

Package: TML (via r-universe)

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Type Package

Title Tropical Geometry Tools for Machine Learning

Version 2.3.0

Description Suite of tropical geometric tools for use in machine learning applications. These methods may be summarized in the following references: Yoshida, et al. (2022) <[arxiv:2209.15045](https://arxiv.org/abs/2209.15045)>, Barnhill et al. (2023) <[arxiv:2303.02539](https://arxiv.org/abs/2303.02539)>, Barnhill and Yoshida (2023) <[doi:10.3390/math11153433](https://doi.org/10.3390/math11153433)>, Aliatimis et al. (2023) <[arXiv:2306.08796](https://arxiv.org/abs/2306.08796)>, Yoshida et al. (2022) <[arXiv:2206.04206](https://arxiv.org/abs/2206.04206)>, and Yoshida et al. (2019) <[doi:10.1007/s11538-018-0493-4](https://doi.org/10.1007/s11538-018-0493-4)>.

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URL <https://github.com/barnhilldave/TML>

BugReports <https://github.com/barnhilldave/TML/issues>

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<i>bw.nn</i>	<i>Nearest neighbor bandwidth calculation</i>
--------------	---

Description

This function finds the bandwidth for an ultrametric based on the tropical distance of the nearest point. The function provides the bandwidth input to `trop.KDE` and was originally used in the `KDETrees` package.

Usage

```
bw.nn(x, prop = 0.2, tol = 1e-06)
```

Arguments

<code>x</code>	matrix; dissimilarity matrix between points in a data set
<code>prop</code>	proportion of observations that defines neighborhood of a point
<code>tol</code>	tolerance for zero bandwidth check

Value

a vector of bandwidths for each tree (row) in `x`

Author(s)

Ruriko Yoshida <ryoshida@nps.edu>

References

Weyenberg, G., Huggins, P., Schardl, C., Howe, D. K., & Yoshida, R. (2014). `kdetrees`: Nonparametric Estimation of Phylogenetic Tree Distributions. In *Bioinformatics*.

<https://github.com/grady/kdetrees/blob/master/R/bw.R>

Examples

```
T1<-Sim_Trees15
T2<-Sim_Trees25
D <- rbind(T1, T2[,])
M <- pw.trop.dist(D, D)
bw.nn(M)
```

cluster.ratio_HC	<i>Ratio of within and between tropical measures for tropical hierarchical clusters</i>
------------------	---

Description

Ratio of within and between cluster tropical measures for a set hierarchical clusters

Usage

```
cluster.ratio_HC(A, V, method = mean)
```

Arguments

A	matrix of tropical points; rows are points
V	list of clusters where each cluster is defined as a matrix
method	method to use for within cluster measure; mean or max

Value

vector of ratios for each cluster

Author(s)

David Barnhill <david.barnhill@nps.edu>

References

David Barnhill, Ruriko Yoshida (2023). Clustering Methods Over the Tropically Convex Sets.

Examples

```
har<-rbind(Sim_points[1:20,],Sim_points[51:70,])  
  
V<-Tropical.HC.AGNES(har, method=mean)  
inds<-V[[2]][[38]]  
cluster.ratio_HC(har, inds, method=mean)
```

cluster.ratio_KM	<i>Ratio of within and between tropical measures for k-means clusters</i>
------------------	---

Description

Ratio of within and between cluster tropical measures for k-means derived clusters

Usage

```
cluster.ratio_KM(A, C, method = mean)
```

Arguments

A	matrix of tropical points; rows are points
C	number of clusters
method	method to use for within cluster measure; mean or max

Value

vector of ratios for each cluster

Author(s)

David Barnhill <david.barnhill@nps.edu>

References

David Barnhill, Ruriko Yoshida (2023). Clustering Methods Over the Tropically Convex Sets.

Examples

```
hars<-Sim_points
cls<-c(rep(1,50),rep(2,50),rep(3,50))
cl_pt<-cbind(hars,cls)

C<-3
cluster.ratio_KM(cl_pt,C,method=mean)
```

convert.to.tree *Create a phylogenetic tree from an ultrametric*

Description

This function constructs a phylogenetic tree from an ultrametric.

Usage

```
convert.to.tree(n, L, u)
```

Arguments

n is the number of leaves
L is a vector of labels (strings) of leaves
u is an ultrametric

Value

A phylogenetic tree of class phylo

Author(s)

Ruriko Yoshida <ryoshida@nps.edu>

Examples

```
um<-Sim_Trees21[1,]  
ll <- 10  
L <- LETTERS[1:10]  
tr<-convert.to.tree(ll, L, um)
```

draw.tpolytope *Draw a 2-D or 3-D tropical polytope*

Description

This command draws a three dimensional tropical polytope

Usage

```
draw.tpolytope.3d(D, col_lines, col_verts, plot = TRUE, tadd = max)
```

```
draw.tpolytope.2d(D, col_lines, col_verts, plot = TRUE, tadd = max)
```

Arguments

D	matrix of vertices of a tropical polytope; rows are the vertices
col_lines	string; color to render the polytope.
col_verts	string; color to render the vertices.
plot	logical; initiate new plot visualization or not.
tadd	function; max indicates max-plus addition, min indicates min-plus addition. Defaults to max

Value

2-D or 3-D rendering of a tropical polytope.

Author(s)

Ruriko Yoshida <ryoshida@nps.edu>

Examples

```
D <-matrix(c(0,0,0,0,0,1,0,0,0,0,1,0,0,0,0,1),4,4,TRUE)
col_lines<-'blue'
col_verts<-'red'
draw.tpolytope.3d(D,col_lines,col_verts,plot=TRUE)
draw.tpolytope.3d(D,col_lines,col_verts,plot=TRUE,tadd=min)
```

```
D <- matrix(c(0,-2,2,0,-2,5,0,2,1,0,1,-1),4,3,TRUE)
col_lines <- 'blue'
col_verts <- 'red'
draw.tpolytope.2d(D,col_lines,col_verts,plot=TRUE)
draw.tpolytope.2d(D,col_lines,col_verts,plot=TRUE,tadd=min)
```

FWpoint.num.w.reg

Modified Fermat-Weber point numerical solver for ultrametrics

Description

Returns a modified Fermat-Weber point of N points using a gradient based numerical method. This method is appropriate for points coming from ultrametrics. The algorithm tries to find a point that minimizes the sum of tropical distances from the samples, but also also tries to find a point that is as close as possible to the space of ultrametrics. The tradeoff between these two objectives is controlled by the penalty parameter. If penalty=0, the method is identical to FWpoint_numerical; it finds the Fermat-Weber point, which may not be an ultrametric. If penalty is very large, the algorithm is trying to find the Fermat-Weber point in the space of ultrametrics.

Usage

```
FWpoint.num.w.reg(datamatrix, penalty = 0)
```

Arguments

`datamatrix` matrix of dimension $N \times e$, where N is the number of observations which lie in \mathbb{R}^e

`penalty` positive real number; the regularization rate

Value

vector; Fermat-Weber point approximation

Author(s)

Georgios Aliatimis <g.aliatimis@lancaster.ac.uk>

References

Aliatimis, Georgios, Ruriko Yoshida, Burak Boyaci and James A. Grant (2023). Tropical Logistic Regression on Space of Phylogenetic Trees

Examples

```
D = matrix(c(0,0,0,0,2,5,0,3,1),3,3,TRUE)
FWpoint.num.w.reg(D,1e4) # (0,2,5/3) not ultrametric
FWpoint.num.w.reg(D,1e4) # (0,5/3,5/3) ultrametric
```

`FWpoint.numerical` *Fermat-Weber point numerical solver*

Description

Returns the Fermat-Weber point of N points using a gradient based numerical method

Usage

```
FWpoint.numerical(datamatrix)
```

Arguments

`datamatrix` matrix of dimension $N \times e$, where N is the number of observations which lie in \mathbb{R}^e .

Value

Fermat-Weber point approximation (vector in \mathbb{R}^e)

Author(s)

Georgios Aliatimis <g.aliatimis@lancaster.ac.uk>

References

Aliatimis, Georgios, Ruriko Yoshida, Burak Boyaci and James A. Grant (2023). Tropical Logistic Regression on Space of Phylogenetic Trees

Examples

```
D = matrix(c(0,0,0,0,2,5,0,3,1),3,3,TRUE)
FWpoint.numerical(D)
```

HAR.TLineSeg	<i>Uniformly sample from a max-plus tropical line segment</i>
--------------	---

Description

This function uses a hit-and-run sampler to uniformly sample from a max-plus tropical line segment

Usage

```
HAR.TLineSeg(D1, D2, tadd = max)
```

Arguments

D1	point in the tropical projective torus
D2	point in the tropical projective torus
tadd	string; max indicates max-plus addition, min indicates min-plus addition. Defaults to max

Value

point on the line segment defined by D1 and D2

Author(s)

Ruriko Yoshida <ryoshida@nps.edu>

References

Yoshida, Ruriko, Keiji Miura and David Barnhill (2022). Hit and Run Sampling from Tropically Convex Sets.

Examples

```
D1 <-c(0,4,2)
D2 <- c(0,7,-1)
HAR.TLineSeg(D1, D2,tadd=max)
HAR.TLineSeg(D1, D2,tadd=min)
```

HAR.TLineSeg.centroid *Gaussian-like Sampling on a max- or min-plus tropical line segment*

Description

This function samples points on a tropical line segment about a location parameter for a given scale parameter defined in terms of tropical distance

Usage

```
HAR.TLineSeg.centroid(D1, D2, m, s, tadd = max)
```

Arguments

D1	point in the tropical projective torus
D2	point in the tropical projective torus
m	location parameter
s	scale parameter
tadd	function; max indicates max-plus addition, min indicates min-plus addition. Defaults to max

Value

point on the line segment defined by D1 and D2 sampled about mu

Author(s)

David Barnhill <david.barnhill@nps.edu>

Examples

```
D1 <-c(0,4,2)
D2 <- c(0,7,-1)
m<-c(0,7,2)
s<-1
HAR.TLineSeg.centroid(D1, D2,m,s)
HAR.TLineSeg.centroid(D1, D2,m,s,tadd=min)
```

`hyper_3D`*2D or 3D rendering of max-plus or min-plus tropical hyperplane*

Description

This function renders a 2D or 3D max-plus or min-plus tropical hyperplane

Usage

```
draw.thyper(D, ext, min.ax, max.ax, plot = FALSE, tadd = max)
```

Arguments

<code>D</code>	point in the tropical projective torus representing the apex of the hyperplane
<code>ext</code>	scalar; indicates how far the hyperplane should extend
<code>min.ax</code>	scalar; value applied to define the minimum limits of the axes of the plot
<code>max.ax</code>	scalar; value applied to define the maximum limits of the axes of the plot
<code>plot</code>	logical; if true produces a new plot otherwise overlays tropical hyperplane on existing plot
<code>tadd</code>	function; max indicates max-plus addition, min indicates min-plus addition. Defaults to max

Value

2D or 3D rendering of max-plus or min-plus tropical hyperplane

Author(s)

David Barnhill <david.barnhill@nps.edu>

Examples

```
# 2D Example
D <-c(0,0,0)
ext<-4
min.ax<- 5
max.ax<- 5
draw.thyper(D,ext,min.ax,max.ax,plot=TRUE)

# 3D Example
D <-c(0,0,0,0)
ext<-4
min.ax<- 5
max.ax<- 5
draw.thyper(D,ext,min.ax,max.ax,plot=TRUE)
draw.thyper(D,ext,min.ax,max.ax,plot=TRUE,tadd=min)
```

lung_fish

Phylogenetic trees based on lung fish data

Description

1290 (non-equidistant) gene trees with 45 leaves originating from lung fish data in matrix form. Also we provide a vector of strings consisting of leaf labels for each species associated with the data set.

Usage

```
lung_fish
```

Format

An object of class `matrix` (inherits from `array`) with 1290 rows and 45 columns.

Source

Liang D, Shen XX, Zhang P. One thousand two hundred ninety nuclear genes from a genome-wide survey support lungfishes as the sister group of tetrapods. *Mol Biol Evol.* 2013 Aug;30(8):1803-7. doi: 10.1093/molbev/mst072. Epub 2013 Apr 14. PMID: 23589454.

max_ins.ball

Calculate the center point and radius of the maximum inscribed ball for a tropical simplex

Description

This function calculates the center point and radius of the maximum inscribed ball for a max- or min-plus tropical simplex

Usage

```
max_ins.ball(A, tadd = max)
```

Arguments

A	matrix of points defining a tropical polytope; rows are the points
tadd	function; max indicates max-plus addition, min indicates min-plus addition. Defaults to max

Value

list containing the radius and center point of a maximum inscribed ball

Author(s)

David Barnhill <david.barnhill@nps.edu>

References

Barnhill, David, Ruriko Yoshida and Keiji Miura (2023). Maximum Inscribed and Minimum Enclosing Tropical Balls of Tropical Polytopes and Applications to Volume Estimation and Uniform Sampling.

Examples

```
P<-matrix(c(0,0,0,0,2,5,0,3,1),3,3,TRUE)
max_ins.ball(P)
max_ins.ball(P,tadd=min)
```

min_enc.ball

Calculate a minimum enclosing ball for a tropical polytope

Description

This function constructs a minimum enclosing ball for a set of points defining a tropical polytope.

Usage

```
min_enc.ball(A)
```

Arguments

A matrix of points defining a tropical polytope. Rows are the points.

Value

list containing center point and radius of minimum enclosing ball of P

Author(s)

David Barnhill <david.barnhill@nps.edu>

References

Barnhill, David, Ruriko Yoshida and Keiji Miura (2023). Maximum Inscribed and Minimum Enclosing Tropical Balls of Tropical Polytopes and Applications to Volume Estimation and Uniform Sampling.

Examples

```
P <-matrix(c(0,0,0,0,3,1,0,2,5),3,3,TRUE)
min_enc.ball(P)
```

normaliz.tree	<i>Normalize a phylogenetic tree</i>
---------------	--------------------------------------

Description

This function normalizes the height of a phylogenetic tree

Usage

```
normaliz.tree(D, h = 1)
```

Arguments

D	numeric vector; ultrametric equidistant tree
h	desired height; defaults to 1

Value

normalized equidistant tree

Author(s)

Ruriko Yoshida <ryoshida@nps.edu>

Examples

```
D <-c(4,4,2)
normaliz.tree(D, h=1)
```

normalize	<i>Normalize a point or set of points in the tropical projective torus</i>
-----------	--

Description

This function normalizes a point or set of points in the tropical projective torus by making the first coordinate zero

Usage

```
normaliz.vector(D)
```

```
normaliz.vectors(D)
```

```
normaliz.polytope(D)
```

```
normaliz.ultrametrics(D)
```

Arguments

D numeric vector in the tropical projective torus or a matrix of points in the tropical projective torus; for matrices, rows are the points

Value

a single or set of normalized points with the first coordinate zero

Author(s)

Ruriko Yoshida <ryoshida@nps.edu>

Examples

```
D <-c(8,4,2)
normaliz.vector(D)

P <-matrix(c(8,4,2,10,1,3,7,2,1),3,3,TRUE)
normaliz.vectors(P)

M<-matrix(c(2,2,2,3,6,4,2,4,7),3,3,TRUE)
normaliz.polytope(M)

M <- Sim_Trees15[1:3,]
normaliz.ultrametrics(M)
```

over_bet_HC	<i>Tropical cluster betweenness measure for each cluster in a set of hierarchical clusters</i>
-------------	--

Description

This function calculates an overall betweenness measure based on tropical distance between a set of clusters derived from tropical hierarchical clustering

Usage

```
over_bet_HC(A, V)
```

Arguments

A matrix of tropical points; rows are points with the last column representing a numbered cluster assignment

V list of clusters defined as matrices derived from agglomerative or divisive hierarchical clustering

Value

vector of betweenness cluster measures

Author(s)

David Barnhill <david.barnhill@nps.edu>

References

David Barnhill, Ruriko Yoshida (2023). Clustering Methods Over the Tropically Convex Sets.

Examples

```
har<-rbind(Sim_points[1:20,],Sim_points[51:70,])

V<-Tropical.HC.AGNES(har, method=mean)
inds<-V[[2]][[38]]
over_bet_HC(har, inds)
```

over_bet_KM	<i>Tropical cluster betweenness measure for a each of a set of k-means derived set of clusters</i>
-------------	--

Description

This function calculates an overall betweenness measure between a set of clusters derived from tropical k-means clustering

Usage

```
over_bet_KM(A, C)
```

Arguments

A	matrix of tropical points; rows are points with the last column representing a numbered cluster assignment
C	number of clusters

Value

betweenness cluster measure

Author(s)

David Barnhill <david.barnhill@nps.edu>

References

David Barnhill, Ruriko Yoshida (2023). Clustering Methods Over the Tropically Convex Sets.

Examples

```
hars<-Sim_points
cls<-c(rep(1,50),rep(2,50),rep(3,50))
cl_pt<-cbind(hars,cls)
```

```
C<-3
over_bet_KM(cl_pt,C)
```

Points.TLineSeg	<i>Sample k equally spaced points on a max- or min-plus tropical line segment</i>
-----------------	---

Description

This function calculates k equally spaced points on a tropical line segment

Usage

```
Points.TLineSeg(D1, D2, k = 20, tadd = max)
```

Arguments

D1	point in the tropical projective torus
D2	point in the tropical projective torus
k	number of points
tadd	function; max indicates max-plus addition, min indicates min-plus addition. Defaults to max

Value

matrix of k equally spaced points on a tropical line segment

Author(s)

Ruriko Yoshida <ryoshida@nps.edu>

Examples

```
D1 <-c(0,4,2)
D2 <- c(0,7,-1)
Points.TLineSeg(D1, D2, k = 5)
Points.TLineSeg(D1, D2, k = 5,tadd=min)
```

`pre.pplot.pro`*Projections of points onto a tropical triangle*

Description

This function produces the a matrix of points projected onto a tropical triangle defined by the column space of a matrix

Usage

```
pre.pplot.pro(S, D)
```

Arguments

S matrix of points representing a tropical polytope; rows are the vertices
D data points in the tropical projective torus

Value

matrix of points representing projections of the points in D (row vectors) onto S

Author(s)

Ruriko Yoshida <ryoshida@nps.edu>

Examples

```
s <- 3 #number of vertices. Here it is a tropical triangle
d <- 3 ## dimension
N <- 100 ## sample size
D <- matrix(rep(0, N*d), N, d)
D[, 1] <- rnorm(N, mean = 5, sd = 5)
D[, 2] <- rnorm(N, mean = -5, sd = 5)
D[, 3] <- rnorm(N, mean = 0, sd = 5)

index <- sample(1:N, s)
S <- D[index,]

DD <- pre.pplot.pro(S, D)
```

prob.class	<i>Estimated probability for binary class assignment</i>
------------	--

Description

Estimates the probability that an observation x belongs to class 1.

Usage

```
prob.class(pars, x)
```

Arguments

pars	vector of parameters, which can be decomposed as two normal vectors and two scaling parameters and has dimension $2 * e + 2$
x	vector of dimension e

Value

real number

Author(s)

Georgios Aliatimis <g.aliatimis@lancaster.ac.uk>

References

Aliatimis, Georgios, Ruriko Yoshida, Burak Boyaci and James A. Grant (2023). Tropical Logistic Regression on Space of Phylogenetic Trees

Examples

```
library(ROCR)
T0 = Sim_Trees15
T1 = Sim_Trees25
D = rbind(T0,T1)
Y = c(rep(0,dim(T0)[1]), rep(1,dim(T1)[1]))
N = length(Y)
set.seed(1)
train_set = sample(N,floor(0.8 * N)) ## 80/20 train-test split
pars <- trop.logistic.regression(D[train_set,],Y[train_set], penalty=1e4)
test_set = (1:N)[-train_set]
Y.hat <- rep(0, length(test_set))
for(i in 1:length(test_set)) Y.hat[i] <- prob.class(pars, D[test_set[i],])
Logit.ROC <- performance(prediction(Y.hat, Y[test_set]), measure="tpr", x.measure="fpr")
plot(Logit.ROC, lwd = 2, main = "ROC Curve for Logistic Regression Model")
print(paste("Logit.AUC=", performance(prediction(Y.hat, Y[test_set]), measure="auc")@y.values))
```

`project.pi` *Project a point on the tropical projective torus onto a tropical polytope*

Description

This function projects points in the tropical projective torus onto a max- or min-plus tropical polytope based on tropical distance

Usage

```
project.pi(D_s, D, tadd = max)
```

Arguments

<code>D_s</code>	matrix where each row is a point defining a tropical polytope
<code>D</code>	point to be projected onto <code>D_s</code>
<code>tadd</code>	function; max indicates max-plus addition, min indicates min-plus addition. Defaults to max.

Value

projection of point `D` onto the tropical polytope defined by `D_s`

Author(s)

David Barnhill <david.barnhill@nps.edu>

Examples

```
D_s <-matrix(c(0,0,0,0,2,5,0,3,1),3,3,TRUE)
D <- c(0,7,-1)
project.pi(D_s,D)
project.pi(D_s,D,tadd=min)
```

`pw.trop.dist` *Constructs the dissimilarity matrix for a set of ultrametrics*

Description

Constructs the dissimilarity matrix based on the tropical distance between points in a dataset

Usage

```
pw.trop.dist(D1, D2)
```

Arguments

D1 matrix of ultrametries
 D2 matrix of ultrametries

Value

matrix; dissimilarity matrix showing the tropical pairwise distance between each point

Author(s)

Ruriko Yoshida <ryoshida@nps.edu>

References

Weyenberg, G., Huggins, P., Schardl, C., Howe, D. K., & Yoshida, R. (2014). kdetrees: Nonparametric Estimation of Phylogenetic Tree Distributions. In Bioinformatics.

Yoshida, Ruriko, David Barnhill, Keiji Miura and Daniel Howe (2022). Tropical Density Estimation of Phylogenetic Trees.

<https://github.com/grady/kdetrees/blob/master/R/dist.diss.R>

Examples

```
T1<-Sim_Trees15
T2<-Sim_Trees25
D <- rbind(T1, T2[,])
pw.trop.dist(D, D)
```

rounding

Remove all tentacles from a tropical simplex

Description

This function removes all tentacles from a tropical simplex. The remaining portion is a full-dimensional tropical polytope known as the trunk of the tropical polytope.

Usage

```
rounding(P)
```

Arguments

P matrix of points defining a tropical simplex. Rows are the points

Value

matrix of points defining only the full-dimensional element (the trunk) of a tropical polytope; rows are points

Author(s)

David Barnhill <david.barnhill@nps.edu>

References

Barnhill, David, Ruriko Yoshida and Keiji Miura (2023). Maximum Inscribed and Minimum Enclosing Tropical Balls of Tropical Polytopes and Applications to Volume Estimation and Uniform Sampling.

Examples

```
P<-matrix(c(0,-1,1,0,0,0,0,1,-1),3,3,TRUE)
BP<-min_enc.ball(P)
RP<-rounding(P)
BRP<-min_enc.ball(RP)
```

sigmoid

Sigmoid function

Description

Returns the sigmoid function valuation

Usage

```
sigmoid(x)
```

Arguments

x real number

Value

sigmoid function value at x

Author(s)

Georgios Aliatimis <g.aliatimis@lancaster.ac.uk>

References

Aliatimis, Georgios, Ruriko Yoshida, Burak Boyaci and James A. Grant (2023). Tropical Logistic Regression on Space of Phylogenetic Trees

Examples

```
sigmoid(0) # 0.5
```

Sim_points	<i>Simulated points over the tropical projective torus</i>
------------	--

Description

150 points generated using Gaussian-like Hit-and-Run sampling with three separate pairs of location and scale parameters

Usage

Sim_points

Format

Sim_points **A 150 x 3 matrix where each row is a point in the:**
tropical projective torus

Sim_Trees1	<i>Six data sets of phylogenetic trees data simulated from the Coalescant model.</i>
------------	--

Description

Six data sets of 1000 gene trees simulated from the Coalescant model based on a specified species with each data set possessing a ratio of species depth to effective population of 0.25, 0.5, 1, 2, 5, and 10.

Usage

Sim_Trees1025

Sim_Trees105

Sim_Trees11

Sim_Trees12

Sim_Trees15

Sim_Trees110

tdets	<i>Calculate the tropical determinant of a square matrix.</i>
-------	---

Description

This function calculates the tropical determinant (or singularity) of a square matrix

Usage

```
tdets(P, tadd = max)
```

Arguments

P	matrix of points defining a tropical polytope. Rows are the points
tadd	function; max indicates max-plus addition, min indicates min-plus addition. Defaults to max

Value

list containing the value of the determinant and reordered matrix P

Author(s)

David Barnhill <david.barnhill@nps.edu>

Examples

```
P<-matrix(c(0,0,0,0,2,5,0,3,1),3,3,TRUE)
tdets(P)
tdets(P,tadd=min)
```

TKmeans	<i>K-means clustering over the tropical projective torus</i>
---------	--

Description

This function performs k-means clustering over the tropical projective torus

Usage

```
TKmeans(A, C, M)
```

Arguments

A	matrix of points defining a tropical polytope; rows are the tropical points
C	number of clusters
M	maximum number of iterations of algorithm to find cluster centroids

Value

list with matrix of observation classified by centroid; matrix of centroid coordinates; number of iterations used

Author(s)

David Barnhill <david.barnhill@nps.edu>

References

David Barnhill, Ruriko Yoshida (2023). Clustering Methods Over the Tropically Convex Sets.

Examples

```
P <-Sim_points
C<-3
M<-10
res<-TKmeans(P,C,M)
try<-res[[1]]
cen<-res[[2]]
plot(try[,2],try[,3],col=try[,4],asp=1)
plot(try[,2],try[,3],col=try[,4],asp=1,xlab='x2',ylab='x3')
points(cen[,2],cen[,3],col=c('purple','hotpink','orange'),pch=19)
```

TLineSeg

Construct a max- or min-plus tropical line segment between two points

Description

This function constructs a max-plus tropical line segment between two points

Usage

```
TLineSeg(D1, D2, tadd = max)
```

Arguments

D1	point in the tropical projective torus
D2	point in the tropical projective torus
tadd	function; max indicates max-plus addition, min indicates min-plus addition. Defaults to max

Value

list of points defining the tropical line segment

Author(s)

Ruriko Yoshida <ryoshida@nps.edu>

Examples

```
D1 <-c(0,4,2)
D2 <- c(0,7,-1)
TLineSeg(D1, D2)
TLineSeg(D1, D2,tadd=min)
```

TML

Tropical Machine Learning in R

Description

TML provides a suite of tools for machine learning application on data over the tropical semiring

tree.to.vector

Phylogenetic tree to vector

Description

A tree is converted to a vector of pairwise distances between leaves. Distance between leaves is defined as the cophenetic distance between them. Normalization is applied so that the maximum distance in the vector output is 1.

Usage

```
tree.to.vector(tree, normalization = TRUE)
```

Arguments

tree phylogenetic tree
normalization logical; normalize the tree if TRUE

Value

vector of pairwise distances in $R^{\binom{m}{2}}$, where m is the number of leaves

Author(s)

Georgios Aliatimis <g.aliatimis@lancaster.ac.uk>

References

Aliatimis, Georgios, Ruriko Yoshida, Burak Boyaci, James A. Grant (2023). Tropical Logistic Regression on Space of Phylogenetic Trees

Examples

```
tree <- ape::read.tree(text='((A:1, B:1):2, (C:1.5, D:1.5):1.5);')
tree.to.vector(tree)
```

trop.ball.vert	<i>Calculate the minimum or entire generating vertex set of a tropical ball using a max- or min-plus algebra</i>
----------------	--

Description

This function calculates the coordinates of the minimum or entire vertex set of a tropical ball in terms of either a max- or min-plus algebra for a given a center point

Usage

```
trop.ball.vert(x, d, tadd = max)
trop.ball.all_vert(x, d)
```

Arguments

x	matrix where each row is a point defining a tropical polytope
d	radius of the tropical ball in terms of tropical distance
tadd	function; max indicates max-plus addition, min indicates min-plus addition, 'all' indicates all vertices. Defaults to 'max'

Value

matrix of normalized tropical points defining the tropical ball. Rows are the points

Author(s)

David Barnhill <david.barnhill@nps.edu>

References

Barnhill, David, Ruriko Yoshida and Keiji Miura (2023). Maximum Inscribed and Minimum Enclosing Tropical Balls of Tropical Polytopes and Applications to Volume Estimation and Uniform Sampling.

Examples

```
x <-c(0,3,7,5)
d <- 2
trop.ball.vert(x,d)
trop.ball.vert(x,d,tadd=min)
trop.ball.all_vert(x,d)
```

trop.dist	<i>Compute the tropical distance</i>
-----------	--------------------------------------

Description

This function computes the tropical distance between two points in the tropical projective torus

Usage

```
trop.dist(D1, D2)
```

Arguments

D1	point in the tropical projective torus
D2	point in the tropical projective torus

Value

tropical distance between D1 and D2

Author(s)

Ruriko Yoshida <ryoshida@nps.edu>

Examples

```
D1 <-c(0,4,2)
D2 <- c(0,7,-1)
trop.dist(D1, D2)
```

trop.FW	<i>Calculate the tropical Fermat-Weber point</i>
---------	--

Description

This function calculates the Fermat-Weber point for a tropical polytope

Usage

```
trop.FW(A)
```

Arguments

A	matrix with normalized tropical points as rows
---	--

Value

numeric vector providing the tropical Fermat-Weber point for the tropical polytope

Author(s)

David Barnhill <david.barnhill@nps.edu>

References

Lin, Bo and Ruriko Yoshida (2016). Tropical Fermat-Weber Points. SIAM J. Discret. Math. 32: 1229-1245.

Examples

```
P <-matrix(c(0,0,0,0,2,5,0,3,1),3,3,TRUE)
trop.FW(P)
```

trop.hyper.dist

Calculate the tropical distance to a max-tropical hyperplane

Description

Calculate the tropical distance to a max-tropical hyperplane

Usage

```
trop.hyper.dist(0, x0, tadd = max)
```

Arguments

0	normal vector of a tropical hyperplane; numeric vector
x0	point of interest; numeric vector
tadd	function; max indicates max-plus addition, min indicates min-plus addition. Defaults to max

Value

tropical distance to max-plus tropical hyperplane

Author(s)

David Barnhill <david.barnhill@nps.edu>

Examples

```
0 <-c(0, -1, -1)
x0 <- c(0, -2, -8)
trop.hyper.dist(0, x0)
trop.hyper.dist(0, x0, tadd=min)
```

```
trop.logistic.regression
```

Tropical Logistic Regression

Description

Performs tropical logistic regression, by finding the optimal statistical parameters for the training dataset (D,Y), where D is the matrix of covariates and Y is the binary response vector

Usage

```
trop.logistic.regression(D, Y, penalty = 0, model_type = "two_species")
```

Arguments

D	matrix of dimension $N \times e$, where N is the number of observations which lie in \mathbb{R}^e
Y	binary vector with dimension N, with each component corresponding to an observation
penalty	scalar; positive real number
model_type	string; options are "two-species" (default), "one-species", "general"

Value

vector; optimal model parameters (two normal vectors and two scaling factors)

Author(s)

Georgios Aliatimis <g.aliatimis@lancaster.ac.uk>

References

Aliatimis, Georgios, Ruriko Yoshida, Burak Boyaci and James A. Grant (2023). Tropical Logistic Regression on Space of Phylogenetic Trees.

Examples

```

library(ROCR)
T0 = Sim_Trees15
T1 = Sim_Trees25
D = rbind(T0,T1)
Y = c(rep(0,dim(T0)[1]), rep(1,dim(T1)[1]))
N = length(Y)
set.seed(1)
train_set = sample(N,floor(0.8 * N)) ## 80/20 train-test split
pars <- trop.logistic.regression(D[train_set,],Y[train_set], penalty=1e4)
test_set = (1:N)[-train_set]
Y.hat <- rep(0, length(test_set))
for(i in 1:length(test_set)) Y.hat[i] <- prob.class(pars, D[test_set[i],])
Logit.ROC <- performance(prediction(Y.hat, Y[test_set]), measure="tpr", x.measure="fpr")
plot(Logit.ROC, lwd = 2, main = "ROC Curve for Logistic Regression Model")
print(paste("Logit.AUC=", performance(prediction(Y.hat, Y[test_set]), measure="auc")@y.values))

```

trop.tri.plot.w.pts *Plotting PCA-derived tropical triangles*

Description

This function conducts tropical PCA to find the best fit tropical triangle given data defined in the tropical projective torus. It employs the vertex HAR with extrapolation sampler to sample points to determine the vertices of the tropical triangle.

Usage

```
trop.tri.plot.w.pts(S, D)
```

Arguments

S	initial set of vertices for the tropical triangle
D	matrix of data where each row is an observation in the tropical projective torus

Value

rendering of tropical triangle saved to current directory

Author(s)

Ruriko Yoshida <ryoshida@nps.edu>

Examples

```

s <- 3 #number of vertices. Here it is a tropical triangle
d <- 3 ## dimension
N <- 100 ## sample size
V <- matrix(c(100, 0, 0, 0, 100, 0, 0, 0, 100, -100, 0, 0, 0, -100, 0, 0, 0, -100), 6, 3, TRUE)
D <- matrix(rep(0, N*d), N, d)
D[, 1] <- rnorm(N, mean = 5, sd = 5)
D[, 2] <- rnorm(N, mean = -5, sd = 5)
D[, 3] <- rnorm(N, mean = 0, sd = 5)
index <- sample(1:N, s)
S <- D[index,]
res <- tropical.PCA.Polytope(S, D, V, I = 1000,50)
DD <- pre.pplot.pro(res[[2]], res[[3]])
trop.tri.plot.w.pts(normaliz.ultrametrics(res[[2]]), DD)

```

trop.Volume

Estimate the volume of a tropical polytope

Description

This function uses tropical HAR with a uniform target distribution to estimate the volume of a tropical polytope

Usage

```
trop.Volume(B, P, x0, s, I, r)
```

Arguments

B	matrix of points defining a minimum enclosing ball for a polytope P; rows are the points
P	matrix of points defining a tropical polytope; rows are the points
x0	initial point used for the HAR sampler
s	number of points to sample from the minimum enclosing ball
I	number of iterations for the HAR sampler
r	radius of the minimum enclosing tropical ball

Value

list containing ratio of points falling in P; volume of the tropical ball; volume estimate of P

Author(s)

David Barnhill <david.barnhill@nps.edu>

References

Barnhill, David, Ruriko Yoshida and Keiji Miura (2023). Maximum Inscribed and Minimum Enclosing Tropical Balls of Tropical Polytopes and Applications to Volume Estimation and Uniform Sampling.

Examples

```
P <-matrix(c(0,0,0,0,3,1,0,2,5),3,3,TRUE)
BR<-min_enc.ball(P)
B<-trop.ball.vert(BR[[1]],BR[[2]])
x0<-c(0,1.5,.4)
S<-200
I<-50
R<-BR[[2]]
trop.Volume(B,P,x0,S,I,R)
```

tropical.centroid *Tropical centroid-based sampling about a center of mass*

Description

This function is a centroid-based HAR sampler about a center of mass denoted by a location parameter with scale parameter in terms of the tropical distance

Usage

```
VE.HAR.centroid(D_s, x0, I = 1, m, s, tadd = max)
```

Arguments

D_s	matrix of vertices of a tropical simplex; each row is a vertex
x0	initial point for sampler, numeric vector
I	number of states in Markov chain
m	location parameter; numeric vector indicating centroid
s	scale parameter; in terms of tropical distance
tadd	function; max indicates max-plus addition, min indicates min-plus addition. Defaults to max

Value

next sampled point from the tropical polytope

Author(s)

David Barnhill <david.barnhill@nps.edu>

Examples

```
D_s <-matrix(c(0,10,10,0,10,0,0,0,10),3,3,TRUE)
x0 <- c(0,0,0)
m <- c(0,5,5)
s <- 1
VE.HAR.centroid(D_s, x0, I = 50,m,s)
VE.HAR.centroid(D_s, x0, I = 50,m,s,tadd=min)
```

tropical.centroid.MH *Centroid-based sampling using Metropolis filter*

Description

This function samples points on a tropical line segment about a location parameter for a given scale parameter defined in terms of tropical distance

Usage

```
trop.centroid.MH(D, x0, m, s, n, I = 50, tadd = max)
trop.centroid.MH.square(D, x0, m, s, n, I = 50, tadd = max)
```

Arguments

D	matrix of vertices of a tropical polytope; each row is a vertex
x0	initial point for sampler, numeric vector
m	location parameter; numeric vector
s	scale parameter; scalar
n	number of points to sample
I	states in Markov chain
tadd	function; max indicates max-plus addition, min indicates min-plus addition. Defaults to max

Value

matrix of n sampled points where each point is a row

Author(s)

David Barnhill <david.barnhill@nps.edu>

References

Yoshida, Ruriko, Keiji Miura and David Barnhill (2022). Hit and Run Sampling from Tropically Convex Sets.

Examples

```

D1 <-matrix(c(0,0,0,0,10,0,0,0,10),3,3,TRUE)
D2 <-matrix(c(0,10,10,0,10,0,0,0,10),3,3,TRUE)
x0 <- c(0,0,0)
m1<-c(0,5,5)
m2<-c(0,-1,1)
s<-1
n<-10
trop.centroid.MH(D1, x0, m1, s, n, I=50)
trop.centroid.MH.square(D1, x0,m1, s, n, I=50)
trop.centroid.MH(D2, x0, m1, s, n, I=50,tadd=min)
trop.centroid.MH.square(D2, x0,m2, s, n, I=50,tadd=min)

```

Tropical.HC.AGNES *Agglomerative (AGNES) tropical hierarchical clustering*

Description

This function performs agglomerative (AGNES) hierarchical clustering over the space of ultrametrics defining the space of equidistant trees

Usage

```
Tropical.HC.AGNES(D, method = mean)
```

Arguments

D matrix of points defining a tropical polytope. Rows are the tropical points
method linkage method: mean, min, or max

Value

list of distances in when merges occur; list of indices of points in each cluster

Author(s)

David Barnhill <david.barnhill@nps.edu>

References

David Barnhill, Ruriko Yoshida (2023). Clustering Methods Over the Tropically Convex Sets.

Examples

```

P <-Sim_points
Tropical.HC.AGNES(P, method=mean)

```

tropical.KDE	<i>Tropical Kernel Density Estimation of Phylogenetic Trees</i>
--------------	---

Description

This function calculates a non-parametric density estimate of a tree over the space of phylogenetic trees on m leaves. It mimics classical kernel density estimation by using a Gaussian kernel in conjunction with tropical distance.

Usage

```
tropical.KDE(D, n, sigma, h = 2)
```

Arguments

D	matrix of phylogenetic tree observations as ultrametrics
n	number of leaves for each tree
sigma	bandwidth parameter based on tropical distance
h	height of the tree

Value

list containing center point and radius of minimum enclosing ball of P

Author(s)

Ruriko Yoshida <ryoshida@nps.edu>

References

Weyenberg, G., Huggins, P., Schardl, C., Howe, D. K., & Yoshida, R. (2014). kdetrees: Nonparametric Estimation of Phylogenetic Tree Distributions. In Bioinformatics.

Yoshida, Ruriko, David Barnhill, Keiji Miura and Daniel Howe (2022). Tropical Density Estimation of Phylogenetic Trees.

Examples

```
T1<-Sim_Trees15
T2<-Sim_Trees25
D <- rbind(T1, T2[1,])
T <- dim(D)[1]
X <- 1:T
M <- pw.trop.dist(D, D)
sigma <- bw.nn(M)
P_5 <- tropical.KDE(D, n, sigma, h = 2)
Q5 <- P_5[T]
```

tropical.PCA	<i>Tropical principal component analysis (PCA) on over tropical projective torus</i>
--------------	--

Description

This function conducts tropical PCA to find the best fit tropical triangle given data defined in the tropical projective torus. It employs the vertex HAR with extrapolation sampler to sample points to determine the vertices of the tropical triangle.

Usage

```
tropical.PCA.Polytope(S, D, V, I = 1, k)
```

Arguments

S	initial set of vertices for the tropical triangle
D	matrix of data where each row is an observation in the tropical projective torus
V	matrix of vertices defining a polytope encompassing D
I	number of iterations to perform
k	number of iterations for the HAR sampler

Value

list with the sum of residuals

Author(s)

Ruriko Yoshida <ryoshida@nps.edu>

References

Page, Robert and others (2020), Tropical principal component analysis on the space of phylogenetic trees, *Bioinformatics*, Volume 36, Issue 17, Pages 4590–4598.

Yoshida, R., Zhang, L. & Zhang, X (2019). Tropical Principal Component Analysis and Its Application to Phylogenetics. *Bull Math Biol* 81, 568–597.

Examples

```
s <- 3 #number of vertices. Here it is a tropical triangle
d <- 3 ## dimension
N <- 100 ## sample size
V <- matrix(c(100, 0, 0, 0, 100, 0, 0, 0, 100, -100, 0, 0, 0, -100, 0, 0, 0, -100), 6, 3, TRUE)
D <- matrix(rep(0, N*d), N, d)
D[, 1] <- rnorm(N, mean = 5, sd = 5)
D[, 2] <- rnorm(N, mean = -5, sd = 5)
D[, 3] <- rnorm(N, mean = 0, sd = 5)
```

```

index <- sample(1:N, s)
S <- D[index,]
DD <- pre.pplot.pro(S, D)
for(i in 1:N)
  DD[i, ] <- normaliz.vector(DD[i, ])

res <- tropical.PCA.Polytope(S, D, V, I = 1000,50)
DD <- pre.pplot.pro(res[[2]], res[[3]])
trop.tri.plot.w.pts(normaliz.ultrametrics(res[[2]]), DD)

```

Trop_ball

*Visualize a Tropical ball in 2D or 3D***Description**

This function constructs a visualization of a 2D or 3D tropical ball

Usage

```

Trop_ball(
  v,
  d,
  a = 1,
  cls = "black",
  cent.col = "black",
  fil = TRUE,
  plt = TRUE,
  bord = "black"
)

```

Arguments

v	center of tropical ball; numeric vector of length 3 or 4
d	radius of tropical ball
a	shading level; 1 is opaque
cls	string indicating color of interior of ball
cent.col	string indicating color of center point
fil	logical for 3D plots; if TRUE 2D facets of 3D ball fill in color of cls parameter
plt	logical; indicates plot a new object; defaults to TRUE; if FALSE, overlays the ball on existing plot
bord	string indicating color of border of ball (only for 2D plots)

Value

2D or 3D visualization of tropical ball

Author(s)

David Barnhill <david.barnhill@nps.edu>

Examples

```
v <-c(0,0,0)
d <- 2
Trop_ball(v,d,a=.1,cls='white',cent.col='black',fil=TRUE,plt=TRUE,bord='black')
v <-c(0,0,0,0)
d <- 2
Trop_ball(v,d,a=1,cls='red',cent.col='black',fil=FALSE,plt=TRUE)
```

trop_wi_dist

Tropical within-cluster measure

Description

This function calculates a within cluster measure by measuring the pairwise tropical distance between points in the cluster.

Usage

```
trop_wi_dist(D1, method = mean)
```

Arguments

D1	matrix of tropical points; rows are points
method	function; metric to measure; mean is the average pairwise tropical distance; max is the maximum pairwise tropical distance

Value

within cluster measure

Author(s)

David Barnhill <david.barnhill@nps.edu>

References

David Barnhill, Ruriko Yoshida (2023). Clustering Methods Over the Tropically Convex Sets.

Examples

```
D<-Sim_points
avg.m<-trop_wi_dist(D, method=mean)
max.m<-trop_wi_dist(D, method=max)
```

Ultrametrics.HAR *Hit-and-Run Sampler for the space of ultrametrics*

Description

This sampler samples a point in the space of ultrametrics where each point represents an equidistant tree on n leaves

Usage

```
Ultrametrics.HAR(x0, n, I = 1, h = 1)
```

Arguments

<code>x0</code>	an equidistant tree defined as ultrametric
<code>n</code>	number of leaves for the equidistant tree
<code>I</code>	number of states in the Markov chain
<code>h</code>	height of phylogenetic tree

Value

point in the space of ultrametrics over n leaves

Author(s)

Ruriko Yoshida <ryoshida@nps.edu>

References

Yoshida, Ruriko, Keiji Miura and David Barnhill (2022). Hit and Run Sampling from Tropically Convex Sets.

Examples

```
x0 <-Sim_Trees15[1,]  
n<-10  
  
Ultrametrics.HAR(x0, n, I = 50, h = 1)
```

VE.HAR	<i>Vertex HAR with extrapolation (VHE) MCMC with uniform target distribution</i>
--------	--

Description

This function samples points uniformly the space defined by a tropical simplex

Usage

```
VE.HAR(D_s, x0, I = 1, tadd = max)
```

Arguments

D_s	matrix of vertices of a tropical simplex; each row is a vertex
x0	initial point for sampler, numeric vector
I	number of states in Markov chain
tadd	function; max indicates max-plus addition, min indicates min-plus addition. Defaults to max

Value

next sampled point from the tropical polytope

Author(s)

David Barnhill <david.barnhill@nps.edu>

References

Yoshida, Ruriko, Keiji Miura and David Barnhill (2022). Hit and Run Sampling from Tropically Convex Sets.

Examples

```
D_s <-matrix(c(0,0,0,0,10,0,0,0,10),3,3,TRUE)
x0 <- c(0,0,0)
VE.HAR(D_s, x0, I = 50)
VE.HAR(D_s, x0, I = 50,tadd=min)
```

```
vector.to.equidistant.tree
```

Vector to equidistant tree

Description

A vector of pairwise distances is used to reconstruct the corresponding equidistant tree

Usage

```
vector.to.equidistant.tree(vec)
```

Arguments

vec vector of pairwise distances in $R^{\binom{m}{2}}$, where m is the number of leaves

Value

equidistant phylogenetic tree

Author(s)

Georgios Aliatimis <g.aliatimis@lancaster.ac.uk>

References

Aliatimis, Georgios, Ruriko Yoshida, Burak Boyaci and James A. Grant (2023). Tropical Logistic Regression on Space of Phylogenetic Trees

Examples

```
vec = c(1/3,1,1,1,1,1/3)
tree = vector.to.equidistant.tree(vec)
plot(tree)
```

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