

Package: tsmp (via r-universe)

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Type Package

Title Time Series with Matrix Profile

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Description A toolkit implementing the Matrix Profile concept that was created by CS-UCR
<<http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>>.

License Apache License (>= 2.0)

URL <https://github.com/matrix-profile-foundation/tsmp>

BugReports <https://github.com/matrix-profile-foundation/tsmp/issues>

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Contents

analyze	3
as.matrixprofile	5
av_apply	6
av_complexity	7
av_hardlimit_artifact	8
av_motion_artifact	9
av_stop_word	10
av_zerocrossing	11
compute	12
discords	13
dist_profile	14
fast_avg_sd	16
fast_movavg	16
fast_movsd	17
find_chains	17
find_discord	18
find_motif	19
find_snippet	21
floss	22
floss_cac	24
floss_extract	25
fluss	26
fluss_cac	27
fluss_extract	28
fluss_score	29
get_data	30
mass_v3	30
min_mp_idx	32
motifs	32
motifs_discords_small	33
mpdist	34
mpx	35
mp_fluss_data	36
mp_gait_data	37
mp_meat_data	37
mp_test_data	38
mp_toy_data	39
mstomp_par	40
plot	42
plot_arcs	45
pmp	47
pmp_upper_bound	48
read	50
remove_class	50
salient_mds	51
salient_score	52

salient_subsequences	53
scrimp	54
sdtc_predict	56
sdtc_score	57
sdtc_train	58
set_data	60
simple_fast	60
stamp_par	61
stompi_update	63
stomp_par	64
tsmp	66
valmod	69
visualize	70
write	71

Index**72**

analyze	<i>Runs an appropriate workflow based on the parameters passed in.</i>
---------	--

Description

The goal of this function is to compute all fundamental algorithms on the provided time series data. See details for more information.

Usage

```
analyze(
  ts,
  windows = NULL,
  query = NULL,
  sample_pct = 1,
  threshold = 0.98,
  n_jobs = 1L
)
```

Arguments

ts	a matrix or a vector. The time series to analyze.
windows	an int or a vector. The window(s) to compute the Matrix Profile. Note that it may be an int for a single matrix profile computation or a vector of int for computing the Pan-Matrix Profile.
query	a matrix or a vector. Optional The query to analyze. Note that when computing the Pan-Matrix Profile the query is ignored!
sample_pct	a numeric. A number between 0 and 1 representing how many samples to compute for the Matrix Profile or Pan-Matrix Profile. When it is 1, the exact algorithm is used. (default is 1.0).

`threshold` a numeric. Correlation threshold. See details. (Default is 0.98).
`n_jobs` an int. The number of cpu cores to use when computing the MatrixProfile. (default is 1).

Details

For now the following is computed:

1. Matrix Profile - exact or approximate based on `sample_pct` given that a single windows is provided. By default is the exact algorithm;
2. Top 3 Motifs;
3. Top 3 Discords;
4. Plot Matrix Profile, Motifs and Discords.

When `windows` is not provided or more than a single window is provided, the Pan-Matrix Profile is computed:

1. Compute the upper bound when a `threshold` is provided (it is, by default);
2. Compute Pan-Matrix Profile for all windows provided, below the upper bound, or a default range when no windows is provided;
3. Top Motifs;
4. Top Discords;
5. Plot Pan-Matrix Profile, motifs and discords.

Value

The appropriate Matrix Profile or Pan-Matrix Profile profile object and also plots the graphics.

References

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

See Also

Other Main API: `compute()`, `discords()`, `motifs()`, `visualize()`

Examples

```
# Matrix Profile
result <- analyze(mp_toy_data$data[, 1], 80)

# Pan Matrix Profile
result <- analyze(mp_toy_data$data[, 1])
```

as.matrixprofile	<i>Convert a TSMP object into another if possible</i>
------------------	---

Description

The base Classes are MatrixProfile and MultiMatrixProfile, but as other functions are used, classes are pushed behind, since the last output normally is the most significant. If you want, for example, to plot the Matrix Profile from a Fluss object, you may use `as.matrixprofile()` to cast it back.

Usage

```
as.matrixprofile(.mp)
as.multimatrixprofile(.mp)
as.pmp(.mp)
as.valmod(.mp)
as.fluss(.mp)
as.chain(.mp)
as.discord(.mp)
as.motif(.mp)
as.multimotif(.mp)
as.arccount(.mp)
as.salient(.mp)
```

Arguments

`.mp` a TSMP object.

Value

Returns the object with the new class, if possible.

Functions

- `as.matrixprofile()`: Cast an object changed by another function back to MatrixProfile.
- `as.multimatrixprofile()`: Cast an object changed by another function back to MultiMatrixProfile.
- `as.pmp()`: Cast an object changed by another function back to PMP.

- `as.valmod()`: Cast an object changed by another function back to MultiMatrixProfile.
- `as.fluss()`: Cast an object changed by another function back to Fluss.
- `as.chain()`: Cast an object changed by another function back to Chain.
- `as.discord()`: Cast an object changed by another function back to Discord.
- `as.motif()`: Cast an object changed by another function back to Motif.
- `as.multimotif()`: Cast an object changed by another function back to MultiMotif.
- `as.arccount()`: Cast an object changed by another function back to ArcCount.
- `as.salient()`: Cast an object changed by another function back to Salient.

Examples

```
w <- 50
data <- mp_gait_data
mp <- tsmp(data, window_size = w, exclusion_zone = 1 / 4, verbose = 0)
mp <- find_motif(mp)
class(mp) # first class will be "Motif"

plot(mp) # plots a motif plot

plot(as.matrixprofile(mp)) # plots a matrix profile plot
```

av_apply

Corrects the matrix profile using an annotation vector

Description

This function overwrites the current Matrix Profile using the Annotation Vector. Use with caution.

Usage

```
av_apply(.mp)
```

Arguments

`.mp` A Matrix Profile with an Annotation Vector.

Value

Returns the input `.mp` object corrected by the embedded annotation vector.

References

- Dau HA, Keogh E. Matrix Profile V: A Generic Technique to Incorporate Domain Knowledge into Motif Discovery. In: Proceedings of the 23rd ACM SIGKDD International Conference on Knowledge Discovery and Data Mining - KDD '17. New York, New York, USA: ACM Press; 2017. p. 125-34.

See Also

Other Annotation vectors: [av_complexity\(\)](#), [av_hardlimit_artifact\(\)](#), [av_motion_artifact\(\)](#), [av_stop_word\(\)](#), [av_zerocrossing\(\)](#)

Examples

```
data <- mp_test_data$train$data[1:1000]
w <- 50
mp <- tsmp(data, window_size = w, verbose = 0)
mp <- av_complexity(mp)
av <- av_apply(mp)
```

av_complexity	<i>Computes the annotation vector that favors complexity</i>
---------------	--

Description

Computes the annotation vector that favors complexity

Usage

```
av_complexity(.mp, data, dilution_factor = 0, apply = FALSE)
```

Arguments

.mp	a Matrix Profile object.
data	a vector or a column matrix of numeric.
dilution_factor	a numeric. (Default is 0). Larger numbers means more dilution.
apply	logical. (Default is FALSE). Applies the Annotation Vector over the Matrix Profile. Use with caution.

Value

Returns the input .mp object with an embedded annotation vector.

References

- Dau HA, Keogh E. Matrix Profile V: A Generic Technique to Incorporate Domain Knowledge into Motif Discovery. In: Proceedings of the 23rd ACM SIGKDD International Conference on Knowledge Discovery and Data Mining - KDD '17. New York, New York, USA: ACM Press; 2017. p. 125-34.

See Also

Other Annotation vectors: [av_apply\(\)](#), [av_hardlimit_artifact\(\)](#), [av_motion_artifact\(\)](#), [av_stop_word\(\)](#), [av_zerocrossing\(\)](#)

Examples

```
data <- mp_test_data$train$data[1:1000]
w <- 50
mp <- tsmf(data, window_size = w, verbose = 0)
av <- av_complexity(mp, apply = TRUE)
```

av_hardlimit_artifact *Computes the annotation vector that suppresses hard-limited artifacts*

Description

Computes the annotation vector that suppresses hard-limited artifacts

Usage

```
av_hardlimit_artifact(.mp, data, apply = FALSE)
```

Arguments

.mp	a Matrix Profile object.
data	a vector or a column matrix of numeric.
apply	logical. (Default is FALSE). Applies the Annotation Vector over the Matrix Profile. Use with caution.

Value

Returns the input .mp object with an embedded annotation vector.

References

- Dau HA, Keogh E. Matrix Profile V: A Generic Technique to Incorporate Domain Knowledge into Motif Discovery. In: Proceedings of the 23rd ACM SIGKDD International Conference on Knowledge Discovery and Data Mining - KDD '17. New York, New York, USA: ACM Press; 2017. p. 125-34.

See Also

Other Annotation vectors: [av_apply\(\)](#), [av_complexity\(\)](#), [av_motion_artifact\(\)](#), [av_stop_word\(\)](#), [av_zerocrossing\(\)](#)

Examples

```
data <- mp_test_data$train$data[1:1000]
w <- 50
mp <- tsmf(data, window_size = w, verbose = 0)
av <- av_hardlimit_artifact(mp, apply = TRUE)
```

av_motion_artifact *Computes the annotation vector that suppresses motion artifacts*

Description

Computes the annotation vector that suppresses motion artifacts

Usage

```
av_motion_artifact(.mp, data, apply = FALSE)
```

Arguments

.mp	a Matrix Profile object.
data	a vector or a column matrix of numeric.
apply	logical. (Default is FALSE). Applies the Annotation Vector over the Matrix Profile. Use with caution.

Value

Returns the input .mp object with an embedded annotation vector.

References

- Dau HA, Keogh E. Matrix Profile V: A Generic Technique to Incorporate Domain Knowledge into Motif Discovery. In: Proceedings of the 23rd ACM SIGKDD International Conference on Knowledge Discovery and Data Mining - KDD '17. New York, New York, USA: ACM Press; 2017. p. 125-34.

See Also

Other Annotation vectors: [av_apply\(\)](#), [av_complexity\(\)](#), [av_hardlimit_artifact\(\)](#), [av_stop_word\(\)](#), [av_zerocrossing\(\)](#)

Examples

```
data <- mp_test_data$train$data[1:1000]
w <- 50
mp <- tsmpp(data, window_size = w, verbose = 0)
av <- av_motion_artifact(mp, apply = TRUE)
```

`av_stop_word`*Computes the annotation vector that suppresses stop-word motifs*

Description

Computes the annotation vector that suppresses stop-word motifs

Usage

```
av_stop_word(  
    .mp,  
    data,  
    stop_word_loc,  
    exclusion_zone = NULL,  
    threshold = 0.1,  
    apply = FALSE  
)
```

Arguments

<code>.mp</code>	a Matrix Profile object.
<code>data</code>	a vector or a column matrix of numeric.
<code>stop_word_loc</code>	an int. The index of stop word location.
<code>exclusion_zone</code>	a numeric. Size of the exclusion zone, based on <code>window_size</code> (default is NULL). See details.
<code>threshold</code>	a numeric. (default is 0.1).
<code>apply</code>	logical. (Default is FALSE). Applies the Annotation Vector over the Matrix Profile. Use with caution.

Details

The function is intended to be generic. However, its parameters (`stop_word_loc`, `exclusion_zone` and `threshold`) are highly dataset dependent.

Value

Returns the input `.mp` object with an embedded annotation vector.

References

- Dau HA, Keogh E. Matrix Profile V: A Generic Technique to Incorporate Domain Knowledge into Motif Discovery. In: Proceedings of the 23rd ACM SIGKDD International Conference on Knowledge Discovery and Data Mining - KDD '17. New York, New York, USA: ACM Press; 2017. p. 125-34.

See Also

Other Annotation vectors: [av_apply\(\)](#), [av_complexity\(\)](#), [av_hardlimit_artifact\(\)](#), [av_motion_artifact\(\)](#), [av_zerocrossing\(\)](#)

Examples

```
data <- mp_test_data$train$data[1:1000]
w <- 50
mp <- tsmf(data, window_size = w, verbose = 0)
av <- av_stop_word(mp, stop_word_loc = 150, apply = TRUE)
```

av_zerocrossing	<i>Computes the annotation vector that favors number of zero crossing</i>
-----------------	---

Description

Computes the annotation vector that favors number of zero crossing

Usage

```
av_zerocrossing(.mp, data, apply = FALSE)
```

Arguments

.mp	a Matrix Profile object.
data	a vector or a column matrix of numeric.
apply	logical. (Default is FALSE). Applies the Annotation Vector over the Matrix Profile. Use with caution.

Value

Returns the input .mp object with an embedded annotation vector.

References

- Dau HA, Keogh E. Matrix Profile V: A Generic Technique to Incorporate Domain Knowledge into Motif Discovery. In: Proceedings of the 23rd ACM SIGKDD International Conference on Knowledge Discovery and Data Mining - KDD '17. New York, New York, USA: ACM Press; 2017. p. 125-34.

See Also

Other Annotation vectors: [av_apply\(\)](#), [av_complexity\(\)](#), [av_hardlimit_artifact\(\)](#), [av_motion_artifact\(\)](#), [av_stop_word\(\)](#)

Examples

```
data <- mp_test_data$train$data[1:1000]
w <- 50
mp <- tsmp(data, window_size = w, verbose = 0)
av <- av_zerocrossing(mp, apply = TRUE)
```

compute

Computes the Matrix Profile or Pan-Matrix Profile

Description

Main API Function

Usage

```
compute(
  ts,
  windows = NULL,
  query = NULL,
  sample_pct = 1,
  threshold = 0.98,
  n_jobs = 1L
)
```

Arguments

ts	a matrix or a vector. The time series to analyze.
windows	an int or a vector. The window(s) to compute the Matrix Profile. Note that it may be an int for a single matrix profile computation or a vector of int for computing the Pan-Matrix Profile.
query	a matrix or a vector. Optional The query to analyze. Note that when computing the Pan-Matrix Profile the query is ignored!
sample_pct	a numeric. A number between 0 and 1 representing how many samples to compute for the Matrix Profile or Pan-Matrix Profile. When it is 1, the exact algorithm is used. (default is 1.0).
threshold	a numeric. Correlation threshold. See details. (Default is 0.98).
n_jobs	an int. The number of cpu cores to use when computing the MatrixProfile. (default is 1).

Details

Computes the exact or approximate Matrix Profile based on the sample percent specified. Currently, MPX and SCRIMP++ are used for the exact and approximate algorithms respectively. See details for more information about the arguments combinations.

When a single windows is given, the Matrix Profile is computed. If a query is provided, AB join is computed. Otherwise the self-join is computed. When multiple windows or none are given, the

Pan-Matrix Profile is computed. If a threshold is set (it is, by default), the upper bound will be computed and the given windows or a default range (when no windows), below the upper bound will be computed.

Value

The profile computed.

References

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

See Also

Other Main API: [analyze\(\)](#), [discords\(\)](#), [motifs\(\)](#), [visualize\(\)](#)

Examples

```
# Matrix Profile
result <- compute(mp_toy_data$data[, 1], 80)

# Pan-Matrix Profile
result <- compute(mp_toy_data$data[, 1])
```

discords

Search for Discord

Description

Search for Discord

Usage

```
discords(
  profile,
  exclusion_zone = profile$ez,
  k = 3L,
  neighbor_count = 10L,
  radius = 3
)
```

Arguments

profile a MatrixProfile or PMP object.

exclusion_zone an int. Number of values to exclude on both sides of the motif to avoid trivial matches. Defaults to the exclusion zone used to compute the (Pan-)Matrix Profile which is found in the profile data structure.

k an int. Number of discords to find. (Default is 3).
neighbor_count an int. Number of neighbors to find. (Default is 3).
radius an int. Set a threshold to exclude matching neighbors with distance > current discord distance * radius. (Default is 3).

References

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

See Also

Other Main API: [analyze\(\)](#), [compute\(\)](#), [motifs\(\)](#), [visualize\(\)](#)

dist_profile	<i>Calculates the distance profile using MASS algorithms</i>
--------------	--

Description

Mueen's Algorithm for Similarity Search is The Fastest Similarity Search Algorithm for Time Series Subsequences under Euclidean Distance and Correlation Coefficient.

Usage

```

dist_profile(
  data,
  query,
  ...,
  window_size = NULL,
  method = "v3",
  index = 1,
  k = NULL,
  weight = NULL,
  paa = 1
)

```

Arguments

data a matrix or a vector.
query a matrix or a vector. See details.
... Precomputed values from the first iteration. If not supplied, these values will be computed.
window_size an int or NULL. Sliding window size. See details.
method method that will be used to calculate the distance profile. See details.
index an int. Index of query window. See details.

k	an int or NULL. Default is NULL. Defines the size of batch for MASS V3. Prefer to use a power of 2. If NULL, it will be set automatically.
weight	a vector of numeric or NULL with the same length of the window_size. This is a MASS extension to weight the query.
paa	a numeric. Default is 1. Factor of PAA reduction (2 == half of size). This is a MASS extension.

Details

This function has several ways to work:

Case 1: You have a small sized query and the data. In this case you only have to provide the first two parameters data and query. Internally the window_size will be get from the query length.

Case 2: You have one or two data vectors and want to compute the join or self-similarity. In this case you need to use the recursive solution. The parameters are data, query, window_size and index. The first iteration don't need the index unless you are starting somewhere else. The query will be the source of a query_window, starting on index, with length of window_size.

The method defines which MASS will be used. Current supported values are: v2, v3, weighted.

Value

Returns the distance_profile for the given query and the last_product for STOMP algorithm and the parameters for recursive call. See details.

References

- Abdullah Mueen, Yan Zhu, Michael Yeh, Kaveh Kamgar, Krishnamurthy Viswanathan, Chetan Kumar Gupta and Eamonn Keogh (2015), The Fastest Similarity Search Algorithm for Time Series Subsequences under Euclidean Distance

Website: <https://www.cs.unm.edu/~mueen/FastestSimilaritySearch.html>

Examples

```
w <- mp_toy_data$sub_len
ref_data <- mp_toy_data$data[, 1]
# minimum example, data and query
nn <- dist_profile(ref_data, ref_data[1:w])
distance_profile <- sqrt(nn$distance_profile)

# data and indexed query
nn <- dist_profile(ref_data, ref_data, window_size = w, index = 10)
distance_profile <- sqrt(nn$distance_profile)

# recursive
nn <- NULL

for (i in seq_len(10)) {
  nn <- dist_profile(ref_data, ref_data, nn, window_size = w, index = i)
}
```

```
# weighted
weight <- c(rep(1, w / 3), rep(0.5, w / 3), rep(0.8, w / 3)) # just an example

nn <- dist_profile(ref_data, ref_data,
  window_size = w, index = 1, method = "weighted",
  weight = weight
)
distance_profile <- sqrt(nn$distance_profile)
```

fast_avg_sd	<i>Fast implementation of moving average and moving standard deviation</i>
-------------	--

Description

This function does not handle NA values

Usage

```
fast_avg_sd(data, window_size, rcpp = FALSE)
```

Arguments

data	a vector or a column matrix of numeric.
window_size	moving sd window size
rcpp	a logical. Uses rcpp implementation.

Value

Returns a list with avg and sd vectors

fast_movavg	<i>Fast implementation of moving average</i>
-------------	--

Description

This function does not handle NA values

Usage

```
fast_movavg(data, window_size)
```

Arguments

data	a vector or a column matrix of numeric.
window_size	moving sd window size

Value

Returns a vector with the moving average

Examples

```
data_avg <- fast_movavg(mp_toy_data$data[, 1], mp_toy_data$sub_len)
```

fast_movsd	<i>Fast implementation of moving standard deviation</i>
------------	---

Description

This function does not handle NA values

Usage

```
fast_movsd(data, window_size, rcpp = FALSE)
```

Arguments

data	a vector or a column matrix of numeric.
window_size	moving sd window size
rcpp	a logical. Uses rcpp implementation.

Value

Returns a vector with the moving standard deviation

Examples

```
data_sd <- fast_movsd(mp_toy_data$data[, 1], mp_toy_data$sub_len)
```

find_chains	<i>Find Time Series Chains</i>
-------------	--------------------------------

Description

Time Series Chains is a new primitive for time series data mining.

Usage

```
find_chains(.mp)
```

Arguments

.mp	a MatrixProfile object.
-----	-------------------------

Value

Returns the input `.mp` object with a new name `chain`. It contains: `chains`, a list of chains found with more than 2 patterns and `best` with the best one.

References

- Zhu Y, Imamura M, Nikovski D, Keogh E. Matrix Profile VII: Time Series Chains: A New Primitive for Time Series Data Mining. Knowl Inf Syst. 2018 Jun 2;1-27.

Website: <https://sites.google.com/site/timeserieschain/>

Examples

```
w <- 50
data <- mp_gait_data
mp <- tsmpr(data, window_size = w, exclusion_zone = 1 / 4, verbose = 0)
mp <- find_chains(mp)
```

find_discord	<i>Search for Discord</i>
--------------	---------------------------

Description

Search for Discord

Usage

```
find_discord(.mp, ...)
```

```
## S3 method for class 'MatrixProfile'
find_discord(
  .mp,
  data,
  n_discords = 1,
  n_neighbors = 3,
  radius = 3,
  exclusion_zone = NULL,
  ...
)
```

```
## S3 method for class 'PMP'
find_discord(
  .mp,
  data,
  n_discords = 1,
  n_neighbors = 3,
  radius = 3,
  exclusion_zone = NULL,
```

```
    ...
  )
```

Arguments

`.mp` a MatrixProfile object.

`...` further arguments to be passed to class specific function.

`data` the data used to build the Matrix Profile, if not embedded.

`n_discords` an int. Number of discords to find. (Default is 1).

`n_neighbors` an int. Number of neighbors to find. (Default is 3).

`radius` an int. Set a threshold to exclude matching neighbors with distance > current discord distance * radius. (Default is 3).

`exclusion_zone` if a number will be used instead of embedded value. (Default is NULL).

Value

For class MatrixProfile, returns the input `.mp` object with a new name `discord`. It contains: `discord_idx`, a vector of discords found

For class PMP, returns the input `.mp` object with a new name `discord`. It contains: `discord_idx`, a vector of discords found

Examples

```
# Single dimension data
w <- 50
data <- mp_gait_data
mp <- tsmf(data, window_size = w, exclusion_zone = 1 / 4, verbose = 0)
mp <- find_discord(mp)
pan <- tsmf(mp_gait_data, window_size = 20:30, mode = "pmp")
mp <- find_discord(pan)
```

find_motif

Search for Motifs

Description

Search for Motifs

Usage

```
find_motif(.mp, ...)
```

```
## S3 method for class 'MatrixProfile'
find_motif(
  .mp,
  data,
```

```

    n_motifs = 3,
    n_neighbors = 10,
    radius = 3,
    exclusion_zone = NULL,
    ...
)

## S3 method for class 'MultiMatrixProfile'
find_motif(
  .mp,
  data,
  n_motifs = 3,
  mode = c("guided", "unconstrained"),
  n_bit = 4,
  exclusion_zone = NULL,
  n_dim = NULL,
  ...
)

## S3 method for class 'PMP'
find_motif(
  .mp,
  data,
  n_motifs = 3,
  n_neighbors = 10,
  radius = 3,
  exclusion_zone = NULL,
  ...
)

```

Arguments

<code>.mp</code>	a MatrixProfile or MultiMatrixProfile object.
<code>...</code>	further arguments to be passed to class specific function.
<code>data</code>	the data used to build the Matrix Profile, if not embedded.
<code>n_motifs</code>	an int. Number of motifs to find. (Default is 3).
<code>n_neighbors</code>	an int. Number of neighbors to find. (Default is 10).
<code>radius</code>	an int. Set a threshold to exclude matching neighbors with distance > current motif distance * radius. (Default is 3).
<code>exclusion_zone</code>	if a number will be used instead of embedded value. (Default is NULL).
<code>mode</code>	a string. Guided or Unconstrained search. Allow partial match. (Default is guided).
<code>n_bit</code>	an int. Bit size for discretization. Ignored on Guided search. (Default is 4).
<code>n_dim</code>	an int. Number of dimensions to use on Guided search instead of embedded value. (Default is NULL).

Value

For class MatrixProfile, returns the input .mp object with a new name motif. It contains: motif_idx, a list of motif pairs found and motif_neighbor a list with respective motif's neighbors.

For class MultiMatrixProfile, returns the input .mp object with a new name motif. It contains: motif_idx, a vector of motifs found and motif_dim a list the dimensions where the motifs were found

For class PMP, returns the input .mp object with a new name motif. It contains: motif_idx, a list of motif pairs found and motif_neighbor a list with respective motif's neighbors.

Examples

```
# Single dimension data
w <- 50
data <- mp_gait_data
mp <- tsmp(data, window_size = w, exclusion_zone = 1 / 4, verbose = 0)
mp <- find_motif(mp)

# Multidimension data
w <- mp_toy_data$sub_len
data <- mp_toy_data$data[1:200, ]
mp <- tsmp(data, window_size = w, mode = "mstomp", verbose = 0)
mp <- find_motif(mp)
pan <- tsmp(mp_gait_data, window_size = 20:30, mode = "pmp")
mp <- find_motif(pan)
```

find_snippet

Time Series Snippets: A New Primitive for Time Series Data Mining

Description

Time Series Snippets tries to solve mainly the common problem of summarization "Show me some representative/typical data". As stated by the original paper, potential uses of snippets are: integrating summarizations of files directly into an operating, production of automatically generated reports, for example, summarize a sleep study and also can be used to support a host of higher-level tasks, including the comparison of massive data collections.

Usage

```
find_snippet(data, s_size, n_snippets = 2L, window_size = s_size/2L)
```

Arguments

data	a matrix or a vector.
s_size	an int. Size of snippet.
n_snippets	an int. Number of snippets to find. (Default is 2).
window_size	an int. The size of the sliding window used to compare the data. Must be smaller than s_size. (Default is s_size / 2).

Details

Motifs vs. snippets: While motifs reward fidelity of conservation, snippets also rewards coverage. Informally, coverage is some measure of how much of the data is explained or represented by a given snippet.

Shapelets vs. snippets: shapelets are defined as subsequences that are maximally representative of a class. Shapelets are supervised, snippets are unsupervised. Shapelets are generally biased to be as short as possible. In contrast, we want snippets to be longer, to intuitively capture the "flavor" of the time series.

Value

Returns the snippet : a list of n_snippets snippets fraction : fraction of each snippet snippetidx : the location of each snippet within time series

References

- Imani S, Madrid F, Ding W, Crouter S, Keogh E. Matrix Profile XIII: Time Series Snippets: A New Primitive for Time Series Data Mining. In: 2018 IEEE International Conference on Data Mining (ICDM). 2018.
- Gharghabi S, Imani S, Bagnall A, Darvishzadeh A, Keogh E. Matrix Profile XII: MPdist: A Novel Time Series Distance Measure to Allow Data Mining in More Challenging Scenarios. In: 2018 IEEE International Conference on Data Mining (ICDM). 2018.

Website: <https://sites.google.com/site/snippetfinder/>

Examples

```
snippets <- find_snippet(mp_fluss_data$walkjogrun$data[1:300], 40, n_snippets = 2)

snippets <- find_snippet(mp_fluss_data$walkjogrun$data, 120, n_snippets = 3)
plot(snippets)
```

floss

Fast Low-cost Online Semantic Segmentation (FLOSS)

Description

Fast Low-cost Online Semantic Segmentation (FLOSS)

Usage

```
floss(
  .mp,
  new_data,
  data_window,
  threshold = 1,
  exclusion_zone = NULL,
  chunk_size = NULL,
  keep_cac = TRUE
)
```

Arguments

<code>.mp</code>	a MatrixProfile object.
<code>new_data</code>	a matrix or vector of new observations.
<code>data_window</code>	an int. Sets the size of the buffer used to keep track of semantic changes.
<code>threshold</code>	a number. (Default is 1). Set the maximum value for evaluating semantic changes. This is data specific. It is advised to check what is 'normal' for your data.
<code>exclusion_zone</code>	if a number will be used instead of embedded value. (Default is NULL).
<code>chunk_size</code>	an int . (Default is NULL). Set the size of new data that will be added to Floss in each iteration if <code>new_data</code> is large. If NULL, the size will be 50. This is not needed if <code>new_data</code> is small, like 1 observation.
<code>keep_cac</code>	a logical. (Default is TRUE). If set to FALSE, the <code>cac_final</code> will contain only values within <code>data_window</code>

Value

Returns the input `.mp` object new names: `cac` the corrected arc count, `cac_final` the combination of `cac` after repeated calls of `floss()`, `floss` with the location of semantic changes and `floss_vals` with the normalized arc count value of the semantic change positions.

References

- Gharghabi S, Ding Y, Yeh C-CM, Kamgar K, Ulanova L, Keogh E. Matrix Profile VIII: Domain Agnostic Online Semantic Segmentation at Superhuman Performance Levels. In: 2017 IEEE International Conference on Data Mining (ICDM). IEEE; 2017. p. 117-26.

Website: <https://sites.google.com/site/onlinesemanticsegmentation/>

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

See Also

Other Semantic Segmentations: [floss_cac\(\)](#), [floss_extract\(\)](#), [fluss_cac\(\)](#), [fluss_extract\(\)](#), [fluss_score\(\)](#), [fluss\(\)](#)

Examples

```

data <- mp_fluss_data$tilt_abp$data[1:1000]
new_data <- mp_fluss_data$tilt_abp$data[1001:1010]
new_data2 <- mp_fluss_data$tilt_abp$data[1011:1020]
w <- 80
mp <- tsmp(data, window_size = w, verbose = 0)
data_window <- 1000
mp <- floss(mp, new_data, data_window)
mp <- floss(mp, new_data2, data_window)

```

floss_cac

*FLOSS - Corrected Arc Counts***Description**

Computes the arc count with edge and 'online' correction (CAC).

Usage

```
floss_cac(.mp, data_window, exclusion_zone = NULL)
```

Arguments

`.mp` a MatrixProfile object.
`data_window` an int. Sets the size of the buffer used to keep track of semantic changes.
`exclusion_zone` if a number will be used instead of embedded value. (Default is NULL).

Details

Original paper suggest using the classic statistical-process-control heuristic to set a threshold where a semantic change may occur in CAC. This may be useful in real-time implementation as we don't know in advance the number of domain changes to look for. Please check original paper (1).

Value

Returns the input `.mp` object a new name `cac` with the corrected arc count and `cac_final` the combination of `cac` after repeated calls of `floss()`.

References

- Gharghabi S, Ding Y, Yeh C-CM, Kamgar K, Ulanova L, Keogh E. Matrix Profile VIII: Domain Agnostic Online Semantic Segmentation at Superhuman Performance Levels. In: 2017 IEEE International Conference on Data Mining (ICDM). IEEE; 2017. p. 117-26.

Website: <https://sites.google.com/site/onlinesemanticsegmentation/>

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

See Also

Other Semantic Segmentations: [floss_extract\(\)](#), [floss\(\)](#), [fluss_cac\(\)](#), [fluss_extract\(\)](#), [fluss_score\(\)](#), [fluss\(\)](#)

Examples

```
data <- mp_fluss_data$tilt_abp$data[1:1000]
new_data <- mp_fluss_data$tilt_abp$data[1001:1010]
w <- 10
mp <- tsmp(data, window_size = w, verbose = 0)
data_window <- 1000
mp <- stmpi_update(mp, new_data, data_window)
mp <- floss_cac(mp, data_window)
```

floss_extract	<i>FLOSS - Extract Segments</i>
---------------	---------------------------------

Description

Extract candidate points of semantic changes.

Usage

```
floss_extract(.mpac, threshold = 1, exclusion_zone = NULL)
```

Arguments

.mpac	a TSMP object of class ArcCount.
threshold	a number. (Default is 1). Set the maximum value for evaluating semantic changes. This is data specific. It is advised to check what is 'normal' for your data.
exclusion_zone	if a number will be used instead of embedded value. (Default is NULL).

Value

Returns the input .mp object a new name floss with the location of semantic changes and floss_vals with the normalized arc count value of the semantic change positions.

References

- Gharghabi S, Ding Y, Yeh C-CM, Kamgar K, Ulanova L, Keogh E. Matrix Profile VIII: Domain Agnostic Online Semantic Segmentation at Superhuman Performance Levels. In: 2017 IEEE International Conference on Data Mining (ICDM). IEEE; 2017. p. 117-26.

Website: <https://sites.google.com/site/onlinesemanticsegmentation/>

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

See Also

Other Semantic Segmentations: [floss_cac\(\)](#), [floss\(\)](#), [fluss_cac\(\)](#), [fluss_extract\(\)](#), [fluss_score\(\)](#), [fluss\(\)](#)

Examples

```
data <- mp_fluss_data$tilt_abp$data[1:1000]
w <- 10
mp <- tsmf(data, window_size = w, verbose = 0)
mp <- fluss_cac(mp)
mp <- fluss_extract(mp, 2)
```

fluss

*Fast Low-cost Unipotent Semantic Segmentation (FLUSS)***Description**

FLUSS is a Domain Agnostic Online Semantic Segmentation that uses the assumption that when few arc are crossing a given index point, means that there is a high probability of semantic change. This function is a wrap to [fluss_cac\(\)](#) and [fluss_extract\(\)](#).

Usage

```
fluss(.mp, num_segments = 1, exclusion_zone = NULL)
```

Arguments

`.mp` a MatrixProfile object.
`num_segments` an int. Number of segments to extract. Based on domain knowledge.
`exclusion_zone` if a number will be used instead of embedded value. (Default is NULL).

Value

Returns the input `.mp` object new names: `cac`, corrected arc count and `fluss` with the location of semantic changes.

References

- Gharghabi S, Ding Y, Yeh C-CM, Kamgar K, Ulanova L, Keogh E. Matrix Profile VIII: Domain Agnostic Online Semantic Segmentation at Superhuman Performance Levels. In: 2017 IEEE International Conference on Data Mining (ICDM). IEEE; 2017. p. 117-26.

Website: <https://sites.google.com/site/onlinesemanticsegmentation/>

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

See Also

Other Semantic Segmentations: [floss_cac\(\)](#), [floss_extract\(\)](#), [floss\(\)](#), [fluss_cac\(\)](#), [fluss_extract\(\)](#), [fluss_score\(\)](#)

Examples

```
data <- mp_fluss_data$tilt_abp$data[1:1000]
w <- 10
mp <- tsmp(data, window_size = w, verbose = 0)
mp <- fluss(mp, 2)
```

fluss_cac	<i>FLUSS - Corrected Arc Counts</i>
-----------	-------------------------------------

Description

Computes the arc count with edge correction (CAC).

Usage

```
fluss_cac(.mp, exclusion_zone = NULL)
```

Arguments

`.mp` a MatrixProfile object.
`exclusion_zone` if a number will be used instead of embedded value. (Default is NULL).

Details

Original paper suggest using the classic statistical-process-control heuristic to set a threshold where a semantic change may occur in CAC. This may be useful in real-time implementation as we don't know in advance the number of domain changes to look for. Please check original paper (1).

Value

Returns the input `.mp` object a new name `cac` with the corrected arc count.

References

- Gharghabi S, Ding Y, Yeh C-CM, Kamgar K, Ulanova L, Keogh E. Matrix Profile VIII: Domain Agnostic Online Semantic Segmentation at Superhuman Performance Levels. In: 2017 IEEE International Conference on Data Mining (ICDM). IEEE; 2017. p. 117-26.

Website: <https://sites.google.com/site/onlinesemanticsegmentation/>

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

See Also

Other Semantic Segmentations: [floss_cac\(\)](#), [floss_extract\(\)](#), [floss\(\)](#), [fluss_extract\(\)](#), [fluss_score\(\)](#), [fluss\(\)](#)

Examples

```
data <- mp_fluss_data$tilt_abp$data[1:1000]
w <- 10
mp <- tsmp(data, window_size = w, verbose = 0)
mp <- fluss_cac(mp)
```

fluss_extract	<i>FLUSS - Extract Segments</i>
---------------	---------------------------------

Description

Extract candidate points of semantic changes.

Usage

```
fluss_extract(.mpac, num_segments = 1, exclusion_zone = NULL)
```

Arguments

`.mpac` a TSMP object of class ArcCount.
`num_segments` an int. Number of segments to extract. Based on domain knowledge.
`exclusion_zone` if a number will be used instead of embedded value. (Default is NULL).

Value

Returns the input `.mp` object a new name `fluss` with the location of semantic changes.

References

- Gharghabi S, Ding Y, Yeh C-CM, Kamgar K, Ulanova L, Keogh E. Matrix Profile VIII: Domain Agnostic Online Semantic Segmentation at Superhuman Performance Levels. In: 2017 IEEE International Conference on Data Mining (ICDM). IEEE; 2017. p. 117-26.

Website: <https://sites.google.com/site/onlinesemanticsegmentation/>

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

See Also

Other Semantic Segmentations: [floss_cac\(\)](#), [floss_extract\(\)](#), [floss\(\)](#), [fluss_cac\(\)](#), [fluss_score\(\)](#), [fluss\(\)](#)

Examples

```
data <- mp_fluss_data$tilt_abp$data[1:1000]
w <- 10
mp <- tsmp(data, window_size = w, verbose = 0)
mp <- fluss_cac(mp)
mp <- fluss_extract(mp, 2)
```

fluss_score	<i>FLUSS - Prediction score calculation</i>
-------------	---

Description

FLUSS - Prediction score calculation

Usage

```
fluss_score(gtruth, extracted, data_size)
```

Arguments

gtruth	an int or vector of int with the ground truth index of segments.
extracted	an int or vector of int with the extracted indexes from <code>fluss_extract()</code> .
data_size	an int. Size of original input data.

Value

Returns the score of predicted semantic transitions compared with the ground truth. Zero is the best, One is the worst.

References

- Gharghabi S, Ding Y, Yeh C-CM, Kamgar K, Ulanova L, Keogh E. Matrix Profile VIII: Domain Agnostic Online Semantic Segmentation at Superhuman Performance Levels. In: 2017 IEEE International Conference on Data Mining (ICDM). IEEE; 2017. p. 117-26.

Website: <https://sites.google.com/site/onlinesemanticsegmentation/>

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

See Also

Other Semantic Segmentations: `floss_cac()`, `floss_extract()`, `floss()`, `fluss_cac()`, `fluss_extract()`, `fluss()`

Examples

```
data <- mp_fluss_data$tilt_abp$data[1:1000]
w <- 10
truth <- c(945, 875)
mp <- tsmp(data, window_size = w, verbose = 0)
mp <- fluss_cac(mp)
mp <- fluss_extract(mp, 2)
score <- fluss_score(truth, mp$fluss, length(data))
```

get_data	<i>Get the data included in a TSMP object, if any.</i>
----------	--

Description

Get the data included in a TSMP object, if any.

Usage

```
get_data(.mp)
```

Arguments

.mp a TSMP object.

Value

Returns the data as matrix. If there is more than one series, returns a list.

Examples

```
mp <- tsmp(mp_toy_data$data[1:200, 1], window_size = 30, verbose = 0)
get_data(mp)
```

mass_v3	<i>Calculates the distance profile using MASS_V3 algorithm</i>
---------	--

Description

Mueen's Algorithm for Similarity Search is The Fastest Similarity Search Algorithm for Time Series Subsequences under Euclidean Distance and Correlation Coefficient.

Usage

```
mass_v3(
  query_window,
  data,
  window_size,
  data_size,
  data_mean,
  data_sd,
  query_mean,
  query_sd,
  k = NULL,
  ...
)
```

Arguments

query_window	a vector of numeric. Query window.
data	a matrix or a vector.
window_size	an int. Sliding window size.
data_size	an int. The length of the reference data.
data_mean	precomputed data moving average.
data_sd	precomputed data moving standard deviation.
query_mean	precomputed query average.
query_sd	precomputed query standard deviation.
k	an int or NULL. Default is NULL. Defines the size of batch. Prefer to use a power of 2.
...	just a placeholder to catch unused parameters.

Details

This is a piecewise version of MASS that performs better when the size of the pieces are well aligned with the hardware.

Value

Returns the distance_profile for the given query and the last_product for STOMP algorithm.

References

- Abdullah Mueen, Yan Zhu, Michael Yeh, Kaveh Kamgar, Krishnamurthy Viswanathan, Chetan Kumar Gupta and Eamonn Keogh (2015), The Fastest Similarity Search Algorithm for Time Series Subsequences under Euclidean Distance

Website: <https://www.cs.unm.edu/~mueen/FastestSimilaritySearch.html>

See Also

[mass_pre\(\)](#) to precomputation of input values.

Examples

```
w <- mp_toy_data$sub_len
ref_data <- mp_toy_data$data[, 1]
query_data <- mp_toy_data$data[, 1]
d_size <- length(ref_data)
q_size <- length(query_data)

pre <- tsmpr::mass_pre(ref_data, query_data, w)

dp <- list()
for (i in 1:(d_size - w + 1)) {
  dp[[i]] <- tsmpr::mass_v3(
    query_data[i:(i - 1 + w)], ref_data,
```

```

        pre$window_size, pre$data_size, pre$data_mean, pre$data_sd,
        pre$query_mean[i], pre$query_sd[i]
    )
}

```

min_mp_idx	<i>Get index of the minimum value from a matrix profile and its nearest neighbor</i>
------------	--

Description

Get index of the minimum value from a matrix profile and its nearest neighbor

Usage

```
min_mp_idx(.mp, n_dim = NULL, valid = TRUE)
```

Arguments

.mp	a MatrixProfile object.
n_dim	number of dimensions of the matrix profile
valid	check for valid numbers

Value

returns a matrix with two columns: the minimum and the nearest neighbor

Examples

```

w <- 50
data <- mp_gait_data
mp <- tsmp(data, window_size = w, exclusion_zone = 1 / 4, verbose = 0)
min_val <- min_mp_idx(mp)

```

motifs	<i>Search for Motifs</i>
--------	--------------------------

Description

Search for Motifs

Usage

```
motifs(  
  profile,  
  exclusion_zone = profile$ez,  
  k = 3L,  
  neighbor_count = 10L,  
  radius = 3  
)
```

Arguments

profile a MatrixProfile or PMP object.

exclusion_zone an int. Number of values to exclude on both sides of the motif to avoid trivial matches. Defaults to the exclusion zone used to compute the (Pan-)Matrix Profile which is found in the profile data structure.

k an int. Number of motifs to find. (Default is 3).

neighbor_count an int. Number of neighbors to find. (Default is 3).

radius an int. Set a threshold to exclude matching neighbors with distance > current discord distance * radius. (Default is 3).

References

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

See Also

Other Main API: [analyze\(\)](#), [compute\(\)](#), [discords\(\)](#), [visualize\(\)](#)

`motifs_discords_small` *Just a synthetic dataset for testing*

Description

Just a synthetic dataset for testing

Usage

```
motifs_discords_small
```

Format

A vector with 875 observations

mpdist

MPdist - Distance between Time Series using Matrix Profile

Description

MPdist is a recently introduced distance measure which considers two time series to be similar if they share many similar subsequences, regardless of the order of matching subsequences. It was demonstrated in that MPdist is robust to spikes, warping, linear trends, dropouts, wandering baseline and missing values, issues that are common outside of benchmark datasets.

Usage

```
mpdist(  
  ref_data,  
  query_data,  
  window_size,  
  type = c("simple", "vector"),  
  thr = 0.05  
)
```

Arguments

ref_data	a matrix or a vector. The reference data
query_data	a matrix or a vector. The query data
window_size	an int. Size of the sliding window.
type	the type of result. (Default is simple). See details.
thr	threshold for MPdist. (Default is 0.05). Don't change this unless you know what you are doing.

Details

MPdist returns the distance of two time series or a vector containing the distance between all sliding windows. If argument type is set to vector, the vector is returned.

Value

Returns the distance of two time series or a vector containing the distance between all sliding windows.

References

- Gharghabi S, Imani S, Bagnall A, Darvishzadeh A, Keogh E. Matrix Profile XII: MPdist: A Novel Time Series Distance Measure to Allow Data Mining in More Challenging Scenarios. In: 2018 IEEE International Conference on Data Mining (ICDM). 2018.

Website: <https://sites.google.com/site/mpdistinfo/>

Examples

```

ref_data <- mp_toy_data$data[, 1]
qe_data <- mp_toy_data$data[, 2]
qd_data <- mp_toy_data$data[150:200, 1]
w <- mp_toy_data$sub_len

# distance between data of same size
deq <- mpdist(ref_data, qe_data, w)

# distance between data of different sizes
ddiff <- mpdist(ref_data, qd_data, w)

# distance vector between data of different sizes
ddvect <- mpdist(ref_data, qd_data, w, type = "vector")

```

mpx	<i>Fast implementation of MP and MPI for internal purposes, without FFT</i>
-----	---

Description

Fast implementation of MP and MPI for internal purposes, without FFT

Usage

```

mpx(
  data,
  window_size,
  query = NULL,
  idx = TRUE,
  dist = c("euclidean", "pearson"),
  n_workers = 1
)

```

Arguments

data	a matrix or a vector. The time series to analyze.
window_size	window size
query	query
idx	compute the profile indexes?
dist	distance measure, Euclidean or Pearson?
n_workers	threads for multi-threading

Value

Returns MP and MPI

Examples

```
mp <- mpx(mp_toy_data$data[1:200, 1], window_size = 30)
```

mp_fluss_data

Original data used in the FLUSS paper

Description

Contains two datasets used in FLUSS paper (1), first is TiltABP from (2), and second is WalkJogRun from PAMAP's dataset (3)

Usage

```
mp_fluss_data
```

Format

A list containing:

data one column matrix with the dataset's data

gtruth a vector with the ground truth of semantic change according to provided dataset

window window size used in original paper

Source

<https://sites.google.com/site/onlinesemanticsegmentation/>

http://www.cs.ucr.edu/~eamonn/time_series_data/

References

- Gharghabi S, Ding Y, Yeh C-CM, Kamgar K, Ulanova L, Keogh E. Matrix Profile VIII: Domain Agnostic Online Semantic Segmentation at Superhuman Performance Levels. In: 2017 IEEE International Conference on Data Mining (ICDM). IEEE; 2017. p. 117-26.
- Heldt, T., Oefinger, M.B., Hoshiyama, M. and Mark, R.G., 2003, September. Circulatory response to passive and active changes in posture. In IEEE Computers in Cardiology, 2003 (pp. 263-266).
- Reiss, A. and Stricker, D., 2012. Introducing a new benchmarked dataset for activity monitoring. In 16th International Symposium on Wearable Computers (ISWC), 2012, pages 108-109. IEEE, 2012.

mp_gait_data

Original data used in the Time Series Chain demo

Description

Original data used in the Time Series Chain demo

Usage

mp_gait_data

Format

A matrix with 904 rows and 1 column with the Y data from an accelerometer

Source

<https://sites.google.com/site/timeserieschain/>

References

- Zhu Y, Imamura M, Nikovski D, Keogh E. Matrix Profile VII: Time Series Chains: A New Primitive for Time Series Data Mining. Knowl Inf Syst. 2018 Jun 2;1-27.

mp_meat_data

Original data used in the Salient Subsequences demo

Description

This is the Meat dataset from UCR Archive modified for Salient discovery. The original data is mixed with Random Walks and the algorithm must pick only the originals.

Usage

mp_meat_data

Format

original is the original dataset with 60+60 observations mixed with 120 random walks:

data 240 time series with length of 448 each.

labels label of each time series, -666 means a random walk.

sub_len size of sliding window.

sub is the original dataset embedded in random walks:

data One time series with length of 107520.

labels label of each original data.

labels_idx starting point where the original data was placed.

sub_len size of sliding window.

Source

http://www.cs.ucr.edu/~eamonn/time_series_data/

References

- Yeh CCM, Van Herle H, Keogh E. Matrix profile III: The matrix profile allows visualization of salient subsequences in massive time series. Proc - IEEE Int Conf Data Mining, ICDM. 2017;579-88.
- Hu B, Rakthanmanon T, Hao Y, Evans S, Lonardi S, Keogh E. Discovering the Intrinsic Cardinality and Dimensionality of Time Series Using MDL. In: 2011 IEEE 11th International Conference on Data Mining. IEEE; 2011. p. 1086-91.

Website: <https://sites.google.com/site/salientsubs/>

mp_test_data

Original data used in the STDS demo

Description

A synthetic dataset base on TRACE dataset and used as Stress Test to STDS algorithm. The TRACE dataset used here is originally from (1), and the version distributed here is from (2)

Usage

mp_test_data

Format

A list of matrices with 215010 rows and 1 dimension:

train\$data training data

train\$label label for training data

test\$data test data

test\$label label for test data

Source

<https://sites.google.com/view/weaklylabeled>

http://www.cs.ucr.edu/~eamonn/time_series_data/

References

- Roverso, D., Multivariate temporal classification by windowed wavelet decomposition and recurrent neural networks, in 3rd ANS Int'l Topical Meeting on Nuclear Plant Instrumentation, Control and Human-Machine Interface, vol. 20, Washington, DC, USA, 2000.
- Yeh C-CM, Kavantzias N, Keogh E. Matrix profile IV: Using Weakly Labeled Time Series to Predict Outcomes. Proc VLDB Endow. 2017 Aug 1;10(12):1802-12.

mp_toy_data

Original data used in the mSTAMP demo

Description

A synthetic dataset with embedded MOTIFS for multidimensional discovery

Usage

mp_toy_data

Format

A list with a matrix with 550 rows and 3 dimensions and an int:

data data with embedded MOTIFS

sub_len size of sliding window

Source

<https://sites.google.com/view/mstamp/>

References

- Yeh CM, Kavantzias N, Keogh E. Matrix Profile VI : Meaningful Multidimensional Motif Discovery.

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

mstomp_par

Multivariate STOMP algorithm Parallel version

Description

Computes the Matrix Profile and Profile Index for Multivariate Time Series.

Usage

```
mstomp_par(
  data,
  window_size,
  exclusion_zone = getOption("tsmp.exclusion_zone", 1/2),
  verbose = getOption("tsmp.verbose", 2),
  must_dim = NULL,
  exc_dim = NULL,
  n_workers = 2
)
```

```
mstomp(
  data,
  window_size,
  exclusion_zone = getOption("tsmp.exclusion_zone", 1/2),
  verbose = getOption("tsmp.verbose", 2),
  must_dim = NULL,
  exc_dim = NULL
)
```

Arguments

data	a matrix of numeric, where each column is a time series. Accepts vector (see details), list and data.frame too.
window_size	an int with the size of the sliding window.
exclusion_zone	a numeric. Size of the exclusion zone, based on window size (default is 1/2).
verbose	an int. See details. (Default is 2).
must_dim	an int or vector of which dimensions to forcibly include (default is NULL).
exc_dim	an int or vector of which dimensions to exclude (default is NULL).
n_workers	an int. Number of workers for parallel. (Default is 2).

Details

The Matrix Profile, has the potential to revolutionize time series data mining because of its generality, versatility, simplicity and scalability. In particular it has implications for time series motif discovery, time series joins, shapelet discovery (classification), density estimation, semantic segmentation, visualization, rule discovery, clustering etc. The MSTOMP computes the Matrix Profile

and Profile Index for Multivariate Time Series that is meaningful for multidimensional MOTIF discovery. It uses the STOMP algorithm that is faster than STAMP but lacks its anytime property.

Although this functions handles Multivariate Time Series, it can also be used to handle Univariate Time Series. `verbose` changes how much information is printed by this function; 0 means nothing, 1 means text, 2 adds the progress bar, 3 adds the finish sound.

Value

Returns a `MultiMatrixProfile` object, a list with the matrix profile `mp`, profile index `pi` left and right matrix profile `lmp`, `rmp` and profile index `lpi`, `rpi`, window size `w`, number of dimensions `n_dim`, exclusion zone `ez`, must dimensions `must` and excluded dimensions `exc`.

If the input has only one dimension, returns the same as `stomp()`.

Functions

- `mstomp_par()`: Parallel version.
- `mstomp()`: Single thread version.

References

- Yeh CM, Kavantzias N, Keogh E. Matrix Profile VI : Meaningful Multidimensional Motif Discovery.
- Zhu Y, Imamura M, Nikovski D, Keogh E. Matrix Profile VII: Time Series Chains: A New Primitive for Time Series Data Mining. *Knowl Inf Syst.* 2018 Jun 2;1-27.

Website: <https://sites.google.com/view/mstamp/>

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

See Also

Other matrix profile computations: `scrimp()`, `stamp_par()`, `stomp_par()`, `tsmp()`, `valmod()`

Examples

```
# using all dimensions
mp <- mstomp(mp_toy_data$data[1:150, ], 30, verbose = 0)

#' # using threads
mp <- mstomp_par(mp_toy_data$data[1:150, ], 30, verbose = 0)

# force using dimensions 1 and 2
mp <- mstomp(mp_toy_data$data[1:200, ], 30, must_dim = c(1, 2))
# exclude dimensions 2 and 3
mp2 <- mstomp(mp_toy_data$data[1:200, ], 30, exc_dim = c(2, 3))
```

plot *Plot a TSMP object*

Description

Plot a TSMP object

Usage

```
## S3 method for class 'ArcCount'
plot(
  x,
  data,
  type = c("data", "matrix"),
  exclusion_zone = NULL,
  edge_limit = NULL,
  threshold = stats::quantile(x$cac, 0.1),
  main = "Arcs Discover",
  xlab = "index",
  ylab = "",
  ...
)

## S3 method for class 'Valmod'
plot(
  x,
  ylab = "distance",
  xlab = "index",
  main = "Valmod Matrix Profile",
  data = FALSE,
  ...
)

## S3 method for class 'MatrixProfile'
plot(
  x,
  ylab = "distance",
  xlab = "index",
  main = "Unidimensional Matrix Profile",
  data = FALSE,
  ...
)

## S3 method for class 'MultiMatrixProfile'
plot(
  x,
  ylab = "distance",
```

```
    xlab = "index",
    main = "Multidimensional Matrix Profile",
    ...
)

## S3 method for class 'SimpleMatrixProfile'
plot(
  x,
  ylab = "distance",
  xlab = "index",
  main = "SiMPle Matrix Profile",
  data = FALSE,
  ...
)

## S3 method for class 'Fluss'
plot(
  x,
  data,
  type = c("data", "matrix"),
  main = "Fast Low-cost Unipotent Semantic Segmentation",
  xlab = "index",
  ylab = "",
  ...
)

## S3 method for class 'Floss'
plot(
  x,
  data,
  type = c("data", "matrix"),
  main = "Fast Low-cost Online Semantic Segmentation",
  xlab = "index",
  ylab = "",
  ...
)

## S3 method for class 'Chain'
plot(
  x,
  data,
  type = c("data", "matrix"),
  main = "Chain Discover",
  xlab = "index",
  ylab = "",
  ...
)
```

```
## S3 method for class 'Discord'
plot(
  x,
  data,
  type = c("data", "matrix"),
  ncol = 3,
  main = "Discord Discover",
  xlab = "index",
  ylab = "",
  ...
)

## S3 method for class 'Snippet'
plot(
  x,
  data,
  ncol = 3,
  main = "Snippet Finder",
  xlab = "index",
  ylab = "",
  ...
)

## S3 method for class 'Motif'
plot(
  x,
  data,
  type = c("data", "matrix"),
  ncol = 3,
  main = "MOTIF Discover",
  xlab = "index",
  ylab = "",
  ...
)

## S3 method for class 'MultiMotif'
plot(
  x,
  data,
  type = c("data", "matrix"),
  ncol = 3,
  main = "Multidimensional MOTIF Discover",
  xlab = "index",
  ylab = "",
  ...
)

## S3 method for class 'Salient'
```

```

plot(x, data, main = "Salient Subsections", xlab = "index", ylab = "", ...)

## S3 method for class 'PMP'
plot(
  x,
  ylab = "distance",
  xlab = "index",
  main = "Unidimensional Matrix Profile",
  data = FALSE,
  ...
)

```

Arguments

x	a Matrix Profile
data	the data used to build the Matrix Profile, if not embedded to it.
type	"data" or "matrix". Choose what will be plotted.
exclusion_zone	if a number will be used instead of Matrix Profile's. (Default is NULL).
edge_limit	if a number will be used instead of Matrix Profile's exclusion zone. (Default is NULL).
threshold	the maximum value to be used to plot.
main	a string. Main title.
xlab	a string. X label.
ylab	a string. Y label.
...	further arguments to be passed to <code>plot()</code> . See <code>par()</code> .
ncol	an int. Number of columns to plot Motifs.

Value

None

Examples

```

mp <- tsmplot(mp_toy_data$data[1:200, 1], window_size = 30, verbose = 0)
plot(mp)

```

plot_arcs

Plot arcs between indexes of a Profile Index

Description

Sometimes may be useful to see where is the nearest neighbor graphically. This is the reasoning behind, for example, FLUSS which uses the arc count to infer a semantic change, and SIMPLE which infer that arcs connect similar segments of a music. See details for a deeper explanation how to use this function.

Usage

```
plot_arcs(
  pairs,
  alpha = NULL,
  quality = 30,
  lwd = 15,
  col = c("blue", "orange"),
  main = "Arc Plot",
  ylab = "",
  xlab = "Profile Index",
  xmin = NULL,
  xmax = NULL,
  ...
)
```

Arguments

<code>pairs</code>	a matrix with 2 columns.
<code>alpha</code>	a numeric. (Default is NULL, automatic). Alpha value for lines transparency.
<code>quality</code>	an int. (Default is 30). Number of segments to draw the arc. Bigger value, harder to render.
<code>lwd</code>	an int. (Default is 15). Line width.
<code>col</code>	a vector of colors. (Default is c("blue", "orange")). Colors for right and left arc, respectively. Accepts one color.
<code>main</code>	a string. (Default is "Arc Plot"). Main title.
<code>ylab</code>	a string. (Default is ""). Y label.
<code>xlab</code>	a string. (Default is "Profile Index"). X label.
<code>xmin</code>	an int. (Default is NULL). Set the minimum value of x axis.
<code>xmax</code>	an int. (Default is NULL). Set the maximum value of x axis.
<code>...</code>	further arguments to be passed to <code>plot()</code> . See <code>par()</code> .

Details

You have two options to use this function. First you can provide just the data, and the function will try its best to retrieve the pairs for plotting. Second, you can skip the first parameters and just provide the `pairs`, which is a matrix with two columns; the first is the starting index, the second is the end index. Two colors are used to allow you to identify the direction of the arc. If you use the `rpi` or `lpi` as input, you will see that these profile indexes have just one direction.

`exclusion_zone` is used to filter out small arcs that may be useless (e.g. you may be interested in similarities that are far away). `edge_limit` is used to filter out spurious arcs that are used connect the beginning and the end of the profile (e.g. silent audio). `threshold` is used to filter indexes that have distant nearest neighbor (e.g. retrieve only the best motifs).

Value

None

Examples

```
plot_arcs(pairs = matrix(c(5, 10, 1, 10, 20, 5), ncol = 2, byrow = TRUE))
```

pmp

*Pan-Matrix Profile***Description**

Computes the Pan-Matrix Profile (PMP) for the given time series.

Usage

```
pmp(
  data,
  window_sizes = seq.int(from = 10, to = length(data)/2, length.out = 20),
  plot = FALSE,
  pmp_obj = NULL,
  n_workers = 1,
  verbose = getOption("tsmp.verbose", 2)
)
```

Arguments

<code>data</code>	a matrix or a vector of numeric.
<code>window_sizes</code>	a vector of the window sizes that will be evaluated. They will be rounded to the lower integer and sorted. (Default is a sequence of 20 values from 10 to half data size).
<code>plot</code>	a logical. If TRUE, every new computation will be plotted. (Default is FALSE).
<code>pmp_obj</code>	a PMP object that may or not contain an upper bound value, and previous computed profiles. The function will add new profiles, not replace. (Default is NULL).
<code>n_workers</code>	an int. Number of workers for parallel. (Default is 1).
<code>verbose</code>	an int. See details. (Default is 2).

Details

The work closest in spirit to ours is VALMOD. The idea of VALMOD is to compute the MP for the shortest length of interest, then use the information gleaned from it to guide a search through longer subsequence lengths, exploiting lower bounds to prune off some calculations. This idea works well for the first few of the longer subsequence lengths, but the lower bounds progressively weaken, making the pruning ineffective. Thus, in the five case studies they presented, the mean value of U/L was just 1.24. In contrast, consider that our termite example in Fig. 15 has a U/L ratio of 240, more than two orders of magnitude larger. Thus, VALMOD is perhaps best seen as finding motifs with some tolerance for a slightly (~25%) too short user-specified query length, rather than a true "motif-of-all-lengths" algorithm. Also note that apart from the shortest length, VALMOD only gives some

information for the other lengths, unlike pmp, which contains exact distances for all subsequences of all lengths.

When just the data is provided, the exploration will be done using the default `window_sizes` that is a sequence of 20 values between 10 and the half data size and the resulting object will have an `upper_bound` equals to `Inf`. If an object is provided by the argument `pmp_obj`, this function will add more information to the resulting object, never changing the values already computed. `verbose` changes how much information is printed by this function; 0 means nothing, 1 means text, 2 adds the progress bar, 3 adds the finish sound.

Talk about upper bound and window sizes

1. `upper_window` will be set to `Inf` on new objects 1.1. `upper_window` will also be used for plot, and for discovery, it must not remove any existing data from the object
2. `window_sizes` is used for plot, it must not remove any mp inside the object 2.1. `window_sizes` tells the function what mp are stored, it may be updated with `as.numeric(names(pmp))`
3. the functions must be capable to handle the data without need to sort by `window_size`, but sort may be useful later(?)

Value

Returns a PMP object.

Examples

```
# Just compute
pan <- pmp(mp_gait_data)
# Compute the upper bound, than add new profiles
pan <- pmp_upper_bound(mp_gait_data)
pan <- pmp(mp_gait_data, pmp_obj = pan)
```

<code>pmp_upper_bound</code>	<i>Pan Matrix Profile upper bound</i>
------------------------------	---------------------------------------

Description

Finds the upper bound for Pan Matrix Profile calculation.

Usage

```
pmp_upper_bound(
  data,
  threshold = getOption("tsmp.pmp_ub", 0.95),
  refine_stepsize = getOption("tsmp.pmp_refine", 0.25),
  return_pmp = TRUE,
  n_workers = 1,
  verbose = getOption("tsmp.verbose", 2)
)
```


Arguments

data	a matrix or a vector of numeric.
threshold	a numeric. Correlation threshold. See details. (Default is 0.95).
refine_stepsize	a numeric. Step size for the last upper bound search. See details. (Default is 0.25).
return_pmp	a logical. If TRUE, returns the computed data as a PMP object, if FALSE, returns just the upper bound value. (Default is TRUE).
n_workers	an int. Number of workers for parallel. (Default is 1).
verbose	verbose an int. See details. (Default is 2).

Details

The Pan Matrix Profile may not give any further information beyond a certain window size. This function starts computing the matrix profile for the window size of 8 and doubles it until the minimum correlation value found is less than the threshold. After that, it begins to refine the upper bound using the refine_stepsize values, until the threshold value is hit.

verbose changes how much information is printed by this function; 0 means nothing, 1 means text, 2 adds the progress bar, 3 adds the finish sound.

Value

Returns a PMP object with computed data, or just the upper bound value if return_pmp is set to FALSE.

References

- Yet to be announced

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

Examples

```
# return the object
pan_matrix <- pmp_upper_bound(mp_gait_data)

# just the upper bound
pan_ub <- pmp_upper_bound(mp_gait_data, return_pmp = FALSE)
```

read	<i>Read TSMP object from JSON file.</i>
------	---

Description

Read TSMP object from JSON file.

Usage

```
read(x, ...)
```

Arguments

x	a character string with the input filename.
...	other arguments to be passed forward.

Examples

```
result <- compute(mp_toy_data$data[, 1], 80)

tempfile <- file.path(tempdir(), "output.json")
write(result, file = tempfile)
result <- read(tempfile)
```

remove_class	<i>Remove a TSMP class from an object</i>
--------------	---

Description

Remove a TSMP class from an object

Usage

```
remove_class(x, class)
```

Arguments

x	a TSMP object
class	character string with the class name

Value

the object without the class

Examples

```
w <- 50
data <- mp_gait_data
mp <- tsmp(data, window_size = w, exclusion_zone = 1 / 4, verbose = 0)
mp <- find_chains(mp)
# Remove the "Chain" class information
mp <- remove_class(mp, "Chain")
```

salient_mds *Convert salient sequences into MDS space*

Description

Convert salient sequences into MDS space

Usage

```
salient_mds(.mp, data, bit_idx = 1)
```

Arguments

.mp	a Matrix Profile object.
data	the data used to build the Matrix Profile, if not embedded.
bit_idx	an int. The index of n_bits used for MDL discretization if more than one was used. (Default is 1).

Value

Returns X,Y values for plotting

References

- Yeh CCM, Van Herle H, Keogh E. Matrix profile III: The matrix profile allows visualization of salient subsequences in massive time series. Proc - IEEE Int Conf Data Mining, ICDM. 2017;579-88.
- Hu B, Rakthanmanon T, Hao Y, Evans S, Lonardi S, Keogh E. Discovering the Intrinsic Cardinality and Dimensionality of Time Series Using MDL. In: 2011 IEEE 11th International Conference on Data Mining. IEEE; 2011. p. 1086-91.

Website: <https://sites.google.com/site/salientsubs/>

Examples

```
# toy example
data <- mp_toy_data$data[, 1]
mp <- tsmp(data, window_size = 30, verbose = 0)
mps <- salient_subsequences(mp, verbose = 0)
mds_data <- salient_mds(mps)
plot(mds_data, main = "Multi dimensional scale")
```

salient_score	<i>Computes the F-Score of salient algorithm.</i>
---------------	---

Description

This score function is useful for testing several values of `n_bits` for MDL discretization and checking against a known set of indexes. This increase the probability of better results on relevant subsequence extraction.

Usage

```
salient_score(.mp, gtruth, verbose = getOption("tsmp.verbose", 2))
```

Arguments

<code>.mp</code>	a Matrix Profile object.
<code>gtruth</code>	a vector of integers with the indexes of relevant subsequences.
<code>verbose</code>	an int. (Default is 2).

Value

Returns a list with `f_score`, `precision`, `recall` and `bits` used in the algorithm.

References

- Yeh CCM, Van Herle H, Keogh E. Matrix profile III: The matrix profile allows visualization of salient subsequences in massive time series. Proc - IEEE Int Conf Data Mining, ICDM. 2017;579-88.
- Hu B, Rakthanmanon T, Hao Y, Evans S, Lonardi S, Keogh E. Discovering the Intrinsic Cardinality and Dimensionality of Time Series Using MDL. In: 2011 IEEE 11th International Conference on Data Mining. IEEE; 2011. p. 1086-91.

Website: <https://sites.google.com/site/salientsubs/>

Examples

```
# toy example
data <- mp_toy_data$data[, 1]
mp <- tsmp(data, window_size = 30, verbose = 0)
mps <- salient_subsequences(mp, n_bits = c(4, 6, 8), verbose = 0)
label_idx <- seq(2, 500, by = 110) # fake data
salient_score(mps, label_idx, verbose = 0)
```

salient_subsequences *Framework for retrieve salient subsequences from a dataset*

Description

In order to allow a meaningful visualization in Multi-Dimensional Space (MDS), this function retrieves the most relevant subsequences using Minimal Description Length (MDL) framework.

Usage

```
salient_subsequences(  
  .mp,  
  data,  
  n_bits = 8,  
  n_cand = 10,  
  exclusion_zone = NULL,  
  verbose = getopt("tsmp.verbose", 2)  
)
```

Arguments

<code>.mp</code>	a TSMP object of class <code>MatrixProfile</code> .
<code>data</code>	the data used to build the Matrix Profile, if not embedded.
<code>n_bits</code>	an int or vector of int. Number of bits for MDL discretization. (Default is 8).
<code>n_cand</code>	an int. number of candidate when picking the subsequence in each iteration. (Default is 10).
<code>exclusion_zone</code>	if a number will be used instead of embedded value. (Default is NULL).
<code>verbose</code>	an int. See details. (Default is 2).

Details

`verbose` changes how much information is printed by this function; 0 means nothing, 1 means text, 2 adds the progress bar, 3 adds the finish sound.

Value

Returns the input `.mp` object with a new name `salient`. It contains: `indexes`, a vector with the starting position of each subsequence, `idx_bit_size`, a vector with the associated bitsize for each iteration and `bits` the value used as input on `n_bits`.

References

- Yeh CCM, Van Herle H, Keogh E. Matrix profile III: The matrix profile allows visualization of salient subsequences in massive time series. Proc - IEEE Int Conf Data Mining, ICDM. 2017;579-88.
- Hu B, Rakthanmanon T, Hao Y, Evans S, Lonardi S, Keogh E. Discovering the Intrinsic Cardinality and Dimensionality of Time Series Using MDL. In: 2011 IEEE 11th International Conference on Data Mining. IEEE; 2011. p. 1086-91.

Website: <https://sites.google.com/site/salientsubs/>

Examples

```
# toy example
data <- mp_toy_data$data[, 1]
mp <- tsmp(data, window_size = 30, verbose = 0)
mps <- salient_subsequences(mp, data, verbose = 0)

# full example
data <- mp_meat_data$sub$data
w <- mp_meat_data$sub$sub_len
mp <- tsmp(data, window_size = w, verbose = 2, n_workers = 2)
mps <- salient_subsequences(mp, data, n_bits = c(4, 6, 8), verbose = 2)
```

scrimp

Anytime univariate SCRIMP++ algorithm

Description

Computes the best so far Matrix Profile and Profile Index for Univariate Time Series. **DISCLAIMER:** This algorithm still in development by its authors. Join similarity, RMP and LMP not implemented yet.

Usage

```
scrimp(
  ...,
  window_size,
  exclusion_zone = getOption("tsmp.exclusion_zone", 1/2),
  verbose = getOption("tsmp.verbose", 2),
  s_size = Inf,
  pre_scrimp = 1/4,
  pre_only = FALSE
)
```

Arguments

...	a matrix or a vector.
window_size	an int. Size of the sliding window.
exclusion_zone	a numeric. Size of the exclusion zone, based on window size (default is 1/2). See details.
verbose	an int. See details. (Default is 2).
s_size	a numeric. for anytime algorithm, represents the size (in observations) the random calculation will occur (default is Inf).
pre_scrimp	a numeric. Set the pre-scrimp step based on window_size, if 0, disables pre-scrimp. (default is 1/4).
pre_only	a logical. Returns only the pre script data. (Default is FALSE).

Details

The Matrix Profile, has the potential to revolutionize time series data mining because of its generality, versatility, simplicity and scalability. In particular it has implications for time series motif discovery, time series joins, shapelet discovery (classification), density estimation, semantic segmentation, visualization, rule discovery, clustering etc. The anytime SCRIMP computes the Matrix Profile and Profile Index in such manner that it can be stopped before its complete calculation and return the best so far results allowing ultra-fast approximate solutions. `verbose` changes how much information is printed by this function; 0 means nothing, 1 means text, 2 adds the progress bar, 3 adds the finish sound. `exclusion_zone` is used to avoid trivial matches.

Value

Returns a MatrixProfile object, a list with the matrix profile `mp`, profile index `pi` left and right matrix profile `lmp`, `rmp` and profile index `lpi`, `rpi`, window size `w` and exclusion zone `ez`.

References

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

See Also

Other matrix profile computations: `mstomp_par()`, `stamp_par()`, `stomp_par()`, `tsmp()`, `valmod()`

Examples

```
mp <- scrimp(mp_toy_data$data[1:200, 1], window_size = 30, verbose = 0)

ref_data <- mp_toy_data$data[, 1]
query_data <- mp_toy_data$data[, 2]
# self similarity
mp <- scrimp(ref_data, window_size = 30, s_size = round(nrow(ref_data) * 0.1))
# join similarity
mp <- scrimp(ref_data, query_data, window_size = 30, s_size = round(nrow(query_data) * 0.1))
```

sdts_predict	<i>Framework for Scalable Dictionary learning for Time Series (SDTS) prediction function</i>
--------------	--

Description

This function trains a model that uses a dictionary to predict state changes. Differently from `fluss()`, it doesn't look for semantic changes (that may be several), but for binary states like "on" or "off". Think for example that a human annotator is pressing a switch any time he thinks that the recorded data is relevant, and releases the switch when he thinks the data is noise. This algorithm will learn the switching points (even better) and try to predict using new data.

Usage

```
sdts_predict(model, data, window_size)
```

Arguments

model	a model created by SDTS training function <code>sdts_train()</code> .
data	a vector of numeric. Time series.
window_size	an int. The average sliding window size.

Value

Returns a vector of logical with predicted annotations.

References

- Yeh C-CM, Kavantzaz N, Keogh E. Matrix profile IV: Using Weakly Labeled Time Series to Predict Outcomes. Proc VLDB Endow. 2017 Aug 1;10(12):1802-12.

Website: <https://sites.google.com/view/weaklylabeled>

See Also

Other Scalable Dictionaries: `sdts_score()`, `sdts_train()`

Examples

```
# This is a fast toy example and results are useless. For a complete result, run the code inside
#' Not run' section below.
w <- c(110, 220)
subs <- 11000:20000
tr_data <- mp_test_data$train$data[subs]
tr_label <- mp_test_data$train$label[subs]
te_data <- mp_test_data$test$data[subs]
te_label <- mp_test_data$test$label[subs]
model <- sdts_train(tr_data, tr_label, w, verbose = 0)
predict <- sdts_predict(model, te_data, round(mean(w)))
```



```
sdts_score(predict, te_label, 1)

windows <- c(110, 220, 330)
model <- sdts_train(mp_test_data$train$data, mp_test_data$train$label, windows, verbose = 0)
predict <- sdts_predict(model, mp_test_data$test$data, round(mean(windows)))
sdts_score(predict, mp_test_data$test$label, 1)
```

sdts_score

Computes the F-Score of a SDTS prediction

Description

Computes the F-Score of a SDTS prediction.

Usage

```
sdts_score(pred, gtruth, beta = 1)
```

Arguments

pred	a vector of logical. Predicted annotation from sdts_predict()
gtruth	a vector of logical. Ground truth annotation.
beta	a numeric. See details. (default is 1).

Details

beta is used to balance F-score towards recall (>1) or precision (<1).

Value

Returns a list with `f_score`, `precision` and `recall`.

References

- Yeh C-CM, Kavantzias N, Keogh E. Matrix profile IV: Using Weakly Labeled Time Series to Predict Outcomes. Proc VLDB Endow. 2017 Aug 1;10(12):1802-12.

Website: <https://sites.google.com/view/weaklylabeled>

See Also

Other Scalable Dictionaries: [sdts_predict\(\)](#), [sdts_train\(\)](#)

Examples

```

# This is a fast toy example and results are useless. For a complete result, run the code inside
#' Not run' section below.
w <- c(110, 220)
subs <- 11000:20000
tr_data <- mp_test_data$train$data[subs]
tr_label <- mp_test_data$train$label[subs]
te_data <- mp_test_data$test$data[subs]
te_label <- mp_test_data$test$label[subs]
model <- sdts_train(tr_data, tr_label, w, verbose = 0)
predict <- sdts_predict(model, te_data, round(mean(w)))
sdts_score(predict, te_label, 1)

windows <- c(110, 220, 330)
model <- sdts_train(mp_test_data$train$data, mp_test_data$train$label, windows)
predict <- sdts_predict(model, mp_test_data$test$data, round(mean(windows)))
sdts_score(predict, mp_test_data$test$label, 1)

```

sdts_train

*Framework for Scalable Dictionary learning for Time Series (SDTS)
training function*

Description

This function trains a model that uses a dictionary to predict state changes. Differently from `fluss()`, it doesn't look for semantic changes (that may be several), but for binary states like "on" or "off". Think for example that a human annotator is pressing a switch any time he thinks that the recorded data is relevant, and releases the switch when he thinks the data is noise. This algorithm will learn the switching points (even better) and try to predict using new data.

Usage

```

sdts_train(
  data,
  label,
  window_size,
  beta = 1,
  pat_max = Inf,
  parallel = FALSE,
  verbose = getOption("tsmp.verbose", 2)
)

```

Arguments

`data` a vector of numeric. Time series.
`label` a vector of logical. Annotations.

window_size	an int or a vector of int. Sliding window sizes.
beta	a numeric. See details. (default is 1).
pat_max	an int. Max number of shape features captured. (default is Inf).
parallel	a logical. Use parallel computation inside (default is FALSE).
verbose	an int. See details. (Default is 2).

Details

beta is used to balance F-score towards recall (>1) or precision (<1). verbose changes how much information is printed by this function; 0 means nothing, 1 means text, 2 adds the progress bar, 3 adds the finish sound.

Value

Returns a list with the learned dictionary score (estimated score), score_hist (history of scores), pattern (shape features), thold (threshold values).

References

- Yeh C-CM, Kavantzias N, Keogh E. Matrix profile IV: Using Weakly Labeled Time Series to Predict Outcomes. Proc VLDB Endow. 2017 Aug 1;10(12):1802-12.

Website: <https://sites.google.com/view/weaklylabeled>

See Also

Other Scalable Dictionaries: [sdts_predict\(\)](#), [sdts_score\(\)](#)

Examples

```
# This is a fast toy example and results are useless. For a complete result, run the code inside
#' Not run' section below.
w <- c(110, 220)
subs <- 11000:20000
tr_data <- mp_test_data$train$data[subs]
tr_label <- mp_test_data$train$label[subs]
te_data <- mp_test_data$test$data[subs]
te_label <- mp_test_data$test$label[subs]
model <- sdts_train(tr_data, tr_label, w, verbose = 0)
predict <- sdts_predict(model, te_data, round(mean(w)))
sdts_score(predict, te_label, 1)

windows <- c(110, 220, 330)
model <- sdts_train(mp_test_data$train$data, mp_test_data$train$label, windows)
predict <- sdts_predict(model, mp_test_data$test$data, round(mean(windows)))
sdts_score(predict, mp_test_data$test$label, 1)
```

set_data	<i>Set/changes the data included in TSMP object.</i>
----------	--

Description

This may be useful if you want to include the data lately or remove the included data (set as NULL).

Usage

```
set_data(.mp, data)
```

Arguments

.mp	a TSMP object.
data	a matrix (for one series) or a list of matrices (for two series).

Value

Returns silently the original TSMP object with changed data.

Examples

```
mp <- tsmp(mp_toy_data$data[1:200, 1], window_size = 30, verbose = 0)
mp <- set_data(mp, NULL)
```

simple_fast	<i>Compute the join similarity for Sound data</i>
-------------	---

Description

Compute the join similarity for Sound data

Usage

```
simple_fast(
  ...,
  window_size,
  exclusion_zone = getOption("tsmp.exclusion_zone", 1/2),
  verbose = getOption("tsmp.verbose", 2)
)
```

Arguments

- ... a matrix of numeric, where each column is a time series. Accepts list and data.frame too. If a second time series is supplied it will be a join matrix profile.
- window_size an int with the size of the sliding window.
- exclusion_zone a numeric. Size of the exclusion zone, based on window size (default is 1/2).
- verbose an int. See details. (Default is 2).

Details

verbose changes how much information is printed by this function; 0 means nothing, 1 means text, 2 adds the progress bar, 3 adds the finish sound.

Value

Returns a SimpleMatrixProfile object, a list with the matrix profile mp, profile index pi, number of dimensions n_dim, window size w and exclusion zone ez.

References

- Silva D, Yeh C, Batista G, Keogh E. Simple: Assessing Music Similarity Using Subsequences Joins. Proc 17th ISMIR Conf. 2016;23-30.
- Silva DF, Yeh C-CM, Zhu Y, Batista G, Keogh E. Fast Similarity Matrix Profile for Music Analysis and Exploration. IEEE Trans Multimed. 2018;14(8):1-1.

Website: <https://sites.google.com/view/simple-fast>

Website: <https://sites.google.com/site/ismir2016simple/home>

Examples

```
w <- 30
data <- mp_toy_data$data # 3 dimensions matrix
result <- simple_fast(data, window_size = w, verbose = 0)
```

stamp_par

Anytime univariate STAMP algorithm Parallel version

Description

Computes the best so far Matrix Profile and Profile Index for Univariate Time Series.

Usage

```

stamp_par(
  ...,
  window_size,
  exclusion_zone = getOption("tsmp.exclusion_zone", 1/2),
  verbose = getOption("tsmp.verbose", 2),
  s_size = Inf,
  n_workers = 2,
  weight = NULL
)

stamp(
  ...,
  window_size,
  exclusion_zone = getOption("tsmp.exclusion_zone", 1/2),
  verbose = getOption("tsmp.verbose", 2),
  s_size = Inf,
  weight = NULL
)

```

Arguments

...	a matrix or a vector. If a second time series is supplied it will be a join matrix profile.
window_size	an int. Size of the sliding window.
exclusion_zone	a numeric. Size of the exclusion zone, based on window size (default is 1/2). See details.
verbose	an int. See details. (Default is 2).
s_size	a numeric. for anytime algorithm, represents the size (in observations) the random calculation will occur (default is Inf).
n_workers	an int. Number of workers for parallel. (Default is 2).
weight	a vector of numeric or NULL with the same length of the window_size. This is a MASS extension to weight the query.

Details

The Matrix Profile, has the potential to revolutionize time series data mining because of its generality, versatility, simplicity and scalability. In particular it has implications for time series motif discovery, time series joins, shapelet discovery (classification), density estimation, semantic segmentation, visualization, rule discovery, clustering etc. The anytime STAMP computes the Matrix Profile and Profile Index in such manner that it can be stopped before its complete calculation and return the best so far results allowing ultra-fast approximate solutions. `verbose` changes how much information is printed by this function; 0 means nothing, 1 means text, 2 adds the progress bar, 3 adds the finish sound. `exclusion_zone` is used to avoid trivial matches; if a query data is provided (join similarity), this parameter is ignored.

Value

Returns a MatrixProfile object, a list with the matrix profile mp, profile index pi left and right matrix profile lmp, rmp and profile index lpi, rpi, window size w and exclusion zone ez.

Functions

- stamp_par(): Parallel version.
- stamp(): Single thread version.

References

- Yeh CCM, Zhu Y, Ulanova L, Begum N, Ding Y, Dau HA, et al. Matrix profile I: All pairs similarity joins for time series: A unifying view that includes motifs, discords and shapelets. Proc - IEEE Int Conf Data Mining, ICDM. 2017;1317-22.
- Zhu Y, Imamura M, Nikovski D, Keogh E. Matrix Profile VII: Time Series Chains: A New Primitive for Time Series Data Mining. Knowl Inf Syst. 2018 Jun 2;1-27.

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

See Also

Other matrix profile computations: [mstomp_par\(\)](#), [scrimp\(\)](#), [stomp_par\(\)](#), [tsmp\(\)](#), [valmod\(\)](#)

Examples

```
mp <- stamp(mp_toy_data$data[1:200, 1], window_size = 30, verbose = 0)

#' # using threads
mp <- stamp_par(mp_toy_data$data[1:200, 1], window_size = 30, verbose = 0)

ref_data <- mp_toy_data$data[, 1]
query_data <- mp_toy_data$data[, 2]
# self similarity
mp <- stamp(ref_data, window_size = 30, s_size = round(nrow(ref_data) * 0.1))
# join similarity
mp <- stamp(ref_data, query_data, window_size = 30, s_size = round(nrow(query_data) * 0.1))
```

stompi_update

Real-time STOMP algorithm

Description

Real-time STOMP algorithm

Usage

```
stompi_update(.mp, new_data, history_size = FALSE)
```

Arguments

`.mp` a TSMP object of class `MatrixProfile`.

`new_data` new data to append to original data.

`history_size` an int or `FALSE`. (Default is `FALSE`). Keep only this amount of data in the object. The value is for the data, not the matrix profile. Notice that the `lmpand lpi` will be inconsistent when repeatedly updating limiting the history size and thus will affect the `mp` and `pi`.

Value

Returns the input `.mp` updated with the new information.

Examples

```
mp <- tsmc(mp_toy_data$data[1:200], 1, window_size = 30, verbose = 0)
mpi <- stompi_update(mp, mp_toy_data$data[201:300], 1)
mp <- tsmc(mp_toy_data$data[1:300], 1, window_size = 30, verbose = 0)
all.equal(mp, mpi, check.attributes = FALSE)
```

stomp_par

Univariate STOMP algorithm

Description

Computes the Matrix Profile and Profile Index for Univariate Time Series.

Usage

```
stomp_par(
  ...,
  window_size,
  exclusion_zone = getOption("tsmp.exclusion_zone", 1/2),
  verbose = getOption("tsmp.verbose", 2),
  n_workers = 2
)

stomp(
  ...,
  window_size,
  exclusion_zone = getOption("tsmp.exclusion_zone", 1/2),
  verbose = getOption("tsmp.verbose", 2)
)
```


Arguments

...	a matrix or a vector. If a second time series is supplied it will be a join matrix profile.
window_size	an int. Size of the sliding window.
exclusion_zone	a numeric. Size of the exclusion zone, based on window size (default is 1/2). See details.
verbose	an int. See details. (Default is 2).
n_workers	an int. Number of workers for parallel. (Default is 2).

Details

The Matrix Profile, has the potential to revolutionize time series data mining because of its generality, versatility, simplicity and scalability. In particular it has implications for time series motif discovery, time series joins, shapelet discovery (classification), density estimation, semantic segmentation, visualization, rule discovery, clustering etc. `verbose` changes how much information is printed by this function; 0 means nothing, 1 means text, 2 adds the progress bar, 3 adds the finish sound. `exclusion_zone` is used to avoid trivial matches; if a query data is provided (join similarity), this parameter is ignored.

Value

Returns a `MatrixProfile` object, a list with the matrix profile `mp`, profile index `pi` left and right matrix profile `lmp`, `rmp` and profile index `lpi`, `rpi`, window size `w` and exclusion zone `ez`.

Functions

- `stomp_par()`: Parallel version.
- `stomp()`: Single thread version.

References

- Zhu Y, Zimmerman Z, Senobari NS, Yeh CM, Funning G. Matrix Profile II : Exploiting a Novel Algorithm and GPUs to Break the One Hundred Million Barrier for Time Series Motifs and Joins. *Icdm*. 2016 Jan 22;54(1):739-48.

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

See Also

Other matrix profile computations: `mstomp_par()`, `scrimp()`, `stamp_par()`, `tsmp()`, `valmod()`

Examples

```
mp <- stomp(mp_toy_data$data[1:200, 1], window_size = 30, verbose = 0)

#' # using threads
mp <- stomp_par(mp_toy_data$data[1:400, 1], window_size = 30, verbose = 0)

ref_data <- mp_toy_data$data[, 1]
```

```

query_data <- mp_toy_data$data[, 2]
# self similarity
mp <- stomp(ref_data, window_size = 30)
# join similarity
mp2 <- stomp(ref_data, query_data, window_size = 30)

```

tsmp

Computes the Matrix Profile and Profile Index

Description

This is a wrap function that makes easy to use all available algorithms to compute the Matrix Profile and Profile Index for multiple purposes.

Usage

```

tsmp(
  ...,
  window_size,
  exclusion_zone = getOption("tsmp.exclusion_zone", 1/2),
  mode = c("stomp", "stamp", "simple", "mstomp", "scrimp", "valmod", "pmp"),
  verbose = getOption("tsmp.verbose", 2),
  n_workers = 1,
  s_size = Inf,
  must_dim = NULL,
  exc_dim = NULL,
  heap_size = 50,
  paa = 1,
  .keep_data = TRUE
)

```

Arguments

...	a matrix or a vector. If a second time series is supplied it will be a join matrix profile (except for <code>mstomp()</code>).
window_size	an int with the size of the sliding window. Use a vector for Valmod.
exclusion_zone	a numeric. Size of the exclusion zone, based on window size (default is 1/2). See details.
mode	the algorithm that will be used to compute the matrix profile. (Default is stomp). See details.
verbose	an int. (Default is 2). See details.
n_workers	an int. Number of workers for parallel. (Default is 1).
s_size	a numeric. for anytime algorithm, represents the size (in observations) the random calculation will occur (default is Inf). See details.

<code>must_dim</code>	an int or vector of which dimensions to forcibly include (default is NULL). See details.
<code>exc_dim</code>	an int or vector of which dimensions to exclude (default is NULL). See details.
<code>heap_size</code>	an int. (Default is 50). Size of the distance profile heap buffer.
<code>paa</code>	an int. (Default is 1). Factor of PAA reduction (2 == half of size)
<code>.keep_data</code>	a logical. (Default is TRUE). Keeps the data embedded to resultant object.

Details

The Matrix Profile, has the potential to revolutionize time series data mining because of its generality, versatility, simplicity and scalability. In particular it has implications for time series motif discovery, time series joins, shapelet discovery (classification), density estimation, semantic segmentation, visualization, rule discovery, clustering etc.

The first algorithm invented was the `stamp()` that using `mass()` as an ultra-fast Algorithm for Similarity Search allowed to compute the Matrix Profile in reasonable time. One of its main feature was its Anytime property which using a randomized approach could return a "best-so-far" matrix that could give us the correct answer (using for example 1/10 of all iterations) almost every time.

The next algorithm was `stomp()` that currently is the most used. Researchers noticed that the dot products do not need to be recalculated from scratch for each subsequence. Instead, we can reuse the values calculated for the first subsequence to make a faster calculation in the next iterations. The idea is to make use of the intersections between the required products in consecutive iterations. This approach reduced the time to compute the Matrix Profile to about 3% compared to `stamp()`, but on the other hand, we lost the Anytime property.

Currently there is a new algorithm that I'll not explain further here. It is called `scrimp()`, and is as fast as `stomp()`, and have the Anytime property. This algorithm is implemented in this package, but still waiting for an article publication.

Further, there is the `mstomp()` that computes a multidimensional Matrix Profile that allows to meaningful MOTIF discovery in Multivariate Time Series. And `simple_fast()` that also handles Multivariate Time Series, but focused in Music Analysis and Exploration.

The `valmod()` uses a new pruning algorithm allowing a similarity search with a range of sliding window sizes.

The `pmp()` is a new concept that creates several profiles from a range of windows.

Some parameters are global across the algorithms:

... One or two time series (except for `mstomp()`). The second time series can be smaller than the first.

window_size The sliding window.

exclusion_zone Is used to avoid trivial matches; if a query data is provided (join similarity), this parameter is ignored.

verbose Changes how much information is printed by this function; 0 means nothing, 1 means text, 2 adds the progress bar, 3 adds the finish sound.

n_workers number of threads for parallel computing (except `simple_fast`, `scrimp` and `valmod`). If the value is 2 or more, the `'_par'` version of the algorithm will be used.

`s_size` is used only in Anytime algorithms: `stamp()` and `scrimp()`. `must_dim` and `exc_dim` are used only in `mstomp()`. `heap_size` is used only for `valmod()` mode can be any of the following: `stomp`, `stamp`, `simple`, `mstomp`, `scrimp`, `valmod`, `pmp`.

Value

Returns the matrix profile `mp` and profile index `pi`. It also returns the left and right matrix profile `lmp`, `rmp` and profile index `lpi`, `rpi` that may be used to detect Time Series Chains. `mstomp()` returns a multidimensional Matrix Profile.

References

- Silva D, Yeh C, Batista G, Keogh E. Simple: Assessing Music Similarity Using Subsequences Joins. Proc 17th ISMIR Conf. 2016;23-30.
- Silva DF, Yeh C-CM, Zhu Y, Batista G, Keogh E. Fast Similarity Matrix Profile for Music Analysis and Exploration. IEEE Trans Multimed. 2018;14(8):1-1.
- Yeh CM, Kavantzias N, Keogh E. Matrix Profile VI : Meaningful Multidimensional Motif Discovery.
- Yeh CCM, Zhu Y, Ulanova L, Begum N, Ding Y, Dau HA, et al. Matrix profile I: All pairs similarity joins for time series: A unifying view that includes motifs, discords and shapelets. Proc - IEEE Int Conf Data Mining, ICDM. 2017;1317-22.
- Zhu Y, Imamura M, Nikovski D, Keogh E. Matrix Profile VII: Time Series Chains: A New Primitive for Time Series Data Mining. Knowl Inf Syst. 2018 Jun 2;1-27.
- Zhu Y, Zimmerman Z, Senobari NS, Yeh CM, Funning G. Matrix Profile II : Exploiting a Novel Algorithm and GPUs to Break the One Hundred Million Barrier for Time Series Motifs and Joins. Icdm. 2016 Jan 22;54(1):739-48.

Website: <https://sites.google.com/view/simple-fast>

Website: <https://sites.google.com/site/ismir2016simple/home>

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

See Also

Other matrix profile computations: `mstomp_par()`, `scrimp()`, `stamp_par()`, `stomp_par()`, `valmod()`

Examples

```
# default with [stomp()]
mp <- tsmp(mp_toy_data$data[1:200, 1], window_size = 30, verbose = 0)

# Anytime STAMP
mp <- tsmp(mp_toy_data$data[1:200, 1], window_size = 30, mode = "stamp", s_size = 50, verbose = 0)

# [mstomp()]
mp <- tsmp(mp_toy_data$data[1:200, ], window_size = 30, mode = "mstomp", verbose = 0)

# [simple_fast()]
mp <- tsmp(mp_toy_data$data[1:200, ], window_size = 30, mode = "simple", verbose = 0)

# parallel with [stomp_par()]
mp <- tsmp(mp_test_data$train$data[1:1000, 1], window_size = 30, n_workers = 2, verbose = 0)
```

 valmod *Variable Length Motif Discovery*

Description

Computes the Matrix Profile and Profile Index for a range of query window sizes

Usage

```
valmod(
  ...,
  window_min,
  window_max,
  heap_size = 50,
  exclusion_zone = getOption("tsmp.exclusion_zone", 1/2),
  lb = TRUE,
  verbose = getOption("tsmp.verbose", 2)
)
```

Arguments

...	a matrix or a vector. If a second time series is supplied it will be a join matrix profile.
window_min	an int. Minimum size of the sliding window.
window_max	an int. Maximum size of the sliding window.
heap_size	an int. (Default is 50). Size of the distance profile heap buffer
exclusion_zone	a numeric. Size of the exclusion zone, based on window size (default is 1/2). See details.
lb	a logical. (Default is TRUE). If FALSE all window sizes will be calculated using STOMP instead of pruning. This is just for academic purposes.
verbose	an int. See details. (Default is 2).

Details

This algorithm uses an exact algorithm based on a novel lower bounding technique, which is specifically designed for the motif discovery problem. `verbose` changes how much information is printed by this function; 0 means nothing, 1 means text, 2 adds the progress bar, 3 adds the finish sound. `exclusion_zone` is used to avoid trivial matches; if a query data is provided (join similarity), this parameter is ignored.

Paper that implements `skimp()` suggests that `window_max / window_min > than 1.24` begins to weakening pruning in `valmod()`.

Value

Returns a Valmod object, a list with the matrix profile mp, profile index pi left and right matrix profile lmp, rmp and profile index lpi, rpi, best window size w for each index and exclusion zone ez. Additionally: evolution_motif the best motif distance per window size, and non-length normalized versions of mp, pi and w: mpnn, pinn and wnn.

References

- Linardi M, Zhu Y, Palpanas T, Keogh E. VALMOD: A Suite for Easy and Exact Detection of Variable Length Motifs in Data Series. In: Proceedings of the 2018 International Conference on Management of Data - SIGMOD '18. New York, New York, USA: ACM Press; 2018. p. 1757-60.

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

See Also

Other matrix profile computations: [mstomp_par\(\)](#), [scrimp\(\)](#), [stamp_par\(\)](#), [stomp_par\(\)](#), [tsmp\(\)](#)

Examples

```
mp <- valmod(mp_toy_data$data[1:200, 1], window_min = 30, window_max = 40, verbose = 0)

ref_data <- mp_toy_data$data[, 1]
query_data <- mp_toy_data$data[, 2]
# self similarity
mp <- valmod(ref_data, window_min = 30, window_max = 40)
# join similarity
mp <- valmod(ref_data, query_data, window_min = 30, window_max = 40)
```

visualize

Plots an object generated from one of the algorithms. In some cases multiple plots will be generated

Description

Plots an object generated from one of the algorithms. In some cases multiple plots will be generated

Usage

```
visualize(profile)
```

Arguments

profile a MatrixProfile or PMP object.

References

Website: <http://www.cs.ucr.edu/~eamonn/MatrixProfile.html>

See Also

Other Main API: [analyze\(\)](#), [compute\(\)](#), [discords\(\)](#), [motifs\(\)](#)

Examples

```
result <- compute(mp_toy_data$data[, 1], 80)
visualize(result)
```

write	<i>Write a TSMP object to JSON file.</i>
-------	--

Description

Write a TSMP object to JSON file.

Usage

```
write(x, ...)

## S3 method for class 'MatrixProfile'
write(x, file, ...)

## S3 method for class 'PMP'
write(x, file, ...)
```

Arguments

x	a MatrixProfile or PMP object. If not, the <code>base::write()</code> function will be called.
...	other arguments to be passed forward.
file	a character string with the output filename.

Examples

```
result <- compute(mp_toy_data$data[, 1], 80)

write(result, file = file.path(tempdir(), "output.json"))
```

Index

- * **Annotation vectors**
 - av_apply, 6
 - av_complexity, 7
 - av_hardlimit_artifact, 8
 - av_motion_artifact, 9
 - av_stop_word, 10
 - av_zerocrossing, 11
- * **Main API**
 - analyze, 3
 - compute, 12
 - discords, 13
 - motifs, 32
 - visualize, 70
- * **Scalable Dictionaries**
 - sdtc_predict, 56
 - sdtc_score, 57
 - sdtc_train, 58
- * **Semantic Segmentations**
 - floss, 22
 - floss_cac, 24
 - floss_extract, 25
 - fluss, 26
 - fluss_cac, 27
 - fluss_extract, 28
 - fluss_score, 29
- * **datasets**
 - motifs_discords_small, 33
 - mp_fluss_data, 36
 - mp_gait_data, 37
 - mp_meat_data, 37
 - mp_test_data, 38
 - mp_toy_data, 39
- * **distance measure**
 - mpdist, 34
- * **hplot**
 - plot, 42
 - plot_arcs, 45
- * **matrix profile computations**
 - mstamp_par, 40
 - scrimp, 54
 - stamp_par, 61
 - stomp_par, 64
 - tsmp, 66
 - valmod, 69
- analyze, 3, 13, 14, 33, 71
- as.arccount (as.matrixprofile), 5
- as.chain (as.matrixprofile), 5
- as.discord (as.matrixprofile), 5
- as.fluss (as.matrixprofile), 5
- as.matrixprofile, 5
- as.motif (as.matrixprofile), 5
- as.multimatrixprofile
 - (as.matrixprofile), 5
- as.multimotif (as.matrixprofile), 5
- as.pmp (as.matrixprofile), 5
- as.salient (as.matrixprofile), 5
- as.valmod (as.matrixprofile), 5
- av_apply, 6, 7–9, 11
- av_complexity, 7, 7, 8, 9, 11
- av_hardlimit_artifact, 7, 8, 9, 11
- av_motion_artifact, 7, 8, 9, 11
- av_stop_word, 7–9, 10, 11
- av_zerocrossing, 7–9, 11, 11
- compute, 4, 12, 14, 33, 71
- discords, 4, 13, 13, 33, 71
- dist_profile, 14
- fast_avg_sd, 16
- fast_movavg, 16
- fast_movsd, 17
- find_chains, 17
- find_discord, 18
- find_motif, 19
- find_snippet, 21
- floss, 22, 25–29
- floss_cac, 23, 24, 26–29

floss_extract, 23, 25, 25, 26–29
fluss, 23, 25, 26, 26, 27–29
fluss(), 56, 58
fluss_cac, 23, 25, 26, 27, 28, 29
fluss_cac(), 26
fluss_extract, 23, 25–27, 28, 29
fluss_extract(), 26, 29
fluss_score, 23, 25–28, 29

get_data, 30

mass(), 67
mass_pre(), 31
mass_v3, 30
min_mp_idx, 32
motifs, 4, 13, 14, 32, 71
motifs_discords_small, 33
mp_fluss_data, 36
mp_gait_data, 37
mp_meat_data, 37
mp_test_data, 38
mp_toy_data, 39
mpdist, 34
mpx, 35
mstomp (mstomp_par), 40
mstomp(), 66–68
mstomp_par, 40, 55, 63, 65, 68, 70

par(), 45, 46
plot, 42
plot(), 45, 46
plot_arcs, 45
pmp, 47
pmp(), 67
pmp_upper_bound, 48

read, 50
remove_class, 50

salient_mds, 51
salient_score, 52
salient_subsequences, 53
scrimp, 41, 54, 63, 65, 68, 70
scrimp(), 67
sdts_predict, 56, 57, 59
sdts_predict(), 57
sdts_score, 56, 57, 59
sdts_train, 56, 57, 58
sdts_train(), 56

set_data, 60
simple_fast, 60
simple_fast(), 67
stamp (stamp_par), 61
stamp(), 67
stamp_par, 41, 55, 61, 65, 68, 70
stomp (stomp_par), 64
stomp(), 41, 67
stomp_par, 41, 55, 63, 64, 68, 70
stompi_update, 63

tsmp, 41, 55, 63, 65, 66, 70

valmod, 41, 55, 63, 65, 68, 69
valmod(), 67
visualize, 4, 13, 14, 33, 70

write, 71