virtual tracks for stratification
gvtrack.create("g_frac", NULL, "kmer.frac", kmer = "G")
gvtrack.create("c_frac", NULL, "kmer.frac", kmer = "C")
gvtrack.create("cg_frac", NULL, "kmer.frac", kmer = "CG")
gvtrack.create("masked_frac", NULL, "masked.frac")

# Define repeat mask
repeats <- gscreen("masked_frac > 0.5",
  intervals = gintervals.all(),
  iterator = 100
)

# Train unstratified model (no stratification)
model_0d <- gsynth.train(
  mask = repeats,
  intervals = gintervals.all(),
  iterator = 200
)

# Train model with 2D stratification (GC content and CG dinucleotide)
model <- gsynth.train(
  list(
    list(
      expr = "g_frac + c_frac",
      breaks = seq(0, 1, 0.025),
      bin_merge = list(list(from = 0.7, to = c(0.675, 0.7)))
    ),
    list(
      expr = "cg_frac",
      breaks = c(0, 0.01, 0.02, 0.03, 0.04, 0.2),
      bin_merge = list(list(from = 0.04, to = c(0.03, 0.04)))
    )
  ),
  mask = repeats,
  intervals = gintervals.all(),
  iterator = 200
)
```

`gtrack.2d.convert_to_indexed`*Convert 2D track to indexed format*

Description

Converts a per-chromosome-pair 2D track (rectangles or points) to indexed format (track.dat + track.idx). This reduces file descriptor usage from $O(N^2)$ to $O(1)$, which is especially beneficial for genomes with many contigs.

Usage

```
gtrack.2d.convert_to_indexed(track = NULL, remove.old = FALSE, force = FALSE)
```

Arguments

| | |
|------------|--|
| track | track name to convert |
| remove.old | Logical. If TRUE, removes old per-chromosome-pair files after successful conversion. Default: FALSE. |
| force | Logical. If TRUE, re-converts even if already in indexed format. Default: FALSE. |

Value

None.

See Also

[gtrack.2d.create](#), [gtrack.2d.import](#), [gtrack.2d.import_contacts](#), [gtrack.convert_to_indexed](#), [gdb.convert_to_indexed](#)

Examples

```
## Not run:  
# Convert a 2D track to indexed format  
gtrack.2d.convert_to_indexed("my_2d_track")  
  
# Convert and remove old per-pair files  
gtrack.2d.convert_to_indexed("my_2d_track", remove.old = TRUE)  
  
# Force re-conversion  
gtrack.2d.convert_to_indexed("my_2d_track", force = TRUE)  
  
## End(Not run)
```

| | |
|------------------|---|
| gtrack.2d.create | <i>Creates a 'Rectangles' track from intervals and values</i> |
|------------------|---|

Description

Creates a 'Rectangles' track from intervals and values.

Usage

```
gtrack.2d.create(  
  track = NULL,  
  description = NULL,  
  intervals = NULL,  
  values = NULL  
)
```

Arguments

| | |
|-------------|--|
| track | track name |
| description | a character string description |
| intervals | a set of two-dimensional intervals |
| values | an array of numeric values - one for each interval |

Details

This function creates a new 'Rectangles' (two-dimensional) track with values at given intervals. 'description' is added as a track attribute.

When multiple databases are connected via [gsetroot](#), the track is created in the current working directory (.misha\$GWD), which defaults to the last connected database. Use [gdir.cd](#) with an absolute path to change where new tracks are created.

Value

None.

See Also

[gtrack.create](#), [gtrack.create_sparse](#), [gtrack.smooth](#), [gtrack.modify](#), [gtrack.rm](#), [gtrack.info](#), [gdir.create](#), [gtrack.attr.get](#)

Examples

```
gdb.init_examples()  
intervs1 <- gintervals.2d(  
  1, (1:4) * 200, (1:4) * 200 + 100,  
  1, (1:4) * 300, (1:4) * 300 + 200
```

```

)
intervs2 <- gintervals.2d(
  "X", (7:10) * 100, (7:10) * 100 + 50,
  2, (1:4) * 200, (1:4) * 200 + 130
)
intervs <- rbind(intervs1, intervs2)
gtrack.2d.create(
  "test_rects", "Test 2d track", intervs,
  runif(dim(intervs)[1], 1, 100)
)
gextract("test_rects", .misha$ALLGENOME)
gtrack.rm("test_rects", force = TRUE)

```

| | |
|------------------|---|
| gtrack.2d.import | <i>Creates a 2D track from tab-delimited file</i> |
|------------------|---|

Description

Creates a 2D track from tab-delimited file(s).

Usage

```
gtrack.2d.import(track = NULL, description = NULL, file = NULL)
```

Arguments

| | |
|-------------|--------------------------------|
| track | track name |
| description | a character string description |
| file | vector of file paths |

Details

This function creates a 2D track track from one or more tab-delimited files. Each file must start with a header describing the columns. The first 6 columns must have the following names: 'chrom1', 'start1', 'end1', 'chrom2', 'start2', 'end2'. The last column is designated for the value and it may have an arbitrary name. The header is followed by a list of intervals and a value for each interval. Overlapping intervals are forbidden.

One can learn about the format of the tab-delimited file by running 'gextract' function on a 2D track with a 'file' parameter set to the name of the file.

If all the imported intervals represent a point (i.e. end == start + 1) a 'Points' track is created otherwise it is a 'Rectangles' track.

'description' is added as a track attribute.

Note: temporary files are created in the directory of the track during the run of the function. A few of them need to be kept simultaneously open. If the number of chromosomes and / or intervals is particularly high, a few thousands files might be needed to be opened simultaneously. Some

operating systems limit the number of open files per user, in which case the function might fail with "Too many open files" or similar error. The workaround could be:

1. Increase the limit of simultaneously opened files (the way varies depending on your operating system).
2. Increase the value of 'gmax.data.size' option. Higher values of 'gmax.data.size' option will increase memory usage of the function but create fewer temporary files.

Value

None.

See Also

[gtrack.rm](#), [gtrack.info](#), [gdir.create](#)

gtrack.2d.import_contacts

Creates a track from a file of inter-genomic contacts

Description

Creates a track from a file of inter-genomic contacts.

Usage

```
gtrack.2d.import_contacts(  
  track = NULL,  
  description = NULL,  
  contacts = NULL,  
  fends = NULL,  
  allow.duplicates = TRUE  
)
```

Arguments

| | |
|------------------|---|
| track | track name |
| description | a character string description |
| contacts | vector of contacts files |
| fends | name of fragment ends file |
| allow.duplicates | if 'TRUE' duplicated contacts are allowed |

Details

This function creates a 'Points' (two-dimensional) track from contacts files. If 'allow.duplicates' is 'TRUE' duplicated contacts are allowed and summed up, otherwise an error is reported.

Contacts (coord1, coord2) within the same chromosome are automatically doubled to include also '(coord2, coord1)' unless 'coord1' equals to 'coord2'.

Contacts may come in one or more files.

If 'fends' is 'NULL' contacts file is expected to be in "intervals-value" tab-separated format. The file starts with a header defining the column names. The first 6 columns must have the following names: 'chrom1', 'start1', 'end1', 'chrom2', 'start2', 'end2'. The last column is designated for the value and it may have an arbitrary name. The header is followed by a list of intervals and a value for each interval. An interval of form (chrom1, start1, end1, chrom2, start2, end2) is added as a point (X, Y) to the resulted track where $X = (start1 + end1) / 2$ and $Y = (start2 + end2) / 2$.

One can see an example of "intervals-value" format by running 'gextract' function on a 2D track with a 'file' parameter set to the name of the file.

If 'fends' is not 'NULL' contacts file is expected to be in "fends-value" tab-separated format. It should start with a header containing at least 3 column names 'fend1', 'fend2' and 'count' in arbitrary order followed by lines each defining a contact between two fragment ends.

| COLUMN | VALUE | DESCRIPTION |
|--------|---------|-----------------------------------|
| fend1 | Integer | ID of the first fragment end |
| fend2 | Integer | ID of the second fragment end |
| count | Numeric | Value associated with the contact |

A fragment ends file is also in tab-separated format. It should start with a header containing at least 3 column names 'fend', 'chr' and 'coord' in arbitrary order followed by lines each defining a single fragment end.

| COLUMN | VALUE | DESCRIPTION |
|--------|-----------------|--|
| fend | Unique integer | ID of the fragment end |
| chr | Chromosome name | Can be specified with or without "chr" prefix, like: "X" or "chrX" |
| coord | Integer | Coordinate |

'description' is added as a track attribute.

Note: temporary files are created in the directory of the track during the run of the function. A few of them need to be kept simultaneously open. If the number of chromosomes and / or contacts is particularly high, a few thousands files might be needed to be opened simultaneously. Some operating systems limit the number of open files per user, in which case the function might fail with "Too many open files" or similar error. The workaround could be:

1. Increase the limit of simultaneously opened files (the way varies depending on your operating system).
2. Increase the value of 'gmax.data.size' option. Higher values of 'gmax.data.size' option will increased memory usage of the function but create fewer temporary files.

Value

None.

See Also

[gtrack.2d.import](#), [gtrack.rm](#), [gtrack.info](#), [gdir.create](#)

`gtrack.array.extract` *Returns values from 'Array' track*

Description

Returns values from 'Array' track.

Usage

```
gtrack.array.extract(
  track = NULL,
  slice = NULL,
  intervals = NULL,
  file = NULL,
  intervals.set.out = NULL
)
```

Arguments

| | |
|--------------------------------|---|
| <code>track</code> | track name |
| <code>slice</code> | a vector of column names or column indices or 'NULL' |
| <code>intervals</code> | genomic scope for which the function is applied |
| <code>file</code> | file name where the function result is to be saved. If 'NULL' result is returned to the user. |
| <code>intervals.set.out</code> | intervals set name where the function result is optionally outputted |

Details

This function returns the column values of an 'Array' track in the genomic scope specified by 'intervals'. 'slice' parameter determines which columns should appear in the result. The columns can be indicated by their names or their indices. If 'slice' is 'NULL' the values of all track columns are returned.

The order inside the result might not be the same as the order of intervals. An additional column 'intervalID' is added to the return value. Use this column to refer to the index of the original interval from the supplied 'intervals'.

If 'file' parameter is not 'NULL' the result is saved to a tab-delimited text file (without 'intervalID' column) rather than returned to the user. This can be especially useful when the result is too big to fit into the physical memory. The resulted file can be used as an input for 'gtrack.array.import' function.

If 'intervals.set.out' is not 'NULL' the result is saved as an intervals set. Similarly to 'file' parameter 'intervals.set.out' can be useful to overcome the limits of the physical memory.

Value

If 'file' and 'intervals.set.out' are 'NULL' a set of intervals with additional columns for 'Array' track column values and 'columnID'.

See Also

[gextract](#), [gtrack.array.get_colnames](#), [gtrack.array.import](#)

Examples

```
gdb.init_examples()
gtrack.array.extract(
  "array_track", c("col3", "col5"),
  gintervals(1, 0, 2000)
)
```

`gtrack.array.get_colnames`

Returns column names of array track

Description

Returns column names of array track.

Usage

```
gtrack.array.get_colnames(track = NULL)
```

Arguments

track track name

Details

This function returns the column names of an array track.

Value

A character vector with column names.

See Also

[gtrack.array.set_colnames](#), [gtrack.array.extract](#), [gvtrack.array.slice](#), [gtrack.info](#)

Examples

```
gtrack.array.get_colnames("array_track")
```

```
gtrack.array.import      Creates an array track from array tracks or files
```

Description

Creates an array track from array tracks or files.

Usage

```
gtrack.array.import(track = NULL, description = NULL, ...)
```

Arguments

| | |
|-------------|---|
| track | name of the newly created track |
| description | a character string description |
| ... | array track or name of a tab-delimited file |

Details

This function creates a new 'Array' track from one or more "sources". Each source can be either another 'Array' track or a tab-delimited file that contains one-dimensional intervals and column values that should be added to the newly created track. One can learn about the exact format of the file by running 'gtrack.array.extract' or 'gextract' functions with a 'file' parameter and inspecting the output file.

There might be more than one source used to create the new track. In that case the new track will contain the columns from all the sources. The equally named columns are merged. Intervals that appear in one source but not in the other are added and the values for the missing columns are set to NaN. Intervals with all NaN values are not added. Partial overlaps between two intervals from different sources are forbidden.

'description' is added as a track attribute.

Value

None.

See Also

[gextract](#), [gtrack.array.extract](#), [gtrack.array.set_colnames](#), [gtrack.rm](#), [gtrack.info](#), [gdir.create](#)

Examples

```
f1 <- tempfile()
gextract("sparse_track", gintervals(1, 5000, 20000), file = f1)
f2 <- tempfile()
gtrack.array.extract("array_track", c("col2", "col3", "col4"),
  gintervals(1, 0, 20000),
  file = f2
)
f3 <- tempfile()
gtrack.array.extract("array_track", c("col1", "col3"),
  gintervals(1, 0, 20000),
  file = f3
)

gtrack.array.import("test_track1", "Test array track 1", f1, f2)
gtrack.array.extract("test_track1", NULL, .misha$ALLGENOME)

gtrack.array.import(
  "test_track2", "Test array track 2",
  "test_track1", f3
)
gtrack.array.extract("test_track2", NULL, .misha$ALLGENOME)

gtrack.rm("test_track1", TRUE)
gtrack.rm("test_track2", TRUE)
unlink(c(f1, f2, f3))
```

```
gtrack.array.set_colnames
      Sets column names of array track
```

Description

Sets column names of array track.

Usage

```
gtrack.array.set_colnames(track = NULL, names = NULL)
```

Arguments

| | |
|-------|------------------------|
| track | track name |
| names | vector of column names |

Details

This sets the column names of an array track.

Value

None.

See Also

[gtrack.array.get_colnames](#), [gtrack.array.extract](#), [gvtrack.array.slice](#), [gtrack.info](#)

Examples

```
old.names <- gtrack.array.get_colnames("array_track")
new.names <- paste("modified", old.names, sep = "_")
gtrack.array.set_colnames("array_track", new.names)
gtrack.array.get_colnames("array_track")
gtrack.array.set_colnames("array_track", old.names)
gtrack.array.get_colnames("array_track")
```

`gtrack.attr.export` *Returns track attributes values*

Description

Returns track attributes values.

Usage

```
gtrack.attr.export(tracks = NULL, attrs = NULL)
```

Arguments

| | |
|---------------------|---------------------------------------|
| <code>tracks</code> | a vector of track names or 'NULL' |
| <code>attrs</code> | a vector of attribute names or 'NULL' |

Details

This function returns a data frame that contains track attributes values. Column names of the data frame consist of the attribute names, row names contain the track names.

The list of required tracks is specified by 'tracks' argument. If 'tracks' is 'NULL' the attribute values of all existing tracks are returned.

Likewise the list of required attributes is controlled by 'attrs' argument. If 'attrs' is 'NULL' all attribute values of the specified tracks are returned. The columns are also sorted then by "popularity" of an attribute, i.e. the number of tracks containing this attribute. This sorting is not applied if 'attrs' is not 'NULL'.

Empty character string in a table cell marks a non-existing attribute.

Value

A data frame containing track attributes values.

See Also

[gtrack.attr.import](#), [gtrack.attr.get](#), [gtrack.attr.set](#)

Examples

```
gdb.init_examples()
gtrack.attr.export()
gtrack.attr.export(tracks = c("sparse_track", "dense_track"))
gtrack.attr.export(attrs = "created.by")
```

| | |
|------------------------------|---|
| <code>gtrack.attr.get</code> | <i>Returns value of a track attribute</i> |
|------------------------------|---|

Description

Returns value of a track attribute.

Usage

```
gtrack.attr.get(track = NULL, attr = NULL)
```

Arguments

| | |
|--------------------|----------------|
| <code>track</code> | track name |
| <code>attr</code> | attribute name |

Details

This function returns the value of a track attribute. If the attribute does not exist an empty string is returned.

Value

Track attribute value.

See Also

[gtrack.attr.import](#), [gtrack.attr.set](#)

Examples

```
gdb.init_examples()
gtrack.attr.set("sparse_track", "test_attr", "value")
gtrack.attr.get("sparse_track", "test_attr")
gtrack.attr.set("sparse_track", "test_attr", "")
```

`gtrack.attr.import` *Imports track attributes values*

Description

Imports track attributes values.

Usage

```
gtrack.attr.import(table = NULL, remove.others = FALSE)
```

Arguments

`table` a data frame containing attribute values
`remove.others` specifies what to do with the attributes that are not in the table

Details

This function makes imports attribute values contained in a data frame 'table'. The format of a table is similar to the one returned by 'gtrack.attr.export'. The values of the table must be character strings. Column names of the table should specify the attribute names, while row names should contain the track names.

The specified attributes of the specified tracks are modified. If an attribute value is an empty string this attribute is removed from the track.

If 'remove.others' is 'TRUE' all non-readonly attributes that do not appear in the table are removed, otherwise they are preserved unchanged.

Error is reported on an attempt to modify a value of a read-only attribute.

Value

None.

See Also

[gtrack.attr.import](#), [gtrack.attr.set](#), [gtrack.attr.get](#), [gdb.get_readonly_attrs](#)

Examples

```
gdb.init_examples()
t <- gtrack.attr.export()
t$newattr <- as.character(1:dim(t)[1])
gtrack.attr.import(t)
gtrack.attr.export(attrs = "newattr")

# roll-back the changes
t$newattr <- ""
gtrack.attr.import(t)
```

| | |
|-----------------|---|
| gtrack.attr.set | <i>Assigns value to a track attribute</i> |
|-----------------|---|

Description

Assigns value to a track attribute.

Usage

```
gtrack.attr.set(track = NULL, attr = NULL, value = NULL)
```

Arguments

| | |
|-------|----------------|
| track | track name |
| attr | attribute name |
| value | value |

Details

This function creates a track attribute and assigns 'value' to it. If the attribute already exists its value is overwritten.

If 'value' is an empty string the attribute is removed.

Error is reported on an attempt to modify a value of a read-only attribute.

Value

None.

See Also

[gtrack.attr.get](#), [gtrack.attr.import](#), [gtrack.var.set](#), [gdb.get_readonly_attrs](#)

Examples

```
gdb.init_examples()
gtrack.attr.set("sparse_track", "test_attr", "value")
gtrack.attr.get("sparse_track", "test_attr")
gtrack.attr.set("sparse_track", "test_attr", "")
```

| | |
|----------------|--|
| gtrack.convert | <i>Converts a track to the most current format</i> |
|----------------|--|

Description

Converts a track (if needed) to the most current format.

Usage

```
gtrack.convert(src.track = NULL, tgt.track = NULL)
```

Arguments

| | |
|-----------|---|
| src.track | source track name |
| tgt.track | target track name. If 'NULL' the source track is overwritten. |

Details

This function converts a track to the most current format. It should be used if a track created by an old version of the library cannot be read anymore by the newer version. The old track is given by 'src.track'. After conversion a new track 'tgt.track' is created. If 'tgt.track' is 'NULL' the source track is overwritten.

Value

None

See Also

[gtrack.create](#), [gtrack.2d.create](#), [gtrack.create_sparse](#)

`gtrack.convert_to_indexed`*Convert a track to indexed format*

Description

Converts a per-chromosome track to indexed format (track.dat + track.idx).

Usage

```
gtrack.convert_to_indexed(track = NULL)
```

Arguments

| | |
|-------|-----------------------|
| track | track name to convert |
|-------|-----------------------|

Details

This function converts a track from the per-chromosome file format to single-file indexed format. The indexed format dramatically reduces file descriptor usage for genomes with many contigs and provides better performance for parallel access.

The function performs the following steps:

1. Validates that all per-chromosome files have consistent metadata
2. Creates track.dat by concatenating all per-chromosome files
3. Creates track.idx with offset/length information for each chromosome
4. Uses atomic operations (fsync + rename) to ensure data integrity
5. Removes the old per-chromosome files after successful conversion

Value

None

See Also

[gtrack.create](#), [gtrack.create_sparse](#), [gtrack.create_dense](#)

Examples

```
## Not run:  
# Convert a track to indexed format  
gtrack.convert_to_indexed("my_track")  
  
## End(Not run)
```

| | |
|-------------|-----------------------|
| gtrack.copy | <i>Copies a track</i> |
|-------------|-----------------------|

Description

Creates a copy of an existing track.

Usage

```
gtrack.copy(src = NULL, dest = NULL)
```

Arguments

| | |
|------|------------------------|
| src | source track name |
| dest | destination track name |

Details

This function creates a copy of a track. The new track is created in the current working directory (.misha\$GWD), which may be in a different database than the source track when multiple databases are connected.

Value

None.

See Also

[gtrack.mv](#), [gtrack.rm](#), [gtrack.exists](#), [gtrack.ls](#)

Examples

```
gdb.init_examples()
gtrack.copy("dense_track", "dense_track_copy")
gtrack.exists("dense_track_copy")
gtrack.rm("dense_track_copy", force = TRUE)
```

| | |
|---------------|--|
| gtrack.create | <i>Creates a track from a track expression</i> |
|---------------|--|

Description

Creates a track from a track expression.

Usage

```
gtrack.create(
    track = NULL,
    description = NULL,
    expr = NULL,
    iterator = NULL,
    band = NULL
)
```

Arguments

| | |
|-------------|---|
| track | track name |
| description | a character string description |
| expr | track expression |
| iterator | track expression iterator. If 'NULL' iterator is determined implicitly based on track expression. |
| band | track expression band. If 'NULL' no band is used. |

Details

This function creates a new track named track. The values of the track are determined by evaluation of 'expr' - a numeric track expression. The type of the new track is determined by the type of the iterator. 'Fixed bin', 'Sparse' or 'Rectangles' track can be created accordingly. 'description' is added as a track attribute.

When multiple databases are connected via [gsetroot](#), the track is created in the current working directory (.misha\$GWD), which defaults to the last connected database. Use [gdir.cd](#) with an absolute path to change where new tracks are created.

Value

None.

See Also

[gtrack.2d.create](#), [gtrack.create_sparse](#), [gtrack.smooth](#), [gtrack.modify](#), [gtrack.rm](#), [gtrack.info](#), [gdir.create](#)

Examples

```

gdb.init_examples()

## Creates a new track that is a sum of values from 'dense' and
## 2 * non-nan values of 'sparse' track. The new track type is
## Dense with a bin size that equals to '70'.
gtrack.create("mixed_track", "Test track",
             "dense_track +
             replace(sparse_track, is.nan(sparse_track), 0) * 2",
             iterator = 70
            )
gtrack.info("mixed_track")
gtrack.rm("mixed_track", force = TRUE)

```

`gtrack.create_dense` *Creates a 'Dense' track from intervals and values*

Description

Creates a 'Dense' track from intervals and values.

Usage

```

gtrack.create_dense(
  track = NULL,
  description = NULL,
  intervals = NULL,
  values = NULL,
  binsize = NULL,
  defval = NaN
)

```

Arguments

| | |
|--------------------------|--|
| <code>track</code> | track name |
| <code>description</code> | a character string description |
| <code>intervals</code> | a set of one-dimensional intervals |
| <code>values</code> | an array of numeric values - one for each interval |
| <code>binsize</code> | bin size of the newly created 'Dense' track |
| <code>defval</code> | default track value for genomic regions not covered by the intervals |

Details

This function creates a new 'Dense' track with values at given intervals. 'description' is added as a track attribute.

Value

None.

See Also

[gtrack.create_sparse](#), [gtrack.import](#), [gtrack.modify](#), [gtrack.rm](#), [gtrack.info](#)

Examples

```
gdb.init_examples()
intervs <- gintervals.load("annotations")
gtrack.create_dense(
  "test_dense", "Test dense track", intervs,
  1:dim(intervs)[1], 50, 0
)
gextract("test_dense", .misha$ALLGENOME)
gtrack.rm("test_dense", force = TRUE)
```

`gtrack.create_dirs` *Create directories needed for track creation*

Description

This function creates the directories needed for track creation. For example, if the track name is 'proj.sample.my_track', this function creates the directories 'proj' and 'sample'. Use this function with caution - a long track name may create a deep directory structure.

Usage

```
gtrack.create_dirs(track, mode = "0777")
```

Arguments

| | |
|-------|-------------------|
| track | name of the track |
| mode | see 'dir.create' |

Value

None.

Examples

```
gdb.init_examples()

# This creates the directories 'proj' and 'sample'
gtrack.create_dirs("proj.sample.my_track")
```

`gtrack.create_pwm_energy`*Creates a new track from PSSM energy function*

Description

Creates a new track from PSSM energy function.

Usage

```
gtrack.create_pwm_energy(  
    track = NULL,  
    description = NULL,  
    pssmset = NULL,  
    pssmid = NULL,  
    prior = NULL,  
    iterator = NULL  
)
```

Arguments

| | |
|--------------------------|---|
| <code>track</code> | track name |
| <code>description</code> | a character string description |
| <code>pssmset</code> | name of PSSM set: 'pssmset.key' and 'pssmset.data' must be presented in 'GROOT/pssms' directory |
| <code>pssmid</code> | PSSM id |
| <code>prior</code> | prior |
| <code>iterator</code> | track expression iterator for the newly created track |

Details

This function creates a new track with values of a PSSM energy function. PSSM parameters (nucleotide probability per position and pluralization) are determined by 'pssmset' key and data files ('pssmset.key' and 'pssmset.data'). These two files must be located in 'GROOT/pssms' directory. The type of the created track is determined by the type of the iterator. 'description' is added as a track attribute.

Value

None.

See Also

[gtrack.create](#), [gtrack.2d.create](#), [gtrack.create_sparse](#), [gtrack.smooth](#), [gtrack.modify](#), [gtrack.rm](#), [gtrack.info](#), [gdir.create](#)

Examples

```
gdb.init_examples()
gtrack.create_pwm_energy("pwm_energy_track", "Test track", "pssm",
    3, 0.01,
    iterator = 100
)
gextract("pwm_energy_track", gintervals(1, 0, 1000))
```

`gtrack.create_sparse` *Creates a 'Sparse' track from intervals and values*

Description

Creates a 'Sparse' track from intervals and values.

Usage

```
gtrack.create_sparse(
    track = NULL,
    description = NULL,
    intervals = NULL,
    values = NULL
)
```

Arguments

| | |
|--------------------------|--|
| <code>track</code> | track name |
| <code>description</code> | a character string description |
| <code>intervals</code> | a set of one-dimensional intervals |
| <code>values</code> | an array of numeric values - one for each interval |

Details

This function creates a new 'Sparse' track with values at given intervals. 'description' is added as a track attribute.

When multiple databases are connected via [gsetroot](#), the track is created in the current working directory (`.misha$GWD`), which defaults to the last connected database. Use [gdir.cd](#) with an absolute path to change where new tracks are created.

Value

None.

See Also

[gtrack.create](#), [gtrack.2d.create](#), [gtrack.smooth](#), [gtrack.modify](#), [gtrack.rm](#), [gtrack.info](#), [gdir.create](#)

Examples

```
gdb.init_examples()
intervs <- gintervals.load("annotations")
gtrack.create_sparse(
  "test_sparse", "Test track", intervvs,
  1:dim(intervvs)[1]
)
gextract("test_sparse", .misha$ALLGENOME)
gtrack.rm("test_sparse", force = TRUE)
```

| | |
|-----------------------------|--|
| <code>gtrack.dataset</code> | <i>Returns the database/dataset path for a track</i> |
|-----------------------------|--|

Description

Returns the path of the database or dataset containing a track.

Usage

```
gtrack.dataset(track = NULL)
```

Arguments

`track` track name or a vector of track names

Details

When datasets are loaded, tracks can come from either the working database or from loaded datasets. This function returns the source path for each track.

Value

Character vector of database/dataset paths. Returns NA for non-existent tracks.

See Also

[gtrack.dbs](#), [gtrack.exists](#), [gtrack.ls](#), [gdataset.ls](#)

Examples

```
gdb.init_examples()
gtrack.dataset("dense_track")
```

| | |
|------------|---|
| gtrack.dbs | <i>Returns the database paths that contain track(s)</i> |
|------------|---|

Description

Returns all database paths that contain a version of a track.

Usage

```
gtrack.dbs(track = NULL, dataframe = FALSE)
```

Arguments

| | |
|-----------|--|
| track | track name or a vector of track names |
| dataframe | return a data frame with columns track and db instead of a named character vector. |

Details

When datasets are loaded, a track may exist in multiple locations (working database and/or datasets). This function computes on-demand and returns all such paths, which is useful for debugging when using `force=TRUE` with `gdataset.load()`.

Value

A named character vector of database paths for each track. If `dataframe` is `TRUE`, returns a data frame with columns `track` and `db`, with multiple rows per track when it appears in multiple databases.

See Also

[gtrack.dataset](#), [gtrack.exists](#), [gtrack.ls](#), [gdataset.ls](#)

Examples

```
gdb.init_examples()
gtrack.dbs("dense_track")
gtrack.dbs(gtrack.ls(), dataframe = TRUE)
```

| | |
|---------------|------------------------------------|
| gtrack.exists | <i>Tests for a track existence</i> |
|---------------|------------------------------------|

Description

Tests for a track existence.

Usage

```
gtrack.exists(track = NULL)
```

Arguments

| | |
|-------|------------|
| track | track name |
|-------|------------|

Details

This function returns 'TRUE' if a track exists in Genomic Database.

Value

'TRUE' if a track exists. Otherwise 'FALSE'.

See Also

[gtrack.ls](#), [gtrack.info](#), [gtrack.create](#), [gtrack.rm](#)

Examples

```
gdb.init_examples()  
gtrack.exists("dense_track")
```

| | |
|------------------------|--|
| gtrack.export_bedgraph | <i>Export a track to bedGraph format</i> |
|------------------------|--|

Description

Exports a track or track expression to a UCSC bedGraph file.

Usage

```
gtrack.export_bedgraph(
  track,
  file,
  intervals = NULL,
  iterator = NULL,
  name = NULL
)
```

Arguments

| | |
|-----------|---|
| track | track name or track expression (character string) |
| file | output file path. If it ends in .gz, output is gzip-compressed. |
| intervals | genomic intervals to export. If NULL (default), the entire genome (.misha\$ALLGENOME) is used. |
| iterator | iterator bin size. If NULL (default), the iterator is determined automatically from the track expression. |
| name | track name for the bedGraph header line. If NULL (default), uses the track parameter value. |

Details

This function evaluates a track expression over the specified genomic intervals and writes the result in standard bedGraph format (4-column, tab-separated: chrom, start, end, value). NaN values are omitted from the output.

The function supports physical tracks, virtual tracks, and arbitrary track expressions (e.g. "dense_track * 2"). 2D tracks are not supported.

If the output file path ends in .gz, the output is gzip-compressed.

Value

NULL (invisible). Called for its side effect of writing a file.

See Also

[gextract](#), [gtrack.export_bigwig](#), [gtrack.info](#)

Examples

```
## Not run:
gdb.init_examples()

# Export a dense track
gtrack.export_bedgraph("dense_track", "/tmp/dense.bedgraph")

# Export with specific intervals
intervs <- gintervals(1, 0, 1000)
gtrack.export_bedgraph("dense_track", "/tmp/dense_chr1.bedgraph",
```

```
    intervals = intervals
)

# Export a track expression
gtrack.export_bedgraph("dense_track * 2", "/tmp/scaled.bedgraph",
    iterator = 100
)

# Export compressed
gtrack.export_bedgraph("dense_track", "/tmp/dense.bedgraph.gz")

## End(Not run)
```

`gtrack.export_bigwig` *Export a track to BigWig format*

Description

Exports a track or track expression to BigWig format by first creating a temporary bedGraph file and then converting it using bedGraphToBigWig (or wigToBigWig as a fallback).

Usage

```
gtrack.export_bigwig(track, file, intervals = NULL, iterator = NULL)
```

Arguments

| | |
|------------------------|---|
| <code>track</code> | track name or track expression (character string) |
| <code>file</code> | output file path (typically ending in <code>.bw</code> or <code>.bigwig</code>). |
| <code>intervals</code> | genomic intervals to export. If NULL (default), the entire genome (<code>.misha\$ALLGENOME</code>) is used. |
| <code>iterator</code> | iterator bin size. If NULL (default), the iterator is determined automatically from the track expression. |

Details

This function requires the UCSC bedGraphToBigWig utility to be installed and available on the system PATH, or bundled with the misha package. If not found, the function will raise an error with installation instructions.

Value

NULL (invisible). Called for its side effect of writing a file.

See Also

[gextract](#), [gtrack.export_bedgraph](#), [gtrack.info](#)

Examples

```

## Not run:
gdb.init_examples()

# Export to BigWig (requires bedGraphToBigWig)
gtrack.export_bigwig("dense_track", "/tmp/dense.bw")

# With specific region
gtrack.export_bigwig("dense_track", "/tmp/dense_chr1.bw",
  intervals = gintervals(1, 0, 1e6)
)

## End(Not run)

```

| | |
|---------------|--|
| gtrack.import | <i>Creates a track from WIG / BigWig / BedGraph / BED / tab-delimited file</i> |
|---------------|--|

Description

Creates a track from WIG / BigWig / BedGraph / BED / tab-delimited file

Usage

```

gtrack.import(
  track = NULL,
  description = NULL,
  file = NULL,
  binsize = NULL,
  defval = NaN,
  attrs = NULL
)

```

Arguments

| | |
|-------------|--|
| track | track name |
| description | a character string description |
| file | file path |
| binsize | bin size of the newly created 'Dense' track or '0' for a 'Sparse' track |
| defval | default track value |
| attrs | a named vector or list of attributes to be set on the track after import |

Details

This function creates a track from WIG / BigWig / BedGraph / tab-delimited file. Zipped files are supported (file name must have '.gz' or '.zip' suffix).

Tab-delimited files must start with a header line with the following column names (tab-separated): 'chrom', 'start', 'end', and exactly one value column name (e.g. 'value'). Each subsequent line provides a single interval: - chrom: chromosome name (e.g. 'chr1') - start: 0-based start coordinate (inclusive) - end: 0-based end coordinate (exclusive) - value: numeric value (floating point allowed); exactly one value column is supported

Columns must be separated by tabs. Coordinates must refer to chromosomes existing in the current genome. Missing values can be specified as 'NaN'.

BED files (.bed/.bed.gz/.bed.zip) are also supported. If the BED 'score' column (5th column) exists and is numeric, it is used as the interval value; otherwise a constant value of 1 is used. For BED inputs, 'binsize' controls the output type: if 'binsize' is 0 the track is 'Sparse'; otherwise the track is 'Dense' with bin-averaged values based on overlaps with BED intervals (and 'defval' for regions not covered).

If 'binsize' is 0 the resulted track is created in 'Sparse' format. Otherwise the 'Dense' format is chosen with a bin size equal to 'binsize'. The values that were not defined in input file file are substituted by 'defval' value.

'description' is added as a track attribute.

When multiple databases are connected via [gsetroot](#), the track is created in the current working directory (.misha\$GWD), which defaults to the last connected database. Use [gdir.cd](#) with an absolute path to change where new tracks are created.

Value

None.

See Also

[gtrack.import_set](#), [gtrack.rm](#), [gtrack.info](#), [gdir.create](#), [gextract](#)

Examples

```
gdb.init_examples()

# Create a simple WIG file for demonstration
temp_file <- tempfile(fileext = ".wig")
writeLines(c(
  "track type=wiggle_0 name=\"example track\"",
  "fixedStep chrom=chr1 start=1 step=1",
  "1.5",
  "2.0",
  "1.8",
  "3.2"
), temp_file)
```

```
# Basic import
gtrack.import("example_track", "Example track from WIG file",
             temp_file,
             binsize = 1
            )
gtrack.info("example_track")
gtrack.rm("example_track", force = TRUE)

# Import with custom attributes
attrs <- c("author" = "researcher", "version" = "1.0", "experiment" = "test")
gtrack.import("example_track_with_attrs", "Example track with attributes",
             temp_file,
             binsize = 1, attrs = attrs
            )

# Check that attributes were set
gtrack.attr.get("example_track_with_attrs", "author")
gtrack.attr.get("example_track_with_attrs", "version")
gtrack.attr.get("example_track_with_attrs", "experiment")

# Clean up
gtrack.rm("example_track_with_attrs", force = TRUE)
```

```
gtrack.import_mappedseq
```

Creates a track from a file of mapped sequences

Description

Creates a track from a file of mapped sequences.

Usage

```
gtrack.import_mappedseq(
  track = NULL,
  description = NULL,
  file = NULL,
  pileup = 0,
  binsize = -1,
  cols.order = c(9, 11, 13, 14),
  remove.dups = TRUE
)
```

Arguments

| | |
|-------------|--------------------------------|
| track | track name |
| description | a character string description |

| | |
|-------------|---|
| file | name of mapped sequences file |
| pileup | interval expansion |
| binsize | bin size of a dense track |
| cols.order | order of sequence, chromosome, coordinate and strand columns in mapped sequences file or NULL if SAM file is used |
| remove.dups | if 'TRUE' the duplicated coordinates are counted only once. |

Details

This function creates a track from a file of mapped sequences. The file can be in SAM format or in a general TAB delimited text format where each line describes a single read.

For a SAM file 'cols.order' must be set to 'NULL'.

For a general TAB delimited text format the following columns must be presented in the file: sequence, chromosome, coordinate and strand. The position of these columns should be specified in 'cols.order' argument. The default value of 'cols.order' is an array of (9, 11, 13, 14) meaning that sequence is expected to be found at column number 9, chromosome - at column 11, coordinate - at column 13 and strand - at column 14. The column indices are 1-based, i.e. the first column is referenced by 1. Chromosome needs a prefix 'chr' e.g. 'chr1'. Valid strand values are '+' or 'F' for forward strand and '-' or 'R' for the reverse strand.

Each read at given coordinate can be "expanded" to cover an interval rather than a single point. The length of the interval is controlled by 'pileup' argument. The direction of expansion depends on the strand value. If 'pileup' is '0', no expansion is performed and the read is converted to a single point. The track is created in sparse format. If 'pileup' is greater than zero, the output track is in dense format. 'binsize' controls the bin size of the dense track.

If 'remove.dups' is 'TRUE' the duplicated coordinates are counted only once.

'description' is added as a track attribute.

'gtrack.import_mappedseq' returns the statistics of the conversion process.

Value

A list of conversion process statistics.

See Also

[gtrack.rm](#), [gtrack.info](#), [gdir.create](#)

| | |
|-------------------|--|
| gtrack.import_set | <i>Creates one or more tracks from multiple WIG / BigWig / BedGraph / tab-delimited files on disk or FTP</i> |
|-------------------|--|

Description

Creates one or more tracks from WIG / BigWig / BedGraph / tab-delimited files on disk or FTP.

Usage

```
gtrack.import_set(
    description = NULL,
    path = NULL,
    binsize = NULL,
    track.prefix = NULL,
    defval = NaN
)
```

Arguments

| | |
|--------------|---|
| description | a character string description |
| path | file path or URL (may contain wildcards) |
| binsize | bin size of the newly created 'Dense' track or '0' for a 'Sparse' track |
| track.prefix | prefix for a track name |
| defval | default track value |

Details

This function is similar to 'gtrack.import' however unlike the latter it can create multiple tracks. Additionally the files can be fetched from an FTP server.

The files are expected to be in WIG / BigWig / BedGraph / tab-delimited formats. One can learn about the format of the tab-delimited file by running 'gextract' function with a 'file' parameter set to the name of the file. Zipped files are supported (file name must have '.gz' or '.zip' suffix).

Files are specified by 'path' argument. 'path' can be also a URL of an FTP server in the form of 'ftp://[address]/[files]'. If 'path' is a URL, the files are first downloaded from FTP server to a temporary directory and then imported to tracks. The temporary directory is created at 'GROOT/downloads'.

Regardless whether 'path' is file path or to a URL, it can contain wildcards. Hence multiple files can be imported (and downloaded) at once.

If 'binsize' is 0 the resulted tracks are created in 'Sparse' format. Otherwise the 'Dense' format is chosen with a bin size equal to 'binsize'. The values that were not defined in input file are substituted by 'defval' value.

The name of a each created track is of '[track.prefix][filename]' form, where 'filename' is the name of the WIG file. For example, if 'track.prefix' equals to "wigs." and an input file name is 'mydata', a track named 'wigs.mydata' is created. If 'track.prefix' is 'NULL' no prefix is appended to the name of the created track.

Existing tracks are not overwritten and no new directories are automatically created.

'description' is added to the created tracks as an attribute.

'gtrack.import_set' does not stop if an error occurs while importing a file. It rather continues importing the rest of the files.

'gtrack.import_set' returns the names of the files that were successfully imported and those that failed.

Value

Names of files that were successfully imported and those that failed.

See Also

[gtrack.import](#), [gwget](#), [gtrack.rm](#), [gtrack.info](#), [gdir.create](#), [gextract](#)

| | |
|--------------------------|--|
| <code>gtrack.info</code> | <i>Returns information about a track</i> |
|--------------------------|--|

Description

Returns information about a track.

Usage

```
gtrack.info(track = NULL, validate = FALSE)
```

Arguments

| | |
|-----------------------|--|
| <code>track</code> | track name |
| <code>validate</code> | if TRUE, validates the track index file integrity (for indexed tracks). Default: FALSE |

Details

Returns information about the track (type, dimensions, size in bytes, etc.). The fields in the returned value vary depending on the type of the track.

Value

A list that contains track properties

See Also

[gtrack.exists](#), [gtrack.ls](#)

Examples

```
gdb.init_examples()  
gtrack.info("dense_track")  
gtrack.info("rects_track")
```

`gtrack.liftover` *Imports a track from another assembly*

Description

Imports a track from another assembly.

Usage

```
gtrack.liftover(
  track = NULL,
  description = NULL,
  src.track.dir = NULL,
  chain = NULL,
  src_overlap_policy = "error",
  tgt_overlap_policy = "auto",
  multi_target_agg = c("mean", "median", "sum", "min", "max", "count", "first", "last",
    "nth", "max.coverage_len", "min.coverage_len", "max.coverage_frac",
    "min.coverage_frac"),
  params = NULL,
  na.rm = TRUE,
  min_n = NULL,
  min_score = NULL
)
```

Arguments

| | |
|---------------------------------|--|
| <code>track</code> | name of a created track |
| <code>description</code> | a character string description |
| <code>src.track.dir</code> | path to the directory of the source track |
| <code>chain</code> | name of chain file or data frame as returned by <code>'gintervals.load_chain'</code> |
| <code>src_overlap_policy</code> | policy for handling source overlaps: "error" (default), "keep", or "discard". "keep" allows one source interval to map to multiple target intervals, "discard" discards all source intervals that have overlaps and "error" throws an error if source overlaps are detected. |
| <code>tgt_overlap_policy</code> | policy for handling target overlaps. One of: |

| Policy | Description |
|--------------------------|---|
| <code>error</code> | Throws an error if any target overlaps are detected. |
| <code>auto</code> | Default. Alias for "auto_score". |
| <code>auto_score</code> | Resolves overlaps by segmenting the target region and selecting the best chain for each segment based on score. |
| <code>auto_longer</code> | Resolves overlaps by segmenting and selecting the chain with the longest span for each segment. Tie-breaker is score. |
| <code>auto_first</code> | Resolves overlaps by segmenting and selecting the chain with the lowest chain_id for each segment. |

| | |
|---------------------|---|
| keep | Preserves all overlapping intervals. |
| discard | Discards any chain interval that has a target overlap with another chain interval. |
| agg | Segments overlaps into smaller disjoint regions where each region contains all contributing chains, allowing for overlapping regions. |
| best_source_cluster | Best source cluster strategy based on source overlap. When multiple chains map a source interval, cluster the best one. |

multi_target_agg

| | |
|-----------|--|
| | aggregation/selection policy for contributors that land on the same target locus. When multiple source intervals map to overlapping regions in the target genome (after applying tgt_overlap_policy), their values must be combined into a single value. |
| params | additional parameters for aggregation (e.g., for "nth" aggregation) |
| na.rm | logical indicating whether NA values should be removed before aggregation (default: TRUE) |
| min_n | minimum number of non-NA values required for aggregation. If fewer values are available, the result will be NA. |
| min_score | optional minimum alignment score threshold. Chains with scores below this value are filtered out. Useful for excluding low-quality alignments. |

Details

This function imports a track located in 'src.track.dir' of another assembly to the current database. Chain file instructs how the conversion of coordinates should be done. It can be either a name of a chain file or a data frame in the same format as returned by 'gintervals.load_chain' function. The name of the newly created track is specified by 'track' argument and 'description' is added as a track attribute.

Note: When passing a pre-loaded chain (data frame), overlap policies cannot be specified - they are taken from the chain's attributes that were set during loading. When passing a chain file path, policies can be specified and will be used for loading. Aggregation parameters (multi_target_agg, params, na.rm, min_n) can always be specified regardless of chain type.

Value

None.

Note

Terminology note for UCSC chain format users: In the UCSC chain format specification, the fields prefixed with 't' (tName, tStart, tEnd, etc.) are called "target" or "reference", while fields prefixed with 'q' (qName, qStart, qEnd, etc.) are called "query". However, misha uses reversed terminology: UCSC's "target/reference" corresponds to misha's "source" (chromsrc, startsrc, endsrc), and UCSC's "query" corresponds to misha's "target" (chrom, start, end).

See Also

[gintervals.load_chain](#), [gintervals.liftover](#)

| | |
|----------------------------|--|
| <code>gtrack.lookup</code> | <i>Creates a new track from a lookup table based on track expression</i> |
|----------------------------|--|

Description

Evaluates track expression and translates the values into bin indices that are used in turn to retrieve values from a lookup table and create a track.

Usage

```
gtrack.lookup(
    track = NULL,
    description = NULL,
    lookup_table = NULL,
    ...,
    include.lowest = FALSE,
    force.binning = TRUE,
    iterator = NULL,
    band = NULL
)
```

Arguments

| | |
|-----------------------------|---|
| <code>track</code> | track name |
| <code>description</code> | a character string description |
| <code>lookup_table</code> | a multi-dimensional array containing the values that are returned by the function |
| <code>...</code> | pairs of track expressions and breaks |
| <code>include.lowest</code> | if 'TRUE', the lowest value of the range determined by breaks is included |
| <code>force.binning</code> | if 'TRUE', the values smaller than the minimal break will be translated to index 1, and the values that exceed the maximal break will be translated to index N-1 where N is the number of breaks. If 'FALSE' the out-of-range values will produce NaN values. |
| <code>iterator</code> | track expression iterator. If 'NULL' iterator is determined implicitly based on track expressions. |
| <code>band</code> | track expression band. If 'NULL' no band is used. |

Details

This function evaluates the track expression for all iterator intervals and translates this value into an index based on the breaks. This index is then used to address the lookup table and create with its values a new track. More than one 'expr'-'breaks' pair can be used. In that case 'lookup_table' is addressed in a multidimensional manner, i.e. 'lookup_table[i1, i2, ...]'.

The range of bins is determined by 'breaks' argument. For example: 'breaks = c(x1, x2, x3, x4)' represents three different intervals (bins): (x1, x2], (x2, x3], (x3, x4].

If 'include.lowest' is 'TRUE' the the lowest value is included in the first interval, i.e. in [x1, x2].

'force.binning' parameter controls what should be done when the value of 'expr' exceeds the range determined by 'breaks'. If 'force.binning' is 'TRUE' then values smaller than the minimal break will be translated to index 1, and the values exceeding the maximal break will be translated to index 'M-1' where 'M' is the number of breaks. If 'force.binning' is 'FALSE' the out-of-range values will produce 'NaN' values.

Regardless of 'force.binning' value if the value of 'expr' is 'NaN' then the value in the track would be 'NaN' too.

'description' is added as a track attribute.

Value

None.

See Also

[glookup](#), [gtrack.2d.create](#), [gtrack.create_sparse](#), [gtrack.smooth](#), [gtrack.modify](#), [gtrack.rm](#), [gtrack.info](#), [gdir.create](#)

Examples

```
gdb.init_examples()

## one-dimensional example
breaks1 <- seq(0.1, 0.2, length.out = 6)
gtrack.lookup(
  "lookup_track", "Test track", 1:5, "dense_track",
  breaks1
)
gtrack.rm("lookup_track", force = TRUE)

## two-dimensional example
t <- array(1:15, dim = c(5, 3))
breaks2 <- seq(0.31, 0.37, length.out = 4)
gtrack.lookup(
  "lookup_track", "Test track", t, "dense_track",
  breaks1, "2 * dense_track", breaks2
)
gtrack.rm("lookup_track", force = TRUE)
```

gtrack.ls

Returns a list of track names

Description

Returns a list of track names in Genomic Database.

Usage

```
gtrack.ls(
  ...,
  db = NULL,
  ignore.case = FALSE,
  perl = FALSE,
  fixed = FALSE,
  useBytes = FALSE
)
```

Arguments

... these arguments are of either form 'pattern' or 'attribute = pattern'

db optional database path to filter tracks. If specified, only tracks from that database are returned.

ignore.case, perl, fixed, useBytes see 'grep'

Details

This function returns a list of tracks whose name or track attribute value match a pattern (see 'grep'). If called without any arguments all tracks are returned.

If pattern is specified without a track attribute (i.e. in the form of 'pattern') then filtering is applied to the track names. If pattern is supplied with a track attribute (i.e. in the form of 'name = pattern') then track attribute is matched against the pattern.

Multiple patterns are applied one after another. The resulted list of tracks should match all the patterns.

When multiple databases are connected, the 'db' parameter can be used to filter tracks to only those from a specific database.

Value

An array that contains the names of tracks that match the supplied patterns.

See Also

[grep](#), [gtrack.exists](#), [gtrack.create](#), [gtrack.rm](#), [gtrack.dataset](#)

Examples

```
gdb.init_examples()

# get all track names
gtrack.ls()

# get track names that match the pattern "den*"
gtrack.ls("den*")
```

```
# get track names whose "created.by" attribute match the pattern
# "create_sparse"
gtrack.ls(created.by = "create_sparse")

# get track names whose names match the pattern "den*" and whose
# "created.by" attribute match the pattern "track"
gtrack.ls("den*", created.by = "track")
```

gtrack.modify

Modifies track contents

Description

Modifies 'Dense' track contents.

Usage

```
gtrack.modify(track = NULL, expr = NULL, intervals = NULL)
```

Arguments

| | |
|-----------|---|
| track | track name |
| expr | track expression |
| intervals | genomic scope for which track is modified |

Details

This function modifies the contents of a 'Dense' track by the values of 'expr'. 'intervals' argument controls which portion of the track is modified. The iterator policy is set internally to the bin size of the track.

Value

None.

See Also

[gtrack.create](#), [gtrack.rm](#)

Examples

```
gdb.init_examples()
intervs <- gintervals(1, 300, 800)
gextract("dense_track", intervs)
gtrack.modify("dense_track", "dense_track * 2", intervs)
gextract("dense_track", intervs)
gtrack.modify("dense_track", "dense_track / 2", intervs)
```

gtrack.mv

Renames or moves a track

Description

Renames a track or moves it to a different namespace within the same database.

Usage

```
gtrack.mv(src = NULL, dest = NULL)
```

Arguments

| | |
|------|------------------------|
| src | source track name |
| dest | destination track name |

Details

This function renames a track or moves it to a different namespace (directory) within the same database. The track cannot be moved to a different database. Use [gtrack.copy](#) followed by [gtrack.rm](#) if you need to move a track between databases.

Value

None.

See Also

[gtrack.copy](#), [gtrack.rm](#), [gtrack.exists](#), [gtrack.ls](#)

Examples

```
gdb.init_examples()
gtrack.create_sparse("test_track", "Test", gintervals(1, 0, 100), 1)
gtrack.mv("test_track", "renamed_track")
gtrack.exists("renamed_track")
gtrack.rm("renamed_track", force = TRUE)
```

| | |
|-------------|-------------------------------------|
| gtrack.path | Returns the path on disk of a track |
|-------------|-------------------------------------|

Description

Returns the path on disk of a track.

Usage

```
gtrack.path(track = NULL)
```

Arguments

track track name or a vector of track names

Details

This function returns the actual file system path where a track is stored. The function works with a single track name or a vector of track names.

Value

A character vector containing the full paths to the tracks on disk.

See Also

[gtrack.exists](#), [gtrack.ls](#), [gintervals.path](#)

Examples

```
gdb.init_examples()
gtrack.path("dense_track")
gtrack.path(c("dense_track", "sparse_track"))
```

| | |
|-----------|-----------------|
| gtrack.rm | Deletes a track |
|-----------|-----------------|

Description

Deletes a track.

Usage

```
gtrack.rm(track = NULL, force = FALSE, db = NULL)
```

Arguments

| | |
|-------|---|
| track | track name |
| force | if 'TRUE', suppresses user confirmation of a named track removal |
| db | optional database path to delete the track from when multiple databases are connected |

Details

This function deletes a track from the Genomic Database. By default 'gtrack.rm' requires the user to interactively confirm the deletion. Set 'force' to 'TRUE' to suppress the user prompt.

Value

None.

See Also

[gtrack.exists](#), [gtrack.ls](#), [gtrack.create](#), [gtrack.2d.create](#), [gtrack.create_sparse](#), [gtrack.smooth](#)

Examples

```
gdb.init_examples()
gtrack.create("new_track", "Test track", "2 * dense_track")
gtrack.exists("new_track")
gtrack.rm("new_track", force = TRUE)
gtrack.exists("new_track")
```

`gtrack.smooth`

Creates a new track from smoothed values of track expression

Description

Creates a new track from smoothed values of track expression.

Usage

```
gtrack.smooth(
  track = NULL,
  description = NULL,
  expr = NULL,
  winsize = NULL,
  weight_thr = 0,
  smooth_nans = FALSE,
  alg = "LINEAR_RAMP",
  iterator = NULL
)
```

Arguments

| | |
|-------------|--|
| track | track name |
| description | a character string description |
| expr | track expression |
| winsize | size of smoothing window |
| weight_thr | smoothing weight threshold |
| smooth_nans | if 'FALSE' track value is always set to 'NaN' if central window value is 'NaN', otherwise it is calculated from the rest of non 'NaN' values |
| alg | smoothing algorithm - "MEAN" or "LINEAR_RAMP" |
| iterator | track expression iterator of 'Fixed bin' type |

Details

This function creates a new 'Dense' track named 'track'. The values of the track are results of smoothing the values of 'expr'.

Each track value at coordinate 'C' is determined by smoothing non 'NaN' values of 'expr' over the window around 'C'. The window size is controlled by 'winsize' and is given in coordinate units (not in number of bins), defining the total regions to be considered when smoothing (on both sides of the central point). Two different algorithms can be used for smoothing:

"MEAN" - an arithmetic average.

"LINEAR_RAMP" - a weighted arithmetic average, where the weights linearly decrease as the distance from the center of the window increases.

'weight_thr' determines the function behavior when some of the values in the window are missing or 'NaN' (missing values may occur at the edges of each chromosome when the window covers an area beyond chromosome boundaries). 'weight_thr' sets the weight sum threshold below which smoothing algorithm returns 'NaN' rather than a smoothing value based on non 'NaN' values in the window.

'smooth_nans' controls what would be the smoothed value if the central value in the window is 'NaN'. If 'smooth_nans' is 'FALSE' then the smoothed value is set to 'NaN' regardless of 'weight_thr' parameter. Otherwise it is calculated normally.

'description' is added as a track attribute.

Iterator policy must be of "fixed bin" type.

Value

None.

See Also

[gtrack.create](#), [gtrack.2d.create](#), [gtrack.create_sparse](#), [gtrack.modify](#), [gtrack.rm](#), [gtrack.info](#), [gdir.create](#)

Examples

```
gdb.init_examples()
gtrack.smooth("smoothed_track", "Test track", "dense_track", 500)
gextract("dense_track", "smoothed_track", gintervals(1, 0, 1000))
gtrack.rm("smoothed_track", force = TRUE)
```

| | |
|----------------|--|
| gtrack.var.get | <i>Returns value of a track variable</i> |
|----------------|--|

Description

Returns value of a track variable.

Usage

```
gtrack.var.get(track = NULL, var = NULL)
```

Arguments

| | |
|-------|---------------------|
| track | track name |
| var | track variable name |

Details

This function returns the value of a track variable. If the variable does not exist an error is reported.

Value

Track variable value.

See Also

[gtrack.var.set](#), [gtrack.var.ls](#), [gtrack.var.rm](#)

Examples

```
gdb.init_examples()
gtrack.var.set("sparse_track", "test_var", 1:10)
gtrack.var.get("sparse_track", "test_var")
gtrack.var.rm("sparse_track", "test_var")
```

| | |
|---------------|---|
| gtrack.var.ls | Returns a list of track variables for a track |
|---------------|---|

Description

Returns a list of track variables for a track.

Usage

```
gtrack.var.ls(  
    track = NULL,  
    pattern = "",  
    ignore.case = FALSE,  
    perl = FALSE,  
    fixed = FALSE,  
    useBytes = FALSE  
)
```

Arguments

track track name
pattern, ignore.case, perl, fixed, useBytes
 see 'grep'

Details

This function returns a list of track variables of a track that match the pattern (see 'grep'). If called without any arguments all track variables of a track are returned.

Value

An array that contains the names of track variables.

See Also

[grep](#), [gtrack.var.get](#), [gtrack.var.set](#), [gtrack.var.rm](#)

Examples

```
gdb.init_examples()  
gtrack.var.ls("sparse_track")  
gtrack.var.set("sparse_track", "test_var1", 1:10)  
gtrack.var.set("sparse_track", "test_var2", "v")  
gtrack.var.ls("sparse_track")  
gtrack.var.ls("sparse_track", pattern = "2")  
gtrack.var.rm("sparse_track", "test_var1")  
gtrack.var.rm("sparse_track", "test_var2")
```

| | |
|---------------|---------------------------------|
| gtrack.var.rm | <i>Deletes a track variable</i> |
|---------------|---------------------------------|

Description

Deletes a track variable.

Usage

```
gtrack.var.rm(track = NULL, var = NULL)
```

Arguments

| | |
|-------|---------------------|
| track | track name |
| var | track variable name |

Details

This function deletes a track variable.

Value

None.

See Also

[gtrack.var.get](#), [gtrack.var.set](#), [gtrack.var.ls](#)

Examples

```
gdb.init_examples()
gtrack.var.set("sparse_track", "test_var1", 1:10)
gtrack.var.set("sparse_track", "test_var2", "v")
gtrack.var.ls("sparse_track")
gtrack.var.rm("sparse_track", "test_var1")
gtrack.var.rm("sparse_track", "test_var2")
gtrack.var.ls("sparse_track")
```

| | |
|----------------|--|
| gtrack.var.set | <i>Assigns value to a track variable</i> |
|----------------|--|

Description

Assigns value to a track variable.

Usage

```
gtrack.var.set(track = NULL, var = NULL, value = NULL)
```

Arguments

| | |
|-------|---------------------|
| track | track name |
| var | track variable name |
| value | value |

Details

This function creates a track variable and assigns 'value' to it. If the track variable already exists its value is overwritten.

Value

None.

See Also

[gtrack.var.get](#), [gtrack.var.ls](#), [gtrack.var.rm](#)

Examples

```
gdb.init_examples()  
gtrack.var.set("sparse_track", "test_var", 1:10)  
gtrack.var.get("sparse_track", "test_var")  
gtrack.var.rm("sparse_track", "test_var")
```

gvtrack.array.slice *Defines rules for a single value calculation of a virtual 'Array' track*

Description

Defines how a single value within an interval is achieved for a virtual track based on 'Array' track.

Usage

```
gvtrack.array.slice(vtrack = NULL, slice = NULL, func = "avg", params = NULL)
```

Arguments

| | |
|--------------|--|
| vtrack | virtual track name |
| slice | a vector of column names or column indices or 'NULL' |
| func, params | see below |

Details

A track (regular or virtual) used in a track expression is expected to return one value for each track interval. 'Array' tracks store multiple values per interval (one for each 'column') and hence if used in a track expression one must define the way of how a single value should be deduced from several ones.

By default if an 'Array' track is used in a track expressions, its interval value would be the average of all column values that are not NaN. 'gvtrack.array.slice' allows to select specific columns and to specify the function applied to their values.

'slice' parameter allows to choose the columns. Columns can be indicated by their names or their indices. If 'slice' is 'NULL' the non-NaN values of all track columns are used.

'func' parameter determines the function applied to the columns' values. Use the following table for a reference of all valid functions and parameters combinations:

func = "avg", params = NULL

Average of columns' values.

func = "max", params = NULL

Maximum of columns' values.

func = "min", params = NULL

Minimum of columns' values.

func = "stdev", params = NULL

Unbiased standard deviation of columns' values.

func = "sum", params = NULL

Sum of columns' values.

func = "quantile", params = [Percentile in the range of [0, 1]]

Quantile of columns' values.

Value

None.

See Also[gvtrack.create](#), [gtrack.array.get_colnames](#), [gtrack.array.extract](#)**Examples**

```

gdb.init_examples()
gvtrack.create("vtrack1", "array_track")
gvtrack.array.slice("vtrack1", c("col2", "col4"), "max")
gextract("vtrack1", gintervals(1, 0, 1000))

```

| | |
|----------------|------------------------------------|
| gvtrack.create | <i>Creates a new virtual track</i> |
|----------------|------------------------------------|

Description

Creates a new virtual track.

Usage

```

gvtrack.create(
  vtrack = NULL,
  src = NULL,
  func = NULL,
  params = NULL,
  dim = NULL,
  sshift = NULL,
  eshift = NULL,
  filter = NULL,
  ...
)

```

Arguments

| | |
|--------|---|
| vtrack | virtual track name |
| src | source (track/intervals). NULL for PWM functions. For value-based tracks, provide a data frame with columns chrom, start, end, and one numeric value column. The data frame functions as an in-memory sparse track and supports all track-based summarizer functions. Intervals must not overlap. |
| func | function name (see above) |
| params | function parameters (see above) |

| | |
|--------|--|
| dim | use 'NULL' or '0' for 1D iterators. '1' converts 2D iterator to (chrom1, start1, end1), '2' converts 2D iterator to (chrom2, start2, end2) |
| sshift | shift of 'start' coordinate |
| eshift | shift of 'end' coordinate |
| filter | genomic mask to apply. Can be: <ul style="list-style-type: none"> • A data.frame with columns 'chrom', 'start', 'end' (intervals to mask) • A character string naming an intervals set • A character string naming a track (must be intervals-type track) • A list of any combination of the above (all will be unified) • NULL to clear the filter |
| ... | additional PWM parameters |

Details

This function creates a new virtual track named 'vtrack' with the given source, function and parameters. 'src' can be either a track, intervals (1D or 2D), or a data frame with intervals and a numeric value column (value-based track). The tables below summarize the supported combinations.

Value-based tracks Value-based tracks are data frames containing genomic intervals with associated numeric values. They function as in-memory sparse tracks without requiring track creation in the database. To create a value-based track, provide a data frame with columns chrom, start, end, and one numeric value column (any name is acceptable). Value-based tracks support all track-based summarizer functions (e.g., avg, min, max, sum, lse, stddev, quantile, nearest, exists, size, first, last, sample, and position functions), but do not support overlapping intervals. They behave like sparse tracks in aggregation: values are aggregated using count-based averaging (each interval contributes equally regardless of length), not coverage-based averaging.

Track-based summarizers

| Source | func | params | Description |
|------------------------------|-----------------------|----------------------|--|
| Track | avg | NULL | Average track value in the iterator interval. |
| Track (1D) | exists | vals (optional) | Returns 1 if any value exists (or specific vals if provided) |
| Track (1D) | first | NULL | First value in the iterator interval. |
| Track (1D) | last | NULL | Last value in the iterator interval. |
| Track | max | NULL | Maximum track value in the iterator interval. |
| Track | min | NULL | Minimum track value in the iterator interval. |
| Dense / Sparse / Array track | nearest | NULL | Average value inside the iterator; for sparse tracks returns nearest value |
| Track (1D) | sample | NULL | Uniformly sampled source value from the iterator interval |
| Track (1D) | size | NULL | Number of non-NaN values in the iterator interval |
| Dense / Sparse / Array track | stddev | NULL | Unbiased standard deviation of values in the iterator interval |
| Dense / Sparse / Array track | sum | NULL | Sum of values in the iterator interval. |
| Dense / Sparse / Array track | lse | NULL | Log-sum-exp of track values in the iterator interval |
| Dense / Sparse / Array track | quantile | Percentile in [0, 1] | Quantile of values in the iterator interval. |
| Dense track | global.percentile | NULL | Percentile of the interval average relative to the full set of values |
| Dense track | global.percentile.max | NULL | Percentile of the interval maximum relative to the full set of values |
| Dense track | global.percentile.min | NULL | Percentile of the interval minimum relative to the full set of values |

Track position summarizers

| Source | func | params | Description |
|------------|---------------------|--------|--|
| Track (1D) | first.pos.abs | NULL | Absolute genomic coordinate of the first value. |
| Track (1D) | first.pos.relative | NULL | Zero-based position (relative to interval start) of the first value. |
| Track (1D) | last.pos.abs | NULL | Absolute genomic coordinate of the last value. |
| Track (1D) | last.pos.relative | NULL | Zero-based position (relative to interval start) of the last value. |
| Track (1D) | max.pos.abs | NULL | Absolute genomic coordinate of the maximum value inside the iterator interval. |
| Track (1D) | max.pos.relative | NULL | Zero-based position (relative to interval start) of the maximum value. |
| Track (1D) | min.pos.abs | NULL | Absolute genomic coordinate of the minimum value inside the iterator interval. |
| Track (1D) | min.pos.relative | NULL | Zero-based position (relative to interval start) of the minimum value. |
| Track (1D) | sample.pos.abs | NULL | Absolute genomic coordinate of a uniformly sampled value. |
| Track (1D) | sample.pos.relative | NULL | Zero-based position (relative to interval start) of a uniformly sampled value. |

For `max.pos.relative`, `min.pos.relative`, `first.pos.relative`, `last.pos.relative`, `sample.pos.relative`, iterator modifiers (including `sshift` / `eshift` and 1D projections generated via `gvtrack.iterator`) are applied before the position is reported. In other words, the returned coordinate is always 0-based and measured from the start of the iterator interval after all modifier adjustments.

Interval-based summarizers

| Source | func | params | Description |
|--------------|-----------------|--|---|
| 1D intervals | distance | Minimal distance from center (default 0) | Signed distance using normalized formula when i |
| 1D intervals | distance.center | NULL | Distance from iterator center to the closest interval |
| 1D intervals | distance.edge | NULL | Edge-to-edge distance from iterator interval to closest |
| 1D intervals | coverage | NULL | Fraction of iterator length covered by source intervals |
| 1D intervals | neighbor.count | Max distance (≥ 0) | Number of source intervals whose edge-to-edge d |

2D track summarizers

| Source | func | params | Description |
|----------|--------------|--------|---|
| 2D track | area | NULL | Area covered by intersections of track rectangles with the iterator interval. |
| 2D track | weighted.sum | NULL | Weighted sum of values where each weight equals the intersection area. |

Motif (PWM) summarizers

| Source | func | Key params | Description |
|-----------------|-------------|---|--------------------------------|
| NULL (sequence) | pwm | pssm, bidirect, prior, extend, spat_* | Log-sum-exp score of motif |
| NULL (sequence) | pwm.max | pssm, bidirect, prior, extend, spat_* | Maximum log-likelihood score |
| NULL (sequence) | pwm.max.pos | pssm, bidirect, prior, extend, spat_* | 1-based position of the best-s |
| NULL (sequence) | pwm.count | pssm, score.thresh, bidirect, prior, extend, strand, spat_* | Count of anchors whose score |

K-mer summarizers

| Source | func | Key params | Description |
|-----------------|------------|----------------------|---|
| NULL (sequence) | kmer.count | kmer, extend, strand | Number of k-mer occurrences whose anchor lies inside the iterator interval. |
| NULL (sequence) | kmer.frac | kmer, extend, strand | Fraction of possible anchors within the interval that match the k-mer. |

Masked sequence summarizers

| Source | func | Key params | Description |
|-----------------|--------------|------------|--|
| NULL (sequence) | masked.count | NULL | Number of masked (lowercase) base pairs in the iterator interval. |
| NULL (sequence) | masked.frac | NULL | Fraction of base pairs in the iterator interval that are masked (lowercase). |

The sections below provide additional notes for motif, interval, k-mer, and masked sequence functions.

Motif (PWM) notes

- `pssm`: Position-specific scoring matrix (matrix or data frame) with columns A, C, G, T; extra columns are ignored.
- `bidirect`: When TRUE (default), both strands are scanned and combined per genomic start (per-position union). The `strand` argument is ignored. When FALSE, only the strand specified by `strand` is scanned.
- `prior`: Pseudocount added to frequencies (default 0.01). Set to 0 to disable.
- `extend`: Extends the fetched sequence so boundary-anchored motifs retain full context (default TRUE). The END coordinate is padded by `motif_length - 1` for all strand modes; anchors must still start inside the iterator.
- Neutral characters (N, n, *) contribute the mean log-probability of the corresponding PSSM column on both strands.
- `strand`: Used only when `bidirect = FALSE`; 1 scans the forward strand, -1 scans the reverse strand. For `pwm.max.pos`, `strand = -1` reports the hit position at the end of the match (still relative to the forward orientation).
- `score.thresh`: Threshold for `pwm.count`. Anchors with log-likelihood \geq `score.thresh` are counted; only one count per genomic start.
- Spatial weighting (`spat_factor`, `spat_bin`, `spat_min`, `spat_max`): optional position-dependent weights applied in log-space. Provide a positive numeric vector `spat_factor`; `spat_bin` (integer > 0) defines bin width; `spat_min/spat_max` restrict the scanning window.
- `pwm.max.pos`: Positions are reported 1-based relative to the final scan window (after iterator shifts and spatial trimming). Ties resolve to the most 5' anchor; the forward strand wins ties at the same coordinate. Values are signed when `bidirect = TRUE` (positive for forward, negative for reverse).

Spatial weighting enables position-dependent weighting for modeling positional biases. Bins are 0-indexed from the scan start. When using `gvtrack.iterator()` shifts (e.g., `sshift = -50`, `eshift = 50`), bins index from the expanded scan window start, not the original interval. Both strands use the same bin at each genomic position. Positions beyond the last bin reuse the final bin's weight. If the window size is not divisible by `spat_bin`, the last bin is shorter (e.g., scanning 500 bp with

40 bp bins yields bins 0-11 of 40 bp plus bin 12 of 20 bp). Use `spat_min` and `spat_max` to restrict scanning to a range divisible by `spat_bin` if needed.

PWM parameters can be supplied either as a single list (`params`) or via named arguments (see examples).

Interval distance notes

`distance`: Given the center 'C' of the current iterator interval, returns 'DC * X/2' where 'DC' is the normalized distance to the center of the interval that contains 'C', and 'X' is the value of the parameter (default: 0). If no interval contains 'C', the result is 'D + X/2' where 'D' is the distance between 'C' and the edge of the closest interval.

`distance.center`: Given the center 'C' of the current iterator interval, returns NaN if 'C' is outside of all intervals, otherwise returns the distance between 'C' and the center of the closest interval.

`distance.edge`: Computes edge-to-edge distance from the iterator interval to the closest source interval, using the same calculation as `gintervals.neighbors`. Returns 0 for overlapping intervals. Distance sign depends on the strand column of source intervals; returns unsigned (absolute) distance if no strand column exists. Returns NA if no source intervals exist on the current chromosome.

For `distance` and `distance.center`, distance can be positive or negative depending on the position of the coordinate relative to the interval and the strand (-1 or 1) of the interval. Distance is always positive if `strand = 0` or if the strand column is missing. The result is NA if no intervals exist for the current chromosome.

Difference between distance functions: The `distance` function measures from the *center* of the iterator interval (a single coordinate point) to the closest *edge* of source intervals when outside, or returns a normalized distance within the interval when inside. The `distance.center` function measures from the center of the iterator interval to the *center* of source intervals. The `distance.edge` function measures *edge-to-edge* distance between intervals, exactly like `gintervals.neighbors`. Use `distance.edge` when you need the same distance computation as `gintervals.neighbors` within a virtual track context.

K-mer notes

- `kmer`: DNA sequence (case-insensitive) to count.
- `extend`: If TRUE (default), counts kmers whose anchor lies in the interval even if the kmer extends beyond it; when FALSE, only kmers fully contained in the interval are considered.
- `strand`: 1 counts forward-strand occurrences, -1 counts reverse-strand occurrences, 0 counts both strands (default). For palindromic kmers, consider using 1 or -1 to avoid double counting.

K-mer parameters can be supplied as a list or via named arguments (see examples).

Modify iterator behavior with 'gvtrack.iterator' or 'gvtrack.iterator.2d'.

Value

None.

See Also

[gvtrack.info](#), [gvtrack.iterator](#), [gvtrack.iterator.2d](#), [gvtrack.array.slice](#), [gvtrack.ls](#), [gvtrack.rm](#)

[gvtrack.iterator](#), [gvtrack.iterator.2d](#), [gvtrack.filter](#)

Examples

```

gdb.init_examples()

gvtrack.create("vtrack1", "dense_track", "max")
gvtrack.create("vtrack2", "dense_track", "quantile", 0.5)
gextract("dense_track", "vtrack1", "vtrack2",
         gintervals(1, 0, 10000),
         iterator = 1000
        )

gvtrack.create("vtrack3", "dense_track", "global.percentile")
gvtrack.create("vtrack4", "annotations", "distance")
gdist(
  "vtrack3", seq(0, 1, l = 10), "vtrack4",
  seq(-500, 500, 200)
)

gvtrack.create("cov", "annotations", "coverage")
gextract("cov", gintervals(1, 0, 1000), iterator = 100)

pssm <- matrix(
  c(
    0.7, 0.1, 0.1, 0.1, # Example PSSM
    0.1, 0.7, 0.1, 0.1,
    0.1, 0.1, 0.7, 0.1,
    0.1, 0.1, 0.7, 0.1,
    0.1, 0.1, 0.7, 0.1,
    0.1, 0.1, 0.7, 0.1
  ),
  ncol = 4, byrow = TRUE
)
colnames(pssm) <- c("A", "C", "G", "T")
gvtrack.create(
  "motif_score", NULL, "pwm",
  list(pssm = pssm, bidirect = TRUE, prior = 0.01)
)
gvtrack.create("max_motif_score", NULL, "pwm.max",
  pssm = pssm, bidirect = TRUE, prior = 0.01
)
gvtrack.create("max_motif_pos", NULL, "pwm.max.pos",
  pssm = pssm
)
gextract(
  c(
    "dense_track", "motif_score", "max_motif_score",
    "max_motif_pos"
  ),
  gintervals(1, 0, 10000),
  iterator = 500
)

```

```

# Kmer counting examples
gvtrack.create("cg_count", NULL, "kmer.count", kmer = "CG", strand = 1)
gvtrack.create("cg_frac", NULL, "kmer.frac", kmer = "CG", strand = 1)
gextract(c("cg_count", "cg_frac"), gintervals(1, 0, 10000), iterator = 1000)

gvtrack.create("at_pos", NULL, "kmer.count", kmer = "AT", strand = 1)
gvtrack.create("at_neg", NULL, "kmer.count", kmer = "AT", strand = -1)
gvtrack.create("at_both", NULL, "kmer.count", kmer = "AT", strand = 0)
gextract(c("at_pos", "at_neg", "at_both"), gintervals(1, 0, 10000), iterator = 1000)

# GC content
gvtrack.create("g_frac", NULL, "kmer.frac", kmer = "G")
gvtrack.create("c_frac", NULL, "kmer.frac", kmer = "C")
gextract("g_frac + c_frac", gintervals(1, 0, 10000),
        iterator = 1000,
        colnames = "gc_content"
)

# Masked base pair counting
gvtrack.create("masked_count", NULL, "masked.count")
gvtrack.create("masked_frac", NULL, "masked.frac")
gextract(c("masked_count", "masked_frac"), gintervals(1, 0, 10000), iterator = 1000)

# Combined with GC content (unmasked regions only)
gvtrack.create("gc", NULL, "kmer.frac", kmer = "G")
gextract("gc * (1 - masked_frac)",
        gintervals(1, 0, 10000),
        iterator = 1000,
        colnames = "gc_unmasked"
)

# Value-based track examples
# Create a data frame with intervals and numeric values
intervals_with_values <- data.frame(
  chrom = "chr1",
  start = c(100, 300, 500),
  end = c(200, 400, 600),
  score = c(10, 20, 30)
)
# Use as value-based sparse track (functions like sparse track)
gvtrack.create("value_track", intervals_with_values, "avg")
gvtrack.create("value_track_max", intervals_with_values, "max")
gextract(c("value_track", "value_track_max"),
        gintervals(1, 0, 10000),
        iterator = 1000
)

# Spatial PWM examples
# Create a PWM with higher weight in the center of intervals
pssm <- matrix(
  c(
    0.7, 0.1, 0.1, 0.1,
    0.1, 0.7, 0.1, 0.1,
  )

```

```

        0.1, 0.1, 0.7, 0.1,
        0.1, 0.1, 0.1, 0.7
    ),
    ncol = 4, byrow = TRUE
)
colnames(pssm) <- c("A", "C", "G", "T")

# Spatial factors: low weight at edges, high in center
# For 200bp intervals with 40bp bins: bins 0, 40, 80, 120, 160
spatial_weights <- c(0.5, 1.0, 2.0, 1.0, 0.5)

gvtrack.create(
  "spatial_pwm", NULL, "pwm",
  list(
    pssm = pssm,
    bidirect = TRUE,
    spat_factor = spatial_weights,
    spat_bin = 40L
  )
)

# Compare with non-spatial PWM
gvtrack.create(
  "regular_pwm", NULL, "pwm",
  list(pssm = pssm, bidirect = TRUE)
)

gextract(c("spatial_pwm", "regular_pwm"),
  gintervals(1, 0, 10000),
  iterator = 200
)

# Using spatial parameters with iterator shifts
gvtrack.create(
  "spatial_extended", NULL, "pwm.max",
  pssm = pssm,
  spat_factor = c(0.5, 1.0, 2.0, 2.5, 2.0, 1.0, 0.5),
  spat_bin = 40L
)
# Scan window will be 280bp (100bp + 2*90bp)
gvtrack.iterator("spatial_extended", sshift = -90, eshift = 90)
gextract("spatial_extended", gintervals(1, 0, 10000), iterator = 100)

# Using spat_min/spat_max to restrict scanning to a window
# For 500bp intervals, scan only positions 30-470 (440bp window)
gvtrack.create(
  "window_pwm", NULL, "pwm",
  pssm = pssm,
  bidirect = TRUE,
  spat_min = 30, # 1-based position
  spat_max = 470 # 1-based position
)
gextract("window_pwm", gintervals(1, 0, 10000), iterator = 500)

```

```

# Combining spatial weighting with window restriction
# Scan positions 50-450 with spatial weights favoring the center
gvtrack.create(
  "window_spatial_pwm", NULL, "pwm",
  pssm = pssm,
  bidirect = TRUE,
  spat_factor = c(0.5, 1.0, 2.0, 2.5, 2.0, 1.0, 0.5, 1.0, 0.5, 0.5),
  spat_bin = 40L,
  spat_min = 50,
  spat_max = 450
)
gextract("window_spatial_pwm", gintervals(1, 0, 10000), iterator = 500)

```

| | |
|----------------|---|
| gvtrack.filter | <i>Attach or clear a genomic mask filter on a virtual track</i> |
|----------------|---|

Description

Attaches or clears a genomic mask filter on a virtual track. When a filter is attached, the virtual track function is evaluated only over the unmasked regions (i.e., regions not covered by the filter intervals).

Usage

```
gvtrack.filter(vtrack = NULL, filter = NULL)
```

Arguments

| | |
|--------|--|
| vtrack | virtual track name |
| filter | genomic mask to apply. Can be: <ul style="list-style-type: none"> • A data.frame with columns 'chrom', 'start', 'end' (intervals to mask) • A character string naming an intervals set • A character string naming a track (must be intervals-type track) • A list of any combination of the above (all will be unified) • NULL to clear the filter |

Details

The filter defines regions to *exclude* from virtual track evaluation. The virtual track function will be evaluated only on the complement of the filter. Once a filter is attached to a virtual track, it applies to **all subsequent extractions** of that virtual track until explicitly cleared with `filter = NULL`.

Order of Operations:

Filters are applied *after* iterator modifiers (sshift/eshift/dim). The order is:

1. Apply iterator modifiers (gvtrack.iterator with sshift/eshift)
2. Subtract mask from the modified intervals

3. Evaluate virtual track function over unmasked regions

Semantics by function type:

- **Aggregations (avg/sum/min/max/stddev/quantile):** Length-weighted over unmasked regions
- **coverage:** Returns (covered length in unmasked region) / (total unmasked length)
- **distance/distance.center:** Unaffected by mask (pure geometry)
- **PWM/kmer:** Masked bases act as hard boundaries; matches cannot span masked regions. **Important:** When extend=TRUE (the default), motifs at the boundaries of unmasked segments can use bases from the adjacent masked regions to complete the motif scoring. For example, if a 4bp motif starts at position 1998 in an unmasked region that ends at 2000, and positions 2000-2002 are masked, the motif will still be scored using the masked bases. In other words, motif matches *starting positions* must be in unmasked regions, but the motif sequence itself can extend into masked regions when extend=TRUE. Set extend=FALSE to prevent any use of masked bases in scoring.

Completely Masked Intervals: If an entire iterator interval is masked, the function returns NA (not 0).

Value

None (invisibly).

See Also

[gvtrack.create](#), [gvtrack.iterator](#), [gvtrack.info](#)

Examples

```
gdb.init_examples()

## Basic usage: Excluding specific regions
gvtrack.create("vtrack1", "dense_track", func = "avg")

# Create intervals to mask (e.g., repetitive regions)
repeats <- gintervals(c(1, 1), c(100, 500), c(200, 600))

# Attach filter - track will be evaluated excluding these regions
gvtrack.filter("vtrack1", filter = repeats)

# Extract values - masked regions are excluded from calculation
result_filtered <- gextract("vtrack1", gintervals(1, 0, 1000))

# Check filter info
gvtrack.info("vtrack1")

# Clear the filter and compare
gvtrack.filter("vtrack1", filter = NULL)
result_unfiltered <- gextract("vtrack1", gintervals(1, 0, 1000))
```

```
## Using multiple filter sources (combined automatically)
centromeres <- gintervals(1, 10000, 15000)
telomeres <- gintervals(1, 0, 1000)
combined_mask <- list(repeats, centromeres, telomeres)

gvtrack.filter("vtrack1", filter = combined_mask)
result_multi_filter <- gextract("vtrack1", gintervals(1, 0, 20000))

## Filters work with iterator modifiers
gvtrack.create("vtrack2", "dense_track", func = "sum")
gvtrack.filter("vtrack2", filter = repeats)
gvtrack.iterator("vtrack2", sshift = -50, eshift = 50)

# Iterator shifts applied first, then mask subtracted
result_shifted <- gextract("vtrack2", gintervals(1, 1000, 2000), iterator = 100)
```

| | |
|--------------|--|
| gvtrack.info | <i>Returns the definition of a virtual track</i> |
|--------------|--|

Description

Returns the definition of a virtual track.

Usage

```
gvtrack.info(vtrack = NULL)
```

Arguments

| | |
|--------|--------------------|
| vtrack | virtual track name |
|--------|--------------------|

Details

This function returns the internal representation of a virtual track.

Value

Internal representation of a virtual track.

See Also

[gvtrack.create](#)

Examples

```
gdb.init_examples()
gvtrack.create("vtrack1", "dense_track", "max")
gvtrack.info("vtrack1")
```

| | |
|------------------|---|
| gvtrack.iterator | <i>Defines modification rules for a one-dimensional iterator in a virtual track</i> |
|------------------|---|

Description

Defines modification rules for a one-dimensional iterator in a virtual track.

Usage

```
gvtrack.iterator(vtrack = NULL, dim = NULL, sshift = 0, eshift = 0)
```

Arguments

| | |
|--------|--|
| vtrack | virtual track name |
| dim | use 'NULL' or '0' for 1D iterators. '1' converts 2D iterator to (chrom1, start1, end1), '2' converts 2D iterator to (chrom2, start2, end2) |
| sshift | shift of 'start' coordinate |
| eshift | shift of 'end' coordinate |

Details

This function defines modification rules for one-dimensional iterator intervals in a virtual track.

'dim' converts a 2D iterator interval (chrom1, start1, end1, chrom2, start2, end2) to a 1D interval. If 'dim' is '1' the interval is converted to (chrom1, start1, end1). If 'dim' is '2' the interval is converted to (chrom2, start2, end2). If 1D iterator is used 'dim' must be set to 'NULL' or '0' (meaning: no conversion is made).

Iterator interval's 'start' coordinate is modified by adding 'sshift'. Similarly 'end' coordinate is altered by adding 'eshift'.

Value

None.

See Also

[gvtrack.create](#), [gvtrack.iterator.2d](#)

Examples

```
gdb.init_examples()

gvtrack.create("vtrack1", "dense_track")
gvtrack.iterator("vtrack1", sshift = 200, eshift = 200)
gextract("dense_track", "vtrack1", gintervals(1, 0, 500))
```

```
gvtrack.create("vtrack2", "dense_track")
gvtrack.iterator("vtrack2", dim = 1)
gextract("vtrack2", gintervals.2d(1, 0, 1000, 1, 0, -1),
        iterator = "rects_track"
    )
```

gvtrack.iterator.2d *Defines modification rules for a two-dimensional iterator in a virtual track*

Description

Defines modification rules for a two-dimensional iterator in a virtual track.

Usage

```
gvtrack.iterator.2d(
  vtrack = NULL,
  sshift1 = 0,
  eshift1 = 0,
  sshift2 = 0,
  eshift2 = 0
)
```

Arguments

| | |
|---------|------------------------------|
| vtrack | virtual track name |
| sshift1 | shift of 'start1' coordinate |
| eshift1 | shift of 'end1' coordinate |
| sshift2 | shift of 'start2' coordinate |
| eshift2 | shift of 'end2' coordinate |

Details

This function defines modification rules for one-dimensional iterator intervals in a virtual track.

Iterator interval's 'start1' coordinate is modified by adding 'sshift1'. Similarly 'end1', 'start2', 'end2' coordinates are altered by adding 'eshift1', 'sshift2' and 'eshift2' accordingly.

Value

None.

See Also

[gvtrack.create](#), [gvtrack.iterator](#)

Examples

```
gdb.init_examples()
gvtrack.create("vtrack1", "rects_track")
gvtrack.iterator.2d("vtrack1", sshift1 = 1000, eshift1 = 2000)
gextract(
  "rects_track", "vtrack1",
  gintervals.2d(1, 0, 5000, 2, 0, 5000)
)
```

gvtrack.ls

Returns a list of virtual track names

Description

Returns a list of virtual track names.

Usage

```
gvtrack.ls(
  pattern = "",
  ignore.case = FALSE,
  perl = FALSE,
  fixed = FALSE,
  useBytes = FALSE
)
```

Arguments

pattern, ignore.case, perl, fixed, useBytes
see 'grep'

Details

This function returns a list of virtual tracks that exist in current R environment that match the pattern (see 'grep'). If called without any arguments all virtual tracks are returned.

Value

An array that contains the names of virtual tracks.

See Also

[grep](#), [gvtrack.create](#), [gvtrack.rm](#)

Examples

```
gdb.init_examples()
gvtrack.create("vtrack1", "dense_track", "max")
gvtrack.create("vtrack2", "dense_track", "quantile", 0.5)
gvtrack.ls()
gvtrack.ls(pattern = "*2")
```

gvtrack.rm

Deletes a virtual track

Description

Deletes a virtual track.

Usage

```
gvtrack.rm(vtrack = NULL)
```

Arguments

vtrack virtual track name

Details

This function deletes a virtual track from current R environment.

Value

None.

See Also

[gvtrack.create](#), [gvtrack.ls](#)

Examples

```
gdb.init_examples()
gvtrack.create("vtrack1", "dense_track", "max")
gvtrack.create("vtrack2", "dense_track", "quantile", 0.5)
gvtrack.ls()
gvtrack.rm("vtrack1")
gvtrack.ls()
```

| | |
|-------|--|
| gwget | <i>Downloads files from FTP server</i> |
|-------|--|

Description

Downloads multiple files from FTP server

Usage

```
gwget(url = NULL, path = NULL)
```

Arguments

| | |
|------|--|
| url | URL of FTP server |
| path | directory path where the downloaded files are stored |

Details

This function downloads files from FTP server given by 'url'. The address in 'url' can contain wildcards to download more than one file at once. Files are downloaded to a directory given by 'path' argument. If 'path' is 'NULL', file are downloaded into 'GROOT/downloads'.

Value

An array of file names that have been downloaded.

See Also

[gtrack.import_set](#)

Examples

```
gdb.init_examples()

outdir <- tempdir()
gwget("ftp://hgdownload.soe.ucsc.edu/goldenPath/hg19/chromosomes/md5sum.txt", path = outdir)
```

| | |
|---------|--|
| gwilcox | <i>Calculates Wilcoxon test on sliding windows over track expression</i> |
|---------|--|

Description

Calculates Wilcoxon test on sliding windows over the values of track expression.

Usage

```
gwilcox(
  expr = NULL,
  winsize1 = NULL,
  winsize2 = NULL,
  maxpval = 0.05,
  onetailed = TRUE,
  what2find = 1,
  intervals = NULL,
  iterator = NULL,
  intervals.set.out = NULL
)
```

Arguments

| | |
|-------------------|---|
| expr | track expression |
| winsize1 | number of values in the first sliding window |
| winsize2 | number of values in the second sliding window |
| maxpval | maximal P-value |
| onetailed | if 'TRUE', Wilcoxon test is performed one tailed, otherwise two tailed |
| what2find | if '-1', lows are searched. If '1', peaks are searched. If '0', both peaks and lows are searched |
| intervals | genomic scope for which the function is applied |
| iterator | track expression iterator of "fixed bin" type. If 'NULL' iterator is determined implicitly based on track expression. |
| intervals.set.out | intervals set name where the function result is optionally outputted |

Details

This function runs a Wilcoxon test (also known as a Mann-Whitney test) over the values of track expression in the two sliding windows having an identical center. The sizes of the windows are specified by 'winsize1' and 'winsize2'. 'gwilcox' returns intervals where the smaller window tested against a larger window gives a P-value below 'maxpval'. The test can be one or two tailed.

'what2find' argument controls what should be searched: peaks, lows or both.

If 'intervals.set.out' is not 'NULL' the result is saved as an intervals set. Use this parameter if the result size exceeds the limits of the physical memory.

Value

If 'intervals.set.out' is 'NULL' a data frame representing the intervals with an additional 'pval' column where P-value is below 'maxpval'.

See Also

[gscreen](#), [gsegment](#)

Examples

```
gdb.init_examples()
gwilcox("dense_track", 100000, 1000,
  maxpval = 0.01,
  what2find = 1
)
```

`print.gsynth.model` *Print summary of a gsynth.model*

Description

Print summary of a gsynth.model

Usage

```
## S3 method for class 'gsynth.model'
print(x, ...)
```

Arguments

| | |
|------------------|--------------------------------|
| <code>x</code> | A gsynth.model object |
| <code>...</code> | Additional arguments (ignored) |

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