Package: zonohedra (via r-universe)

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Description Computes a zonohedron from real vector generators. The package also computes zonogons (2D zonotopes) and zonosegs (1D zonotopes). An elementary S3 class for matroids is included, which supports matroids with rank 3, 2, and 1. Optimization methods are taken from Heckbert (1985) https://www.cs.cmu.edu/~ph/zono.ps.gz.

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Description

This package deals with *zonohedra*, which are zonotopes of dimension 3. It also handles *zonogons* (2D zonotopes) and *zonosegs* (1D zonotopes).

The term zonoseg ("zonotope" + "segment") is my own personal term; I could not find an alternative term. It is a linear image of the unit cube $[0,1]^n$ in the real numbers, and a compact segment of reals.

S3 classes

```
Z class(Z)
zonohedron "zonohedron" "zonotope" "list"
zonogon "zonogon" "zonotope" "list"
zonoseg "zonoseg" "zonotope" "list"
```

For example, the section() returns very diffferent things for a zonohedron and a zonogon, and so section.zonohedron() and section.zonogon() are coded and documented separately. A section for a zonoseg does not make sense, so section.zonoseg() is undefined.

Terminology

For a convex polytope, a *supporting hyperplane* is a hyperplane that intersect the polytope's boundary but *not* its interior.

A zonotope is a convex polytope. A zonohedron has supporting planes, and a zonogon has supporting lines.

In the package **zonohedra**, a *zonotope* mean a zonotope of dimension 3, 2, or 1.

A *face* of a zonotope is the intersection of the boundary of the zonotope with some supporting hyperplane. A *d-face* is a face of dimension *d*. So a *0-face* is a *vertex*, and a *1-face* is an *edge*.

A *facet* of a zonotope is a face whose dimension is 1 less than the dimension of the zonotope. A facet is a maximal proper face.

A zonohedron has 0-faces (vertices), 1-faces (edges), and 2-faces (facets).

A zonogon has 0-faces (vertices) and 1-faces (edges). Since the dimension of an edge is 1 less than the dimension of the zonogon, an edge of a zonogon is also a facet of a zonogon.

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boundarypgramdata compute data about specific parallelograms in the boundary of a zono- hedron

Description

The boundary of a zonohedron is the union of parallelograms, where some of them may be facets, and some may be tiles in the standard tiling of more complex facets. The edges of each parallelogram are given by a pair of distinct simplified generators. If a zonohedron has n of these generators, then there are n(n-1)/2 such pairs. For each pair of generators, there are 2 parallelograms which are antipodal to each other. This function computes data about one parallelogram from this antipodal pair. The total number of parallelograms is n(n-1).

Usage

```
boundarypgramdata( x, gndpair, cube=FALSE )
```

Arguments

a **zonohedron** object as returned by the constructor zonohedron()

gndpair an Mx2 integer matrix. Each row of gndpair must contain a pair of points in

the ground set of the simplified matroid of the zonohedron x. The 1st point must

be less that the 2nd point.

gndpair can also be a numeric vector that can be converted to such a matrix, by

row.

cube if TRUE, then a point of the cube that maps to the center of the given parallelo-

gram is returned, see Value.

Value

boundarypgramdata() returns a data. frame with M rows and these columns:

gndpair the given gndpair

hyperplaneidx the index of the hyperplane in the simplified matroid of x that contains gndpair

center the center of the parallelogram relative to the center of the zonohedron. For the

antipodal parallelogram, multiply this by -1.

transitions the number of transitions in pcube - a point in the n-cube that maps to center.

This is a non-negative even integer.

And if cube is TRUE, then this column is added:

pcube a point in the n-cube that maps to center. For the antipodal parallelogram,

subtract this from 1; the number of transitions is the same.

If a row of gndpair has an invalid pair, the other columns are filled with NAs.

In case of global error, the function returns NULL.

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WARNING

In this version of the package, when the parallelogram is a tile in a more complex facet, the function may return incorrect results. This will be fixed in a future version.

See Also

zonohedron()

genlist

zonohedra generators useful for testing and plotting

Description

```
classics.genlist 13 classic zonohedra generators colorimetry.genlist 4 sets of Color Matching Functions (each set is a 3xN matrix)
```

Format

Each is an S3 class **genlist** object organized as a list of 3xN matrices (N varies). The list must have names, preferably short names or abbreviations. Each matrix can have optional attributes "shortname" and "fullname" which are useful when printing with print.genlist().

Note

Making these S3 class **genlist** makes it possible to easily print a short summary using print.genlist().

For colorimetry.genlist[[2]] a few remarks are in order. These generators come from the xyz CIE color matching functions of 1931, from 360 to 830 nm with 1 nm step. From 699 to 830 nm, the angles between the generators only differ by a few microradians, and it apparent that the designers tapered all 3 functions identically in that nm range. For an illustration of this in the chromaticity domain, see Burns, Figure 10. When the zonohedron is constructed from these 132 generators, with the default options, all these generators a 'collapsed' into a single one. In the original matroid these 132 *points* form a *multiple group*, and in the simplified matroid they are collapsed to a single *point*, labeled with 699.

Source

```
David Eppstein. Zonohedra and Zonotopes.
```

https://www.ics.uci.edu/~eppstein/junkyard/ukraine/ukraine.html

Colour & Vision Research Laboratory. University College London. http://www.cvrl.org

References

ASTM E 308 - 01. Standard Practice for Computing the Colors of Objects by Using the CIE System. Table 1

Scott A Burns. **The location of optimal object colors with more than two transitions**. Color Research & Application. Vol. 46. No. 6. pp 1180-1193. 2021.

Günther Wyszecki and W.S. Stiles. Color Science: Concepts and Methods, Quantitative Data and Formulae. Second Edition. Wiley-Interscience. 1982. Table I(3.3.1). pp. 723-735.

See Also

```
print.genlist()
```

Examples

```
# get the names of 3 sets of color matching functions
names(colorimetry.genlist)
# [1] "xyz1931.5nm" "xyz1931.1nm" "lms2000.1nm"
# print zonohedra metrics associated with 3 sets of color matching functions
colorimetry.genlist
                    fullname generators vertices edges facets
                                                               area
                                                                      volume pointed
# xyz1931.5nm xyz at 5nm step
                            81 5100 10146 5048 1582.722 4070.345 TRUE
# xyz1931.1nm xyz at 1nm step
                                    471 112910 225720 112812 39586.707 509434.149 TRUE
# lms2000.1nm lms at 1nm step 441 146642 292860 146220 22736.652 181369.085
                                                                                TRUE
# ciexyzjv.5nm xyz at 5nm step (1978)
                                            8012 16020 8010 1553.535 3951.899
                                      90
```

 $\begin{array}{ll} \textit{getImportant Metrics about a Zonohedron, and Print and Summarize} \\ & \textit{Them} \end{array}$

Description

Get some important zonohedron metrics; for most some computation is needed.

The print() function prints nicely formatted facts about a zonohedron, including its matroid.

The summary() function prints a single-line summary, formatted as a row in a data frame.

```
## S3 method for class 'zonohedron'
getmetrics( x )

## S3 method for class 'zonohedron'
print( x, trans2=FALSE, matroid=TRUE, ... )

## S3 method for class 'zonohedron'
summary( object, ... )
```

getmetrics.zonohedron 7

Arguments

x a zonohedron object as returned by the constructor zonohedron()
trans2 if TRUE then print extra metrics on the 2-transition surface associated with x
matroid if TRUE then print extra information about the matroid associated with x
object a zonohedron object as returned by the constructor zonohedron()
... for print() further arguments are ignored; for summary() the further arguments can be *more* zonohedron objects, which are summarized by adding more rows to the same data frame; see Examples.

Value

getmetrics.zonohedron() returns a list with these items:

vertices the number of vertices edges the number of edges

facets the number of facets (2D faces); all of them are zonogons

area the sum of the areas of all the facets

volume as a polytope

All of these are always positive.

print.zonohedron() returns TRUE or FALSE.

summary.zonohedron() returns a data frame, see Examples.

See Also

```
genlist, zonohedron(),
```

Examples

```
zono = zonohedron( classics.genlist[['BD']] )
zono
# zonohedron:
                                     Bilinski dodecahedron
# fullname:
# generators (original):
# generators with multiples:
                                     0
# generators (simplified):
# number of facets:
                                     12 [6 antipodal facet-pairs]
# facets that contain 0:
                                     4
                                          { 1 3 4 6 }
# number of edges:
                                     24
                                     0.809017 2.118034 1.309017
# center:
# pointed:
                                     TRUE
# salient:
                                     TRUE
# area:
                                      38.83282
# volume:
                                      16.94427
# matroid:
```

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```
# ground set:
                       4 points {1 2 3 4}
                            {1 2} {1 3} {1 4} {2 3} {2 4} {3 4}
# hyperplanes:
# rank:
# loops:
                       0
                          {}
# multiple groups:
                       0
                          {}
# uniform:
                       TRUE
# paving:
                       TRUE
# simple:
                       TRUE
# This matroid is constructed from a 3x4 real matrix.
                      2
                               3
             1
# [1,] 1.000000 1.618034 0.000000 -1.000000
# [2,] 1.618034 0.000000 1.000000 1.618034
# [3,] 0.000000 1.000000 1.618034 0.000000
summary( zono )
                fullname generators vertices edges facets
                                                              area
                                                                    volume
# 1 Bilinski dodecahedron
                                          14
                                                24
                                                       12 38.83282 16.94427
zono4 = zonohedron( classics.genlist[['RI']] )
zono7 = zonohedron( classics.genlist[['T0']] )
summary( zono, zono4, zono7 )
                fullname generators vertices edges facets
                                                              area
                                                                    volume
# 1 Bilinski dodecahedron
                                                      12 38.83282 16.94427
                               4
                                         14
                                                24
                                  5
                                                       20 64.72136 42.36068
# 2 rhombic icosahedron
                                          22
                                                40
                                                36
# 3 truncated octahedron
                                          24
                                                       14 53.56922 32.00000
                                  6
```

grpDuplicated

Grouping by duplicated elements

Description

grpDuplicated() is a generic function that takes an indexed set of "elements", and outputs an integer vector with the same length. The "elements" can be components of a vector, or the row vectors or column vectors of a matrix. In the output vector, a component is 0 if and only if the corresponding element is unique. When the element is unique, it forms a *singleton group*. Output components have equal positive integer values if and only if the corresponding elements are identical to each other. These elements form a *non-singleton group*, and the positive integer is called the *group number*.

The number of singleton groups is equal to #(zeros), which is equal to the #(elements) - #(duplicated elements).

The number of non-singleton groups is equal to max(output vector).

The number of all groups is equal to #(zeros) + max(output vector).

```
## Default S3 method:
grpDuplicated( x, ... )
```

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```
## S3 method for class 'matrix'
grpDuplicated( x, MARGIN=1, ... )
```

Arguments

x a vector or matrix of atomic mode "numeric", "integer", "logical", "complex", "character" or "raw".

MARGIN an integer scalar, the matrix margin to be held fixed, as in apply. MARGIN=1 means that it looks for duplicated rows, and MARGIN=2 means that it looks for duplicated columns. Other values are invalid.

... arguments for particular methods.

Details

The implementation is based on std::unordered_map in C++11, which uses a hash-table.

Value

The return value is an integer vector with all elements ranging from 0 to K, where K is the number of non-singleton groups.

For vector x the elements are the vector components, and the output is the same length as the input. For a matrix x with MARGIN=1, the elements are the rows of the matrix and the output has length nrow(x).

For a matrix x with MARGIN=2, the elements are the columns of the matrix and the output has length ncol(x).

The 'ngroups' attribute of the returned vector is set to an integer 3-vector. The 1st component is the total number of groups, the 2nd component is the number of singleton groups, and the 3rd component is the number of non-singleton groups K.

Note

The templated C++ function that does the real work is taken from the package **uniqueAtomMat** by Long Qu, but the returned vector is slightly modified by Glenn Davis.

Author(s)

Long Qu and Glenn Davis

Source

```
https://github.com/cran/uniqueAtomMat/
```

The package uniqueAtomMat was removed from CRAN by its author Long Qu.

Examples

```
set.seed(0)
# test a numeric vector
x = rnorm(7)
```

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```
y = rnorm(5)
grpDuplicated(c(x,y,rev(x)))
## [1] 7 6 5 4 3 2 1 0 0 0 0 0 1 2 3 4 5 6 7
## attr(,"ngroups")
  [1] 12 5 7
# test a numeric matrix, both rows and columns
A = matrix( rnorm(3*7), 3, 7)
B = matrix(rnorm(3*5), 3, 5)
    the columns of cbind(A,B,A) have the duplicates one would expect
grpDuplicated( cbind(A,B,A), MARGIN=2 )
## [1] 1 2 3 4 5 6 7 0 0 0 0 0 1 2 3 4 5 6 7
   attr(,"ngroups")
## [1] 12 5 7
# but the rows of cbind(A,B,A) are unique
grpDuplicated( cbind(A,B,A), MARGIN=1 )
## [1] 0 0 0
## attr(,"ngroups")
## [1] 3 3 0
```

inside

test points for being inside a zonotope

Description

Test points for being inside a zonotope. The boundary points are considered to be inside.

Usage

```
## S3 method for class 'zonotope'
inside( x, p )
```

Arguments

x a **zonotope** object - a **zonohedron**, **zonogon**, or **zonoseg**

p an NxM numeric matrix, where M is the dimension of the zonotope. The points to be tested are in the rows. p can also be a numeric vector that can be converted to such a matrix, by row.

Details

The given zonotope is viewed as the intersection of *slabs*; there is a slab for each hyperplane in the simplified matroid. For each slab a signed distance to boundary of the slab is computed. For points outside the slab the distance is positive, for points on the boundary, the distance is 0, and for points in the interior of the slab the distance is negative. The distance to the zonotope is computed as the maximum over all these slab distances, and the *critical hyperplane* index is recorded. A point is inside iff the zonotope distance ≤ 0 .

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Value

inside.zonotope() returns a data.frame with N rows and these columns:

p the given point

whether the point is inside the zonotope. For boundary points, inside is TRUE.

the distance from the point to the zonotope. For boundary points, distance is

0. When distance > 0, it may be larger than the true mathematical distance, so
distance is really a pseudo-distance.

idxhyper the index of the *critical hyperplane* in the simplified matroid. This is the index
of the slab where the maximum slab distance was taken. For a **zonoseg** there is
only 1 hyperplane (the empty set) so this is always 1.

If the row names of p are unique, they are copied to the row names of the output. In case of error, the function returns NULL.

See Also

```
inside2trans()
```

Examples

inside2trans

test points against a 2-transition surface

Description

This function tests points for being inside the 2-transition surface associated with a zonohedron.

```
inside2trans( x, p )
```

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Arguments

x a **zonohedron** object
p an Nx3 numeric matrix. The points to be tested are in the rows. p can also be a

numeric vector that can be converted to such a matrix, by row.

Details

If the surface has no self-intersections, the the definition of whether a point p is "inside" is fairly straightforward: it is where the linking number of p and the surface is non-zero. In fact, if it is non-zero then it must be +1 or -1. The *linking number* is analogous the *winding number* in 2D, for more discussion see **Note**.

Unfortunately, there is currently no test for whether the surface *has* self-intersections, For a bad surface with self-intersections, the linking number might be any integer. Since there is no such test, we simply use the same non-zero linking number rule always.

The computed linkingnumber is returned so that the user can apply the non-zero rule, or the evenodd rule, as appropriate for their situation. These 2 rules are analogous to the two winding number rules used for polygons in computer graphics, see **Point in polygon**.

The case where a point is *on* the surface (i.e. the distance to the surface is 0) is problematic. The linkingnumber is then undefined, and we currently set inside to be undefined as well. Thus inside should be interpreted as *strictly inside*. However, in some situations, the user may want to consider inside to be TRUE in this problematic case. Or the user may want to consider points that are within a very small epsilon of the surface, where roundoff might have occurred, to have inside=FALSE or inside=NA. So the both the computed linkingnumber and distance are returned so the user can use them to make their own definition of what "inside" means.

Value

inside2trans() returns a data.frame with N rows and these columns:

р	the given point
distance	the distance from the point to the surface. This is the true Euclidean distance, and not a "pseudo-distance" as in the case of <code>inside()</code> . If the point is on the surface, the distance should be 0 up to numerical precision.
linkingnumber	the linking number of the point and the surface. If the point is <i>on</i> the surface (distance==0), the (mathematical) linking number is undefined, and the computed linkingnumber is NA (integer).
inside	whether the point is inside the surface; a logical. This is currently set to linkingnumber != 0. If the linkingnumber is NA (integer), then inside is NA (logical).
timecalc	the time to do the calculations, in seconds

If the row names of p are unique, they are copied to the row names of the output. In case of error, the function returns NULL.

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Note

The standard definition of the *linking number* of a point and a surface uses intersections with rays, see the vignette The 2-Transition Subcomplex and the 2-Transition Surface for the precise definition. This is fine in theory, but in practice does not handle well the case when the ray intersects the boundary of a parallelogram. So this function uses an integral formula for the degree of a *linking map* that reduces to summing the signed area of a lot of spherical triangles, see **Spivak** p. 75 and **Guillemin and Pollack** p. 188.

References

Guillemin, Victor and Alan Pollack. Differential Topology. Prentice-Hall. 1974.

Point in polygon — **Wikipedia, The Free Encyclopedia**. https://en.wikipedia.org/w/index.php?title=Point_in_polygon&oldid=1139808558. 2023.

Spivak, Michael. *A Comprehensive Introduction to Differential Geometry*. Volume 1. 3rd edition. Publish or Perish. 1999.

See Also

inside()

invertboundary

invert points on the boundary of a zonohedron

Description

A zonohedron Z is the image of a linear map $[0,1]^n \to Z \subset \mathbf{R}^3$, from the n-cube to 3D space. For a point on the boundary of the zonohedron, this function computes a point in the unit cube that maps to it. All coordinates of the point in the cube are 0 or 1, except for two of them. The point is not necessarily unique.

Usage

```
## S3 method for class 'zonohedron'
invertboundary( x, point, tol=5.e-14 )
```

Arguments

Х	a zonohedron object as returned by the constructor zonohedron()
point	Mx3 matrix with points on the boundary of x in the rows. Such a matrix typically is returned by raytrace.zonohedron() or section.zonohedron(). point can also be a numeric vector that can be converted to such a matrix, by row.
tol	points that are not within tol of the boundary are skipped, see Details

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Details

Given the boundary point, the function determines the facet that contains it. The pcube coordinates of the *base vertex* of this facet are all 0 or 1, and fairly easy to determine. If the facet is a parallelogram, the other two coordinates are fairly easy to determine too. If the facet is a zonogon with K generators, with K>2, then the unknown K coordinates are calculated using invert.zonogon(). Because of floating point behaviour, coordinates can be slightly negative or slightly more than 1. After the calculation, they are clamped to [0,1].

Value

invertboundary.zonohedron() returns a data.frame with M rows and these columns:

point the given boundary point

distance signed distance to the boundary of x; for successful inversion its absolute value

 $is \le tol$

facetidx index of the facet pair that contains the point

sign sign of the facet pair; either +1 or -1

pcube a point in the unit n-cube that maps to the given boundary point; all coordinates

of pcube are 0 or 1, except for 2 of them.

transitions the number of transitions in pcube - a non-negative even integer

If a point point cannot be inverted, e.g. because distance is too large, the other columns are all NA

If the row names of point are unique, they are copied to the row names of the output. The column names of pcube are copied from the ground set of the associated matroid.

In case of global error, the function returns NULL.

See Also

zonohedron(), section.zonohedron(), raytrace.zonohedron(), invert.zonogon()

lintransform linear transformations of zonotopes, and vector matroids

Description

These functions perform straightforward linear transformations on the generators of a zonotope, and the column vectors of a vector matroid

```
## S3 method for class 'zonohedron'
lintransform( x, W )
## S3 method for class 'zonogon'
lintransform( x, W )
## S3 method for class 'matroid'
lintransform( x, W )
```

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Arguments

W

X	x can be a vector matroid object, as returned from the constructor matroid()
	that takes a matrix as input.

An invertible matrix that matches the rank of x. This invertibility is verified. We can also be a scalar; it is then replaced by that scalar multiplied by the identity matrix of the appropriate rank.

Value

If x is a zonohedron (or zonogon), lintransform(x) returns the zonohedron (or zonogon) whose generators are the generators of x with the matrix W applied on the left side. This function is optimized - it is *not* necessary to transform the generators and start all over again.

If x is a vector matroid, lintransform(x) returns the matroid whose generators are the generators of x with the matrix W applied on the left side. If x is a matroid, but *not* a vector matroid, it returns the original matroid and prints a warning message.

In case of error, e.g. invalid W, the function prints an error message and returns NULL.

References

Matroid - Wikipedia. https://en.wikipedia.org/w/index.php?title=Matroid&oldid=1086234057

See Also

```
rank(), matroid()
```

matroid matroid construction

Description

Construct a matroid from a matrix, or from explicit list of hyperplanes

```
## S3 method for class 'matrix'
matroid( x, e0=0, e1=1.e-6, e2=1.e-10, ground=NULL, ... )
## S3 method for class 'list'
matroid( x, ground=NULL, ... )
```

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Arguments

Х

x can be a numeric matrix with 3, 2, or 1 rows whose columns determine the matroid. The matrix must be either square or "wide", i.e. more columns than rows. The matrix must be *full-rank*, i.e. the rank must be equal to the number of rows, which is then the rank of the constructed matroid. Such a matroid is often called a *column matroid* or *vector matroid*.

x can also be a list of vectors of positive integers, which are thought of as sets, and are the hyperplanes of the matroid. The hyperplanes are checked that they satisfy the matroid hyperplane axioms. The rank of the constructed matroid is determined automatically, and must be 3, 2, or 1.

ground

The *ground set* of the matroid - a vector of positive integers in strictly increasing order.

When x is a matrix, length(ground) must be equal to ncol(x). The point ground[i] corresponds to the *i'th* column of x. If ground is NULL, the column names of x are converted to such a vector if possible. If this is not possible, ground is set to 1:ncol(x).

When x is a list, every set in the list must be a subset of ground. If ground is NULL, it is set to the union of all the sets in x. For technical reasons, when the rank is 1, ground is required and cannot be NULL, see **Details**.

e0

threshold, in the L^{∞} norm, for a column vector of x to be considered 0, and thus that the corresponding point in the matroid is a loop. Since the default is e0=0, by default a column vector must be exactly 0 to become a loop.

e1

threshold, in a pseudo-angular sense, for column vectors to be multiples of each other, and thus members of a group of multiple (aka parallel) points in the matroid. This tolerance is only used when the rank is 2 or 3.

e2

threshold, in a pseudo-angular sense, for the planes spanned by pairs of column vectors to be considered coincident, and thus the columns to be in the same hyperplane of the matroid. This tolerance is used when the rank is 3.

... not used

Details

It was mentioned above that the tolerances e1 and e2 are *pseudo-angular*. Specifically, vectors are normalized to the L^2 unit sphere and the distance between them is computed in the L^{∞} norm.

Matroids are well-known to have many cryptomorphic definitions, e.g. independent sets, bases, circuits, rank function, closure operator, flats, and hyperplanes. See **Matroid - Wikipedia**. In this package, matroids can only be constructed from hyperplanes, but there are functions rank() and is_independent() that can be used *after* construction.

Checking that the hyperplanes satisfy the matroid hyperplane axioms is made easier by the fact that all simple matroids of rank 3 or less are *paving matroids*, see **Paving Matroid - Wikipedia**

Rank 1 matroids are extremely simple - the loops form the single hyperplane (possibly empty),

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and the non-loops form a multiple group. If ground=NULL the non-loops are unknown, so this is why ground is required when the rank is 1.

Value

matroid() returns an object with S3 class 'matroid'.

In case of error, e.g. invalid x or computed hyperplanes, the function prints an error message and returns NULL.

Note

The ground set of positive integers should not be too sparse; otherwise performance may suffer.

When x is a matrix with 3 rows, it may happen that the computed hyperplanes do not satisfy the axioms for a matroid. In that case, the user will be prompted to try reducing tolerance e2. Getting the expected hyperplanes may require some *a priori* knowledge of the expected hyperplanes. For best results, the matrix should be given with maximum precision.

References

```
Matroid - Wikipedia.
```

```
https://en.wikipedia.org/w/index.php?title=Matroid&oldid=1086234057
```

Paving Matroid - Wikipedia.

```
https://en.wikipedia.org/w/index.php?title=Paving_matroid&oldid=1021966244
```

See Also

```
rank(), simplify(), getsimplified()
```

matroid-getters

matroid get functions

Description

get some important members of a matrix

```
## S3 method for class 'matroid'
getground( x )

## S3 method for class 'matroid'
gethyperplane( x )

## S3 method for class 'matroid'
getmatrix( x )
```

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```
## S3 method for class 'matroid'
getloop( x )
## S3 method for class 'matroid'
getmultiple( x )
```

Arguments

x a matroid object, as returned from the constructor matroid()

Value

getground() returns an vector of positive integers in strictly increasing order = the ground set of the matroid x.

gethyperplane() returns a list of vectors of positive integers = the hyperplanes of the matroid. If x is the simplification of an "original matroid", the "lmdata" attribute of the returned list is set to the *loop* and *multiple group* data of the "original hyperplanes". These hyperplanes can be recovered using unsimplify().

If x was constructed from a matrix, these hyperplanes are sorted in decreasing order by length. The non-trivial hyperplanes come first, followed by the trivial hyperplanes. A hyperplane is *trivial* iff it is independent in the matroid. For a matroid of rank 3, a hyperplane is trivial iff it has 2 points.

getmatrix() returns the matrix passed to the matroid.matrix() constructor, or NULL if the list constructor was used. The column names are labeled with the ground set.

getloop() returns an integer vector with the loops of x. If x is simple, it is the empty vector.

getmultiple() returns a list of integer vectors - the multiple groups of x. If x is simple, it is the empty list.

References

Matroid - Wikipedia. https://en.wikipedia.org/w/index.php?title=Matroid&oldid=1086234057

See Also

```
rank(), simplify(), unsimplify(), getsimplified(),
```

Examples

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matroid-props

matroid properties

Description

get some important boolean properties of a matrix, see Matroid - Wikipedia for the definitions.

Usage

```
## S3 method for class 'matroid'
is_simple( x )
## S3 method for class 'matroid'
is_uniform( x )
## S3 method for class 'matroid'
is_paving( x )
```

Arguments

Х

a matroid object, as returned from the constructor matroid()

Value

is_simple() returns a logical. A matroid is *simple* iff it has no loops and no multiple groups.

is_uniform() returns a logical. A matroid is *uniform* iff all the hyperplanes have the same size, which is the rank-1.

is_paving() returns a logical. For the definition of *paving* see **Paving Matroid - Wikipedia**. This property is important because the hyperplane axioms are fairly easy to check.

References

```
Matroid - Wikipedia. https://en.wikipedia.org/w/index.php?title=Matroid&oldid=1086234057 Paving Matroid - Wikipedia. https://en.wikipedia.org/w/index.php?title=Paving_matroid&oldid=1021966244
```

```
matroid()
```

20 minkowskisum

Minkowski sum of Two zonotopes
Titille Wall sum of Two Loneropes

Description

A zonotope can be viewed as a Minkowski sum of line segments, with one endpoint at 0. Therefore, the Minkowski sum of two zonotopes (in the same dimension) is also a zonotope.

Usage

```
## S3 method for class 'zonotope'
minkowskisum( zono1, zono2, e0=0, e1=1.e-6, e2=1.e-10, ground=NULL, ... )
## S3 method for class 'zonotope'
zono1 %+% zono2
```

Arguments

zono1	a zonotope object - a zonohedron, a zonogon, or a zonoseg
zono2	a zonotope object with the same dimension as zono1
e0	see zonohedron()
e1	see zonohedron()
e2	see zonohedron()
ground	the ground set of the returned zonotope. If ground is NULL, it is set to the ground set of zono1 followed by the ground set of zono2 translated sufficiently to not intersect that of zono1.
	not used

Details

After verifying that zono1 and zono2 are the same dimension, it takes the 2 matrices, cbinds them, and passes the new matrix to the appropriate constructor, along with the other arguments. There are no special optimizations.

Value

minkowskisum() returns a zonotope of the same dimension as zono1 and zono2.

%+% is a more convenient binary operator that calls minkowskisum(), but without the flexibility of the extra arguments.

In case of error, the function returns NULL.

References

Zonohedron - Wikipedia.

https://en.wikipedia.org/wiki/Zonohedron.

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See Also

```
zonohedron(), zonogon(), zonoseg()
```

plot2trans

plot the 2-transition surface associated with a zonohedron

Description

The 2-transition surface has the topology of a sphere and is contained in the zonohedron. All the facets are parallelograms. The surface is centrally symmetric, with the same center as the zonohedron. The surface may have self-intersections.

Usage

Arguments

X	a zonohedron object as returned by the constructor zonohedron()
type	a character string with what parts to draw. If type contains 'e', then draw the edges. If type contains 'f', then draw filled facets. If type contains 'p', then draw points at the centers of the facets.
ecol	The color to use when drawing the edges.
econc	If TRUE then draw the concave edges in red, and with extra thickness
fcol	The color to use when drawing the facets.
falpha	The opacity to use when drawing the facets.
level	An integer vector which is a subvector of $0: (M-2)$, where M is the number of simplified generators. Only the facets and edges at the specified levels are drawn. When level=NULL then <i>all</i> facets and edges are drawn. This argument does not affect the drawing of points.
normals	If TRUE then draw the unit facet normals.
both	if FALSE then draw only one half of the centrally symmetric surface. Otherwise draw both halves (the default).
bgcol	the background color
add	If TRUE then add to the current 3D plot. If there is no current 3D plot, it is an error.
	not used

Details

Facets and regular edges are drawn with rgl::quads3d(). Concave edges are drawn with rgl::segments3d(). Points are drawn with rgl::points3d().

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Value

The function returns TRUE; or FALSE in case of error.

Note

The package **rgl** is required for 3D plots. A large black point is drawn at 0, a large white point at the "white point", and a 50% gray point at the center. A line from the black point to the white point is also drawn.

See Also

```
zonohedron(), plothighertrans(), plot.zonohedron()
```

plothighertrans

plot abundant and deficient parallelograms

Description

The 2-transition surface associated with a zonohedron is a topological sphere and is contained in the zonohedron. The surface is centrally symmetric, with the same center as the zonohedron. The surface may have self-intersections. For this function, the surface is required to be strictly starshaped at the center. For the definition of *strictly starshaped* see the vignette The 2-Transition Subcomplex and the 2-Transition Surface.

The 2-transition surface is a union of parallelograms. Each parallelogram has a unit normal that defines a linear functional.

If a 2-transition parallelogram is in the *interior* of the zonohedron then the functional is not maximized on the parallelogram, and there is a corresponding similar parallelogram on the boundary of the zonohedron where the functional *is* maximized. The first parallelogram (in the surface) is called *deficient* because the functional is not maximized, and the second parallelogram (in the boundary) is called *abundant* because the number of corresponding transitions across this parallelogram is more than 2.

If the 2-transition parallelogram is on the boundary, then it is called *coincident*. The coincident parallelograms are ignored and not drawn in this function.

Because of this 1-1 correspondence between deficient parallelograms (in the 2-transition surface) and the abundant parallelograms (in the boundary of the zonohedron), the area of these two surfaces are the same.

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Arguments

Х	a zonohedron object as returned by the constructor zonohedron()
abalpha	The opacity to use when drawing the abundant parallelograms. If abalpha=0 then they are not drawn.
defcol	The color to use when drawing the deficient parallelograms
defalpha	The opacity to use when drawing the deficient parallelograms. If defalpha=0 (the default), then they are not drawn.
ecol	The color to use when drawing the edges. If ecol=NA (the default), then they are not drawn.
connections	If TRUE then draw segments between centers of the deficient parallelograms in the 2-transition surface, and centers of the the corresponding abundant parallel- ograms in the zonohedron boundary
bgcol	the background color
both	if FALSE then draw only one half of the centrally symmetric boundary. Otherwise draw both halves. This affects edges and parallelograms.
• • •	not used

Details

Connections are drawn with rgl::segments3d() and rgl::points3d(). parallelograms and edges are drawn with rgl::quads3d(). Both parallelograms and edges are drawn unlit (lit=FALSE). The parallelograms are colored by the number of transitions using the color codes in **Burns**, up to 10 transitions.

A large black point is drawn at 0, a large white point at the "white point", and a 50% gray point at the center. A line from the black point to the white point is also drawn.

Value

The function returns TRUE when successful; or FALSE in case of error.

WARNING

This function currently only works when the 2-transition surface is starshaped at the center. This excludes many of the classic zonohedra.

Note

The package **rgl** is required for 3D plots.

References

Scott A Burns. **The location of optimal object colors with more than two transitions**. Color Research & Application. Vol. 46. No. 6. pp 1180-1193. 2021.

```
zonohedron(), plot.zonohedron(), plot2trans()
```

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plotpolygon	plot the generator polygon associated with a pointed zonohedron
plotpolygon	plot the generator polygon associated with a pointed zonohedron

Description

A zonohedron is *pointed* iff there is a vector \mathbf{n} so the inner product of all the zonohedron generators with \mathbf{n} is positive. In other terminology, it is *pointed* iff there is an open halfspace that contains all the generators.

When \mathbf{n} exists, a neighborhood of 0 can be cut by a plane orthogonal to \mathbf{n} and the intersection is a polygon. Since \mathbf{n} is not unique, the polygon is only unique up to a 2D projective transformation.

Usage

```
plotpolygon( x, normal=NULL, points=TRUE, labels=TRUE )
```

Arguments

x	a zonohedron object as returned by the constructor zonohedron(). It must be pointed.
normal	the vector n to use - a non-zero numeric vector of length 3. If it is given, the validity is checked and if invalid it is an error. If it is NULL, a few canonical normals are first tested for validity. If they are invalid, then a valid one is computed.
points	If TRUE then draw the vertices of the polygon
labels	If TRUE then draw labels, taken from the ground set of the simplified matroid, near the vertices

Details

The selected normal vector \mathbf{n} is added to the title of the plot.

Value

The function returns TRUE; or FALSE in case of error.

```
zonohedron(), plot.zonohedron()
```

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print

Print Basic Facts about a Matroid

Description

The function prints a nicely formatted summary of a matroid, including the ground set, the rank, loops, multiple groups, and some boolean properties. It prints the number of hyperplanes, broken down by their size. If it is a vector matroid, and its matrix is not too large, it prints that matrix. If the matroid is not simple, it also prints the simplified matroid.

Usage

```
## S3 method for class 'matroid'
print( x, ... )
```

Arguments

x a matroid object as returned by the constructor matroid()
... further arguments ignored, but required by the generic print()

Value

The function returns TRUE or FALSE.

See Also

```
matroid()
```

print.genlist

Print Basic Metrics for Each Zonohedron Generated by the Matrices in a **genlist** object

Description

An S3 class **genlist** object is organized as a named list of 3xN matrices, when N varies. The print() method constructs a zonohedron object from each matrix and then prints some basic metrics about each zonohedron, as a data frame. If the matrix has the "fullname" attribute, it is added as a column. The names of the list are copied to the rownames of the data frame.

```
## S3 method for class 'genlist'
print( x, full=TRUE, ... )
```

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Arguments

```
x a genlist object

full if TRUE, include area and volume columns
... not used
```

Details

```
print.genlist() uses summary.zonohedron().
```

Value

The function returns TRUE or FALSE.

See Also

```
genlist, summary.zonohedron()
```

Examples

```
# print zonohedra metrics associated with 3 sets of color matching functions
colorimetry.genlist
                    fullname generators vertices edges facets
                                                                   area
                                                                            volume
# xyz1931.5nm xyz at 5nm step
                              81
                                         5100 10146 5048 1582.722
                                                                          4070.345
# xyz1931.1nm xyz at 1nm step
                                    471 112910 225720 112812 39586.707 509434.149
# lms2000.1nm lms at 1nm step
                                    441 146642 292860 146220 22736.652 181369.085
names(classics.genlist)
            "RD" "BD"
                        "RI"
                             "RHD" "RT" "TO" "TRD" "TC" "RE"
# [1] "C"
                                                                  "RH" "TI"
                                                                             "TSR"
print( classics.genlist, full=FALSE )
                                   fullname generators vertices edges facets
# C
                                                    3
                                                            8
# RD
                       rhombic dodecahedron
                                                             14
                                                                   24
                                                                         12
# BD
                      Bilinski dodecahedron
                                                    4
                                                             14
                                                                   24
                                                                         12
# RI
                        rhombic icosahedron
                                                    5
                                                             22
                                                                   40
                                                                         20
                                                    5
# RHD
                                                                   28
              rhombo-hexagonal dodecahedron
                                                            18
                                                                         12
                    rhombic triacontahedron
                                                     6
# RT
                                                             32
                                                                   60
                                                                         30
# TO
                       truncated octahedron
                                                    6
                                                            24
                                                                  36
                                                                         14
# TRD
             truncated rhombic dodecahedron
                                                    7
                                                            32
                                                                  48
                                                            48
# TC
                    truncated cuboctahedron
                                                    9
                                                                  72
                                                                         26
# RE
                   rhombic enneacontahedron
                                                   10
                                                            92
                                                                 180
                                                                         90
# RH
                 rhombic hectotriadiohedron
                                                            134
                                                                  264
                                                   12
                                                                        132
                truncated icosidodecahedron
# TI
                                                   15
                                                            120
                                                                 180
                                                                         62
# TSR truncated small rhombicosidodecahedron
                                                   21
                                                            240
                                                                  360
                                                                        122
```

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rank

Rank and Independence

Description

calculate the rank of any subset of a matroid, or determine whether any subset is independent

Usage

```
rank( x, subs )
is_independent( x, subs )
```

Arguments

x a matroid object, as returned from the constructor matroid()
subs a list of integer vectors, representing subsets of the ground set of x. subs can
also be an integer vector, which is put into a list of length 1.

Value

rank(x, subs) returns an integer vector with the same length as the list subs. The i'th value is the rank of the i'th set in subs. If a set is not a subset of the ground set of x, the value is NA, and a warning message is printed.

is_independent(x, subs) returns a logical vector with the same length as the list subs. The i'th value is the independence of i'th set in x. It is equal to TRUE iff the rank of the subset is equal to the cardinality of the subset.

For both functions the names are copied from input to output.

References

Matroid - Wikipedia. https://en.wikipedia.org/w/index.php?title=Matroid&oldid=1086234057

See Also

```
matroid()
```

Examples

```
# make a matroid with rank 3
mat = matroid( classics.genlist[['RT']] )
# the ground set itself should have rank 3
rank( mat, getground(mat) )
## [1] 3
```

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```
# single points should have rank 1 (there are no loops)
rank( mat, as.list(getground(mat)) )
## [1] 1 1 1 1 1 1
# all hyperplanes should have rank 2
rank( mat, gethyperplane(mat) )
## [1] 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2
# a point not in the ground set should have rank NA
# and the emtpy set should have rank 0
rank( mat, list(100L,integer(0)) )
## 1 of 1 subsets are not a subset of ground.
## [1] NA 0
```

raytrace2trans

compute the intersection of a ray and the 2-transition surface associated with a zonohedron

Description

The open ray with basepoint b and non-zero direction d is the set of the form b + td where t > 0.

This function computes the intersection of an open ray and the 2-transition surface associated with a zonohedron. The linking number of the surface and b must be ± 1 . This is verified at the beginning, and if not true, then it is an error. The linking number condition implies that an intersection exists for every ray based at b. Note also that the condition implies that b is not on the surface. For discussion of uniqueness, see **Details**. For the definition of *linking number* see The 2-Transition Subcomplex and the 2-Transition Surface.

The 2-transition surface is a union of parallelograms. The surface is symmetric about the center of the zonhedron, so each parallelogram has an antipodal parallelogram. Each parallelogram is specified by an ordered pair of distinct generators from the *simplified* matroid associated with the zonohedron. Thus, if there are N generators, there are N(N-1) parallelograms. Swapping the generators of a parallelogram changes it to the antipodal parallelogram.

The 2-transition surface has two *poles* - the point 0 and the sum of all the generators. It is OK for the ray to pass through one of these poles.

Usage

```
raytrace2trans( x, base, direction, invert=FALSE, plot=FALSE, tol=1.e-12, ... )
```

Arguments

x a **zonohedron** object as returned by the constructor zonohedron()

base a numeric 3-vector - the basepoint of all the rays. The surface must be strictly starshaped at base, and this is verified.

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direction a numeric Mx3 matrix with M non-zero directions in the rows. The basepoint

and these directions define M rays.

direction can also be a numeric vector that can be converted to such a matrix,

by row.

invert if TRUE, then compute a point in the unit cube that maps to the point on the

2-transition surface associated with x, and add it as a column in the returned

 ${\tt data.frame}$

plot if TRUE, the computed rays, up to the boundary, are *added* to an existing plot

of the zonohedron x, see plot.zonohedron(). The segments are drawn in the

color red. If there is no open 3D plot, a warning is issued.

tol the tolerance for being strictly starshaped, and for intersection with a *pole*.

... not used

Details

The function is designed for the situation when the intersection of the ray and the surface exists and is unique. This is guaranteed for all ray directions d when the surface is strictly starshaped at b. This condition is checked at the beginning of the function, and if false then a warning is issued that the intersection point may not be unique. For the definition of *strictly starshaped* see The 2-Transition Subcomplex and the 2-Transition Surface.

For finding a parallelogram of intersection, a brute-force search is used; all parallelograms are searched until the first one that intersects the ray is found. To speed things up, the 3D problem is reduced to 2D, and the search is programmed in plain C.

If plot is TRUE, the rays are drawn in red using rgl::segments3d() and rgl::points3d().

Value

raytrace2trans() returns a data. frame with M rows and these columns:

base the given basepoint - this is the same in every row

direction the given direction

gndpair the 2 generators of the parallelogram that the ray intersects, taken from the

ground set of the simplified matroid. If the ray passes through a pole, both

of these are NA.

alpha the 2 coordinates of the intersection point within the parallelogram

ray parameter of the intersection with the parallelogram, always positive point the point on the surface; the intersection of the ray and the parallelogram

iters the number of parallelograms searched, until the desired one was found. If the

ray intersects a pole, this is 0.

timetrace the computation time for the given ray, in seconds. This does not include the

initial preprocessing time.

And if invert is TRUE, then this column is added:

pcube a point in the unit cube that maps to point. This point in the cube always has 2

transitions.

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If base and direction in a row cannot be processed, the rest of the row is NA.

If the row names of direction are unique, they are copied to the row names of the output.

In case of error, the function returns NULL.

Note

The package **rgl** is required for 3D plotting.

See Also

```
zonohedron(), plot.zonohedron(), section2trans(), raytrace.zonohedron()
```

section2trans	compute the intersection of a plane and the 2-transition surface associated with a zonohedron

Description

In general, the 2-transition surface may be highly non-convex, possibly with self-intersections. The intersection of a plane and the 2-transition surface is a union of polygons, possibly with self-intersections and intersecting each other. This function computes one of those polygons. If there are other polygons, it issues a warning and does not try to compute them.

Usage

```
section2trans( x, normal, beta, invert=FALSE, plot=FALSE, tol=1.e-12, ... )
```

Arguments

Х	a zonohedron object as returned by the constructor zonohedron()
normal	a non-zero numeric 3-vector - the normal of all the planes
beta	a numeric M-vector of plane constants. The equation of the k'th plane k is: $< x, normal > = beta[k]$.
invert	if TRUE, then compute a point in the unit cube that maps to the point on the 2-transition surface, and add it as a column in the returned data.frame
plot	if TRUE, the polygons formed by the the intersection of the planes and the 2-transition surface. <i>added</i> to an existing 3D plot of the zonohedron x, see plot.zonohedron(). The polygons are drawn in red.
tol	a small positive number, used as the tolerance for the plane intersecting the interior of each parallelogram, see Details .
	not used

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Details

The function is designed for the situation when the intersection of a plane and the surface is a single polygon.

Given a plane, the function finds all the parallelograms of the surface whose interiors intersect the plane. Each intersection is a line segment. For each parallelogram it associates one of the endpoints of the segment. The parallelograms are put in polygon order by picking an arbitrary one as the starting point, and then "marching" from one to the next using the canonical parallelogram adjacency relation. After returning to the starting point, if there are other parallelograms remaining, it means that there are other polygons in the section and a warning is issued.

Value

section2trans() returns a list of length M (=length(beta)), and the i'th item in the list is a data frame with these columns:

point a Px3 matrix with the P points of the i'th polygon in the rows. If the plane does

not intersect the 2-transition surface, then P=0 and the matrix has 0 rows. The row names of point are the indexes of the facets that contain the vertices of the

polygon; see Details.

gndpair the 2 indexes from the ground set that generates the parallelogram containing

point. See **Details** for a description of the "marching parallelogram" procedure.

And if invert is TRUE, then this column is added:

pcube a point in the unit cube that maps to point. This point in the cube always has 2

transitions.

The names of the returned list are readable strings that contain normal and beta[i].

In case of error, the function returns NULL.

Note

The package **rgl** is required for 3D plotting.

See Also

zonohedron(), plot.zonohedron(), section.zonohedron(), raytrace2trans()

simplify simplify and unsimplify

Description

A *simple matroid* has no loops and no multiple groups. Simplification is the process of removing all loops, and every point except one from each multiple group. The result is a simple matroid. The functions below simplify a matroid, or an explicit list of hyperplanes.

The hyperplanes can be *unsimplified* if the original loops and multiple groups are known.

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Usage

```
## S3 method for class 'matroid'
getsimplified( x, ... )

## S3 method for class 'list'
simplify( x, ground=NULL, ... )

## S3 method for class 'list'
unsimplify( x, loop=NULL, multiple=NULL, ground=NULL, ... )
```

Arguments

x can be a matroid object, as returned from the constructor matroid()

x can also be a list of vectors of positive integers, which are thought of as sets. All must be subsets of ground. They do not have to satisfy the matroid hyperplane axioms. For a definition of *loop* and *multiple group* in this case, see

Details.

ground The ground set of the sets in x (both simplify() and unsimplify()). It must be

a vector of positive integers in strictly increasing order (not verified). If ground

is NULL, it is set to the union of the sets in x.

loop a vector of positive integers, the loops, to add to the list x; loop must be disjoint

from ground (verified). If loop is NULL, the function looks for loop in the attribute data attr(x, 'lmdata'). If there is no such attribute, loop is set to the

empty set.

multiple a list of vectors of positive integers, the multiple groups, to add to the list

x; these groups must be pairwise disjoint and disjoint from loop (not verified). Each group must intersect the ground set in exactly one point (verified). If multiple is NULL, the function looks for multiple in the attribute data attr(x,'lmdata'). If there is no such attribute, multiple is set to the

empty list.

... not used

Details

First consider the case when x is a list of vectors of positive integers. Each vector represents a subset of the ground set. They are not required to satisfy the hyperplane axioms, but by abuse of language we will call them hyperplanes in this paragraph. A *loop* is a point (an integer in the ground set) that is in every hyperplane. Imagine now that all loops have been removed. Say that two points p and q are *multiples* iff for every hyperplane H, $p \in H$ iff $q \in H$. This is an equivalence relation, and the *multiple groups* are the equivalence classes with more than one point. For computation it is convenient to think of a boolean *incidence matrix*. There is a column for each point in the ground set, and a row for each hyperplane. An entry is TRUE iff the point is in the hyperplane. A *loop* is then a column of all TRUEs. A *multiple group* is a maximal set of duplicate columns. This is basically how simplify() is implemented, except with optimizations that avoid computing the very large incidence matrix.

Now consider the case when x is a matroid object. When x was constructed, the simplification

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of x was computed (with help from the *previous* simplify()) and stored as a member of x (unless x was already simple). So in this case getsimplified(x) does not do any real work and only takes microseconds.

These functions are accelerated with C/C++.

Value

If x is a matroid, getsimplified(x) returns x when x is simple, and a member of x when x is not simple. It does not do any real work.

If x is a list, simplify(x) returns a list of the same length, but with all loops removed, and every point except one from each multiple group removed. The integer that remains is the smallest one in the group. The order of the sets is preserved. It also sets the 'lmdata' attribute of the returned list to a list of 2 objects - the loop and multiple group data found in x.

If x is a list, unsimplify(x) returns a list of the same length, but with the loops and multiples added back. The order of the sets is preserved.

In case of error, e.g. invalid x etc., the function prints an error message and returns NULL.

References

Matroid - Wikipedia. https://en.wikipedia.org/w/index.php?title=Matroid&oldid=1086234057

See Also

```
rank(), matroid()
```

Examples

```
# an example using simplify.list() and unsimplify.list()
# get the matrix for CIE XYZ at 5 nm step size
mat3x81 = colorimetry.genlist[[1]]

# create the matroid
mat5 = matroid( mat3x81 )

# test for simplicity
is_simple(mat5)
## [1] FALSE

# get the list of hyperplanes, and simplify
hyper = gethyperplane( mat5 )
hypersimple = simplify( hyper )

# print the loop and multiple data found
attr(hypersimple, 'lmdata')

# unsimplify and compare to the originals
# the list attr(hypersimple, 'lmdata') is 'secretly' used in unsimplify()
```

34 spherize

```
identical( unsimplify(hypersimple), hyper )
## [1] TRUE
```

spherize

spherize a zonotope

Description

The input is a zonotope with a best-fit ellipsoid (or ellipse for a **zonogon**) with axes that may have very different lengths. The function computes a *spherizing matrix* W, and then transforms the zonotope so its boundary is close to a sphere.

Usage

```
## S3 method for class 'zonotope'
spherize( x, method='ZCA', ... )
```

Arguments

x a zonotope object - a **zonohedron**, a **zonogon**, or a **zonoseg**.

method for computing the matrix W, either 'ZCA' or 'PCA-COR'. Matching is partial and

case-insensitive.

... not used

Details

The 2 methods are taken from Kessy, et. al..

Value

After computing the matrix W, the function return lintransform(x, W). The "sphering" attribute is set to W.

If x is a 1D **zonoseg**, sphering is not really possible, so the function prints a warning message and returns x. In case of error, the function returns NULL.

References

Agnan Kessy, Alex Lewin, Korbinian Strimmer. **Optimal whitening and decorrelation**. https://arxiv.org/abs/1512.00809 v4 2016.

```
lintransform()
```

support 35

Description

Compute the classical support function for a zonotope. It also computes a point on the boundary where the linear functional is maximized, and the dimension of the face where the supporting hyperplane intersects the zonotope.

Usage

```
## S3 method for class 'zonotope'
support( x, direction, tol=5.e-15 )
```

Arguments

x a zonotope object - a **zonohedron**, a **zonogon**, or a **zonoseg**

direction an NxM matrix with N directions in the rows. If x is a **zonohedron**, M must be

3. If x is a **zonogon**, M must be 2. If x is a **zonoseg**, M must be 1. direction can also be a vector that can be converted to such a matrix, by row. The direction

is normal to the supporting hyperplane

tol the tolerance for determining whether the supporting hyperplane intersects a

face with positive dimension. This does not affect the value of the support func-

tion. For a zonoseg, tol is ignored.

Value

The function returns a data. frame with N rows and these columns:

direction the given direction

value the value of the support function of x, in the given direction

argmax a point on the boundary of x where the functional max is taken. This point is the

center of the face where the supporting hyperplane intersects the zonotope.

dimension of the face where the supporting hyperplane intersects the zonotope. 0 means a

vertex, 1 means an edge, and 2 means a 2-face.

If direction is 0, the other columns are NA. If the rownames of direction are unique, they are copied to the row names of the output.

In case of error, the function returns NULL.

References

Wikipedia - Support function

```
https://en.wikipedia.org/wiki/Support_function
```

```
zonoseg()
```

36 symmetrize

|--|

Description

The input is a zonotope whose matroid is simple. The function adds new generators that creates a new zonotope that is a translate of the original, and has center of symmetry at 0.

Usage

```
## S3 method for class 'zonotope'
symmetrize( x, e0=0, e1=1.e-6, e2=1.e-10, ... )
```

Arguments

x	a zonotope object - a zonohedron , a zonogon , or a zonoseg . this zonotope must be simple.	The matroid of
e0	see zonohedron()	
e1	see zonohedron()	
e2	see zonohedron()	
	not used	

Details

Each generator g (a column of the matrix) is replace by 2 generators: g/2 and -g/2. The new set of generators correponds to a *star* at 0, from Sec 2-8 of **Coxeter**.

The new ground points are obtained by translating the original ground points by the their maximum.

Value

The function returns a zonotope that is a translate of the original, and has center of symmetry at 0. In case of error, the function returns NULL.

References

Coxeter, H.S.M. Regular Polytopes. Dover Publications. 1973.

```
zonohedron(), zonogon(), zonoseg()
```

transitionsdf 37

transitionsdf	summarize the number of transitions and associated data, over all parallelograms in the boundary of a zonohedron

Description

The 2-transition surface is a union of parallelograms. For this function, the surface is required to be strictly starshaped at the center. For the definition of *strictly starshaped* see The 2-Transition Subcomplex and the 2-Transition Surface.

Each parallelogram has a unit normal that defines a linear functional.

If the 2-transition parallelogram is in the *interior* of the zonohedron then the functional is not maximized on the parallelogram, and there is a corresponding similar parallelogram on the boundary of the zonohedron where the functional *is* maximized. The first parallelogram (in the surface) is called *deficient* because the functional is not maximized, and the second parallelogram (in the boundary) is called *abundant* because the number of corresponding transitions across this parallelogram is more than 2. The difference between the functional values is called the *deficit*.

If the 2-transition parallelogram is on the boundary, then it is called *coincident*. It is also called *non-deficient* and the deficit is 0.

Usage

```
transitionsdf( x, trans2=TRUE )
```

Arguments

X	a zonohedron	object as	returned	by the	constructor	<pre>zonohedron().</pre>	The 2-
	4	1	441	4 1	1		

transition surface must be strictly starshaped.

trans2 if TRUE, then include metrics on the non-deficient (coincident) parallelograms,

with 2 transitions. This is always the first row of the returned data frame.

if FALSE, then data on the non-deficient parallelograms is not included, and the returned data frame only has data on the deficient parallelograms, with more

than 2 transitions.

Value

transitionsdf() returns a data.frame with a row for each number of transitions found, plus a final row with totals on appropriate columns. The columns are:

transitions	the number of transitions, a positive even integer, in increasing order.
parallelograms	the number of parallelograms with the given number of transitions
area	the min and max of the area of the parallelograms with the given number of transitions
area.sum	the total area of the parallelograms with the given number of transitions

38 zonogon

deficit the min and max of the deficit of the parallelograms with the given number of	parallelograms with the given number of
---	---

transitions. When there are 2 transitions the deficit should be exactly 0, but is usually slightly non-0 due to truncation. When there are more than 2 transitions

the deficit is positive.

example the 2 generators (from the ground set of the simplified matroid) of the parallel-

ogram with the maximum area

In case of error, the function returns NULL.

Note

Because of the 1-1 correspondence between similar parallelograms, the surface areas of the 2-transition surface and the boundary of the zonohedron are equal.

See Also

```
zonohedron(), plot.zonohedron()
```

zonogon

zonogon construction

Description

Construct a zonogon from a numeric matrix with 2 rows.

Usage

```
zonogon( mat, e0=0, e1=1.e-6, ground=NULL ) polarzonogon( n, m=n, ground=NULL )
```

mat	a numeric 2xM matrix, where $2 \le M$. The matrix must have rank 2 (verified). The M columns are the generators of the zonogon.
e0	threshold for a column of mat to be considered 0, in the L^{∞} norm. Since the default is e0=0, by default a column must be exactly 0 to be considered 0.
e1	threshold, in a pseudo-angular sense, for non-zero column vectors to be multiples of each other, and thus members of a group of multiple (aka parallel) points in the associated matroid. It OK for a column to be a negative multiple of another.
ground	The <i>ground set</i> of the associated matroid of rank 2 - an integer vector in strictly increasing order, or NULL. When ground is NULL, it is set to 1:ncol(mat). If ground is not NULL, length(ground) must be equal to ncol(mat). The point ground[i] corresponds to the <i>i'th</i> column of mat.

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n	an integer ≥ 3 . The generators are computed as n equally spaced points on the unit circle, starting at $(1,0)$.
m	an integer with $2 \le m \le n$. When $m < n$, only the first m points are used as generators of the zonogon.

Details

polarzonogon() is useful for testing. The term *polar zonogon* is my own, and based on the *polar zonohedron* in *Chilton & Coxeter*. It it loads the matrix mat and passes it to zonogon(). When m=n the zonogon is a regular 2n-gon. When m<n the zonogon is a has 2m vertices, but is not necessarily regular. The generators correspond to the n'th-roots of unity.

Value

zonogon() and polarzonogon() return a list with S3 class 'zonogon'. In case of error, e.g. invalid mat, the functions print an error message and returns NULL.

Note

The ground set of positive integers should not be too sparse; otherwise performance may suffer.

References

B. L. Chilton and H. S. M. Coxeter. **Polar Zonohedra**. The American Mathematical Monthly. Vol 70. No. 9. pp. 946-951. 1963.

See Also

```
zonohedron(), zonoseg(),
```

zonogon-getmetrics

get important metrics about a zonogon, and print basic facts about a zonogon.

Description

Get some important zonogon metrics; some computation is used. Also print the data, and more.

Usage

```
## S3 method for class 'zonogon'
getmetrics( x )
## S3 method for class 'zonogon'
print( x, ... )
```

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Arguments

```
x a zonogon object ... not used
```

Details

print.zonogon() prints some basic information about the zonogon, and the associated matroid.

Value

```
getmetrics() returns a list with these items:
```

vertices the number of vertices

perimeter the sum of the lengths of the all edges

area as a polygon

All of these are always positive.

print.zonogon() returns TRUE or FALSE.

See Also

```
zonogon(), getmetrics.zonohedron()
```

zonogon-invert

invert points in a zonogon

Description

A zonogon Z is the image of a linear map $[0,1]^n \to Z \subset \mathbb{R}^2$, from the n-cube to the plane. For a point in a zonogon, this function finds a point in the unit cube that maps to it. There are infinitely many such points in general (unless n=2), but this function picks a specific point using the standard tiling, see **Details**.

Usage

```
## S3 method for class 'zonogon'
invert( x, z, tol=0, plot=FALSE, ... )
```

Arguments

X	a zonogon object as returned by the constructor zonogon()

z a numeric Mx2 matrix, with M points in the rows. z can also be a numeric vector

that can be converted to such a matrix, by row.

tol points that are within tol of the boundary are still processed, see **Details**

plot if TRUE then the points in z are *added* to an existing plot of the zonogon x, using

a red X symbol, see plot.zonogon(). If there is no plot open, a warning is

issued.

... not used

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Details

The given points are first tested for being inside the zonogon, using inside() and the given tol. If any are outside, a warning is issued. When the corresponding point pcube is computed, it is clamped to the unit cube, so the inversion error may be as large as tol.

Inversion is not unique in general. For this function, the *standard tiling* of the zonogon by parallelograms is computed; it is an example of a *zonotopal tiling*. It is a *regular zonotopal tiling* because it arises from the projection of a zonohedron onto the plane, see *Ziegler*. The function plot.zonogon() has an option to plot this tiling. Given the point z, the function determines a parallelogram that contains the point. The pcube coordinates of the *base* of this parallelogram are all 0 or 1, and the coordinates of z *within* the parallelogram are in [0,1]. Thus, all coordinates of pcube are 0 or 1, except possibly for 2 of them.

Value

invert.zonogon() returns a data.frame with M rows and these columns:

z the given point

pcube a point in the unit cube that maps to z. Every pcube has all coordinates 0 or 1,

except possibly for the 2 given by hyper, see **Details**.

hyper the 2 indexes of the generators of the parallelogram that contains z, in the sim-

plified matroid. These 2 coordinates in pcube are not 0 or 1 in general.

hyperidx the index of the parallelogram that contains z

If a point z cannot be inverted, the other columns are all NA, and a warning message is printed.

If the row names of z are unique, they are copied to the row names of the output. The column names of pcube are copied from the ground set of the associated matroid.

In case of error, the function returns NULL.

References

Ziegler, G.M. Lectures on Polytopes. Graduate Texts in Mathematics. Springer New York. 2007.

See Also

```
zonogon(), inside(), plot.zonogon()
```

```
# make a zonogon with 5 generators
pz20 = polarzonogon( 20, 5 )

# make 7 random points in the zonogon
set.seed(0)
pcube = matrix( runif(5*7), 5, 7 )
z = t( getmatrix(pz20) %*% pcube )

# invert these 7 points back to the cube
```

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```
invert( pz20, z )
                          pcube.1
                                     pcube.2
                                                pcube.3
                                                           pcube.4
                                                                       pcube.5
     z.1
                   z.2
# 1 2.0676319 1.6279807 0.00000000 0.70030526 1.00000000 1.00000000 0.01553241
# 2 2.4031738 1.9658035 0.00000000 0.96572153 1.00000000 1.00000000 0.28450140
# 3 0.9230336 1.0885446 0.00000000 0.00000000 0.39548689 1.00000000 0.04948838
# 4 2.5242122 1.7395069 0.16540765 1.000000000 1.000000000 1.00000000 0.03542132
# 5 2.2598725 1.0601592 0.38111324 1.00000000 1.00000000 0.20192029 0.00000000
# 6 1.1387813 1.2636700 0.00000000 0.00000000 0.65250505 1.00000000 0.07478012
# 7 1.6315341 1.0777737 0.00000000 0.64210923 1.00000000 0.36039509 0.00000000
    hyper.1 hyper.2 hyperidx
# 1
         2
                 5
# 2
         2
                 5
                          7
# 3
         3
                 5
                           9
         1
                 5
                           4
# 5
         1
                 4
                          3
# 6
         3
                 5
                          9
# 7
         2
```

zonogon-plot

plot a zonogon

Description

Plot a zonogon object, with many options.

Usage

X	a zonogon object as returned by the constructor zonogon()
orientation	if TRUE then draw the edges with orientation arrows. Otherwise just draw unoriented line segments.
normals	if TRUE then draw an outward-pointing unit normal on each edge
elabels	if TRUE then label each edge with its generator
tiling	if TRUE then draw the standard tiling of the zonogon by parallelograms
tlabels	if TRUE then label each parallelogram in the tiling with its generators. If tiling is FALSE then this is ignored.
trans2	if TRUE then draw the image of the 2-transition subcomplex of the unit cube $[0,1]^n$, in the color blue.
	trans2 can also be an integer 2-vector defining a range of levels of the subcom-
	plex, where the $level$ of a vertex of the n -cube is the number of 1s. Both integers
	should be between 0 and n .

zonogon-raytrace 43

```
trans2type which part of the 2-transition subcomplex to draw. It can be 'BP' for bandpass (aka Type 1), 'BS' for bandstop (aka Type 2), or 'both' for both.
...
```

Details

A white dot is plotted at the center of the zonogon. A suitable is title is added above the plot. If the zonogon was returned from spherize.zonotope() the string "[spherized]" is added to the title.

Value

The function returns TRUE; or FALSE in case of error.

See Also

Description

The open ray with basepoint b and non-zero direction d is the set of the form b + td where t > 0.

This function computes the intersection of an open ray and the boundary of a zonogon Z. The basepoint is normally required to be in the interior of Z, but an exception is made if the basepoint is 0, and on the boundary of Z, and the direction points into the interior of Z. In these two cases the intersection of the open ray and the boundary of Z is unique. In the second case, the basepoint is also allowed to be the sum of all the generators - the so-called *white point* of Z.

Usage

```
## S3 method for class 'zonogon'
raytrace( x, base, direction, plot=FALSE, ... )
```

x	a zonogon object as returned by the constructor zonogon()
base	a numeric 2-vector - the basepoint of all the rays. base must either be in the interior of x, or 0 or the <i>white point</i> and on the boundary of x.
direction	a numeric Mx2 matrix with M non-zero directions in the rows. The basepoint and these directions define M rays. direction can also be a numeric vector that can be converted to such a matrix, by row.
plot	if TRUE, the computed rays, up to the boundary, are <i>added</i> to an existing plot of the zonogon x, see plot.zonogon(). The segments are drawn in the color red. If there is no open plot, a warning is issued. not used
• • •	not used

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Value

raytrace.zonogon() returns a data.frame with M rows and these columns:

base the given basepoint - this is the same in every row

direction the given direction

facetidx the index of the facet (an edge) where ray exits the zonogon

sign of the facet, either +1 or -1

ray parameter of the intersection with the exit facet, always positive point the point on the boundary; the intersection of the ray and the facet

timetrace the computation time, in seconds

If base and direction in a row cannot be processed, the rest of the row is NA.

If the row names of direction are unique, they are copied to the row names of the output.

In case of error, the function returns NULL.

make a zonogon with 5 generators

See Also

```
zonogon(), plot.zonogon(), section.zonohedron()
```

```
pz20 = polarzonogon( 20, 5 )
   make 4 random directions
set.seed(0)
dir = matrix(rnorm(4*2), 4, 2)
# use basepoint in the interior of the zonogon
raytrace( pz20, c(0.5,0.5), dir )
# base.1 base.2 direction.1 direction.2 facetidx sign
                                                    tmax boundary.1 boundary.2 timetrace
# 1 0.5 0.5 1.2629543 0.4146414 4 -1 2.0503073 3.08944438 1.35014236 7.680000e-05
# 2 0.5 0.5 -0.3262334 -1.5399500 1 -1 0.3246859 0.39407664 0.00000000 4.649995e-05
    0.5 0.5 1.3297993 -0.9285670 2 -1 0.4868719 1.14744192 0.04790678 4.310103e-05
    0.5 0.5 1.2724293 -0.2947204 2 -1 0.9354693 1.69031851 0.22429808 4.149997e-05
# use basepoint at 0 - on the boundary of the zonogon
raytrace( pz20, c(0,0), dir )
# base.1 base.2 direction.1 direction.2 facetidx sign tmax boundary.1 boundary.2 timetrace
           0 1.2629543 0.4146414 4 -1 2.192481 2.7690037 0.9090936 0.0001216
# 1
# 2
            0 -0.3262334 -1.5399500
                                        NA NA
                                                 NA
                                                             NA
                                                                      NA
                                                                               NA
# 3
       0
          0 1.3297993 -0.9285670 NA NA
                                                    NA
                                                             NA
                                                                      NA
                                                                               NA
# 4
       0
            0 1.2724293 -0.2947204 NA NA
                                                    NA
                                                             NA
                                                                      NA
                                                                               NA
```

zonogon-section 45

zonogon-section	compute the intersection of a line and the boundary of a zonogon
G	

Description

Generically, a line intersects the boundary of a zonogon in 2 points. Computing those 2 points is the chief goal of this function.

For a supporting line, the intersection is a face of the zonogon, but in this function only one point of intersection is computed and returned.

Usage

```
## S3 method for class 'zonogon'
section( x, normal, beta, tol=1.e-10, plot=FALSE, ... )
```

Arguments

X	a zonogon object as returned by the constructor zonogon()
normal	a non-zero numeric 2-vector - the normal of all the lines
beta	a numeric M-vector of line-constants. The equation of the k'th line k is: $<$ x, normal> = beta[k].
tol	a small positive number, used as the tolerance for the line being considered a supporting line
plot	if TRUE, the line segments formed by the intersection of the lines and the zonogon are <i>added</i> to an existing plot of the zonogon x, see plot.zonogon(). The segments are drawn in dashed linestyle and the color red. boundary1 and boundary2 are plotted as points. If there is no open plot, a warning is issued.
	not used

Value

section.zonogon() returns a data.frame with M rows and these columns:

normal the given normal vector - this is the same in every row

beta the given line constant

boundary1 the 1st intersection point - a 2-vector boundary2 the 2nd intersection point - a 2-vector

Regarding orientation, if normal is considered "north" then boundary1 is on the "west" and boundary2 is on the "east".

If a line is a supporting line of the zonogon, then boundary1 is some point in the boundary face (vertex or edge), and boundary2 is NA. If a line does not intersect the zonogon, both boundary1 and boundary2 are NA.

If the names of beta are unique, they are copied to the row names of the output.

In case of error, the function returns NULL.

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See Also

```
zonogon(), plot.zonogon(), section.zonohedron()
```

Examples

```
# make a zonogon with 5 generators
pz20 = polarzonogon( 20, 5 )
section( pz20, normal=c(1,1), beta=-1:5 )
# normal.1 normal.2 beta boundary1.1 boundary1.2 boundary2.1 boundary2.2
# 1
         1 1 -1 NA NA NA NA
         1 1 0 -2.220446e-16 0.000000e+00 NA NA
1 1 1 2.452373e-01 7.547627e-01 1.0000000 0.0000000
1 1 2 6.203838e-01 1.379616e+00 1.7547627 0.2452373
1 1 3 1.095537e+00 1.904463e+00 2.3796162 0.6203838
# 2
# 3
# 4
# 5
# 6
                  1 4 1.674729e+00 2.325271e+00 2.9044629 1.0955371
          1
          1 1
# 7
                          5 2.420068e+00 2.579932e+00 3.3252706 1.6747294
```

zonohedron

zonohedron construction

Description

Construct a zonohedron from a numeric matrix with 3 rows. Also construct some special zonohedra useful for testing.

Usage

```
zonohedron( mat, e0=0, e1=1.e-6, e2=1.e-10, ground=NULL )
polarzonohedron( n, m=n, height=pi, ground=NULL )
regularprism( n, m=n, axis=c(0,0,1), ground=NULL )
```

mat	a numeric $3xM$ matrix, where $3 \le M$. The matrix must have rank 3 (verified). The M columns are the generators of the zonohedron.
e0	threshold for a column of mat to be considered 0, in the L^∞ norm. Since the default is e0=0, by default a column must be exactly 0 to be considered 0.
e1	threshold, in a pseudo-angular sense, for non-zero column vectors to be multiples of each other, and thus members of a group of multiple (aka parallel) points in the associated matroid. It OK for a column to be a negative multiple of another.
e2	threshold, in a pseudo-angular sense, for the planes spanned by pairs of column vectors to be considered coincident, and thus the columns to be in the same hyperplane of the associated matroid.

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ground	The <i>ground set</i> of the associated matroid of rank 3 - an integer vector in strictly increasing order, or NULL.
	When ground is NULL, it is set to 1:ncol(mat). If ground is not NULL, length(ground) must be equal to ncol(mat). The point ground[i] corresponds to the <i>i'th</i> column of mat.
n	an integer ≥ 3 . The generators are computed as n equally spaced points on a circle. See Details for more on this circle.
m	an integer with $2 \le m \le n$. When $m < n$, only the first m points are used as generators of the zonohedron.
height	the z value at the apex of the zonohedron, which is the sum of all the generators. The z value of all the generators is set to make this happen. When height=pi, as $n \to \infty$ the zonohedron converges to the interior of the surface of revolution of the curve $x = sin(z)$ for $z \in [0, \pi]$, see Chilton & Coxeter .
axis	the axis of the regular prism. It must be a 3-vector with z value non-zero.

Details

In zonohedron(), the contruction of the *zones* (or *belts*) is optimized by following the procedure in *Heckbert*. The key step is sorting face normals that all lie on a great circle of the unit sphere.

For polarzonohedron() the circle is centered at (0,0,height/n) and parallel to the xy-plane. The radius is height/n.

For regularprism() the circle is the unit circle in the xy-plane. The 3-vector axis is added as column m+1 of the matrix. The returned zonohedron is the Minkowski sum of a zonogon and the line segment defined by axis. If m < n, the zonogon may not be regular.

Both of these functions are useful for testing. They load the matrix mat and pass it to zonohedron().

Value

zonohedron() and polarzonohedron() return a list with S3 class 'zonohedron'. In case of error, e.g. invalid mat, the functions print an error message and returns NULL.

Note

The ground set of positive integers should not be too sparse; otherwise performance may suffer.

References

B. L. Chilton and H. S. M. Coxeter. **Polar Zonohedra**. The American Mathematical Monthly. Vol 70. No. 9. pp. 946-951. 1963.

Paul Heckbert. **An Efficient Algorithm for Generating Zonohedra**. 3-D Technical Memo 11. 24 February 1985. Computer Graphics Lab. New York Institute of Technology

See Also

zonohedron(), zonoseg(),

48 zonohedron-plot

Description

Plot a **zonohedron** object in 3D, with many options.

Usage

x	a zonohedron object as returned by the constructor zonohedron()
type	a string of letter with what parts to draw. If type contains an 'p', then draw a point at the center of each facet. If type contains an 'e', then draw the edges. If type contains an 'f', then draw filled facets.
pcol	The color to use when drawing points. It can be a vector of 2 colors, and then when both is TRUE, the first color is used for one half, and the second color is used for the antipodal half. When pcol is NULL, it is set to c('black', 'red').
ecol	A vector of colors to use when drawing the edges. Let N be the number of <i>simplified</i> generators of the zonohedron. Each edge is parallel to exactly one of the generators, so this divides the edges into N <i>zones</i> , or <i>belts</i> . ecol can be a vector of N colors, one for each zone. If ecol is shorter than N, it is extended to length N using the last color. If ecol is longer than N, the extra colors are ignored. If ecol is NULL, it is set to rainbow(N).
ewd	width of the edges, in pixels
etcol	color of the <i>tiling edges</i> , for the standard tiling of the facets by parallelograms. This only applies to facets that are <i>not</i> parallelograms. The default etol=NA means do not draw these edges.
fcol	A vector of colors to use when drawing the facets. The 1st color is used for parallelograms, the next color for hexagons, etc. For facets with more edges than colors available, the last color is used. If fcol is NULL, it is set to c('blue', 'red', 'yellow', 'green', 'orange', 'purple').
falpha	opacity of the facets
normals	if TRUE then draw an outward-pointing unit normal from each facet
bgcol	the background color
both	if FALSE then draw only one half of the centrally symmetric boundary. Otherwise draw both halves. This affects points, edges, and facets.
	not used

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Details

Points are drawn with rgl::points3d(). Edges are drawn with rgl::segments3d(). Edges of the tiles are drawn with rgl::quads3d(). Facets are drawn with rgl::quads3d(); facets with more than 4 edges are split into trapezoids. Facet normals are drawn with rgl::arrow3d().

Value

The function returns TRUE; or FALSE in case of error.

Note

The package **rgl** is required for 3D plots. A large black point is drawn at 0, a 50% gray point at the center, and a large white point at the "white point" (which is 2*center).

A line from the black point to the white point is also drawn.

See Also

```
zonohedron(), spherize.zonotope()
```

zonohedron-raytrace

compute the intersection of a ray, based in the interior of a zonohedron, and the boundary of that zonohedron

Description

The open ray with basepoint b and non-zero direction d is the set of the form b + td where t > 0.

This function computes the intersection of an open ray and the boundary of a zonohedron Z. The basepoint is normally required to be in the interior of Z, but an exception is made if the basepoint is 0, and on the boundary of Z, and the direction points into the interior of Z. In these two cases the intersection of the open ray and the boundary of Z is unique. In the second case, the basepoint is also allowed to be the sum of all the generators - the so-called *white point* of Z.

Usage

```
## S3 method for class 'zonohedron'
raytrace( x, base, direction, invert=FALSE, plot=FALSE, ... )
```

Arguments

x a **zonohedron** object as returned by the constructor zonohedron()

base a numeric 3-vector - the basepoint of all the rays. base must either be in the

interior of x, or 0 or the white point and on the boundary of x.

direction a numeric Mx3 matrix with M non-zero directions in the rows. The basepoint

and these directions define M rays.

direction can also be a numeric vector that can be converted to such a matrix,

by row.

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invert if TRUE, then compute a point in the unit cube that maps to the point on the

boundary of x and add it as a column in the returned data. frame

plot if TRUE, the computed rays, up to the boundary, are *added* to an existing plot

of the zonohedron x, see plot.zonohedron(). The segments are drawn in the

color red. If there is no open plot, a warning is issued.

... not used

Details

If plot is TRUE, the rays are drawn with rgl::segments3d().

Value

raytrace.zonohedron() returns a data.frame with M rows and these columns:

base the given basepoint - this is the same in every row

direction the given direction

facetidx the index of the facet (a zonogon) where ray exits the zonohedron

sign of the facet, either +1 or -1

ray parameter of the intersection with the exit facet, always positive

point the point on the boundary; the intersection of the ray and the facet

timetrace the computation time, in seconds

And if invert is TRUE, then these columns are added:

distance signed distance to the boundary of x

pcube a point in the unit cube that maps to boundary

transitions the number of transitions in pcube - a non-negative even integer

If base and direction in a row cannot be processed, the rest of the row is NA.

If the row names of direction are unique, they are copied to the row names of the output.

In case of error, the function returns NULL.

Note

The package **rgl** is required for 3D plotting.

See Also

zonohedron(), plot.zonohedron(), section.zonohedron(), invertboundary(), raytrace.zonogon()

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Examples

```
make a regular prism, a regular 20-gon extruded 1 unit along z-axis
rp10 = regularprism( 10 )
    make 7 random directions
set.seed(0)
dir = matrix(rnorm(7*3),7,3)
# use basepoint in the interior of the zonohedron
raytrace( rp10, c(0.5,0.5,0.5), dir )
   base.1 base.2 base.3 direction.1 direction.2 direction.3 facetidx sign
# 1
       0.5
             0.5 0.5 1.262954285 -0.294720447 -0.299215118 1 1 1.6710386
# 2
       0.5
              1 1.2150348

      0.5
      0.5
      0.5
      1.329799263
      2.404653389
      0.252223448
      6
      -1 0.8724774

      0.5
      0.5
      0.5
      1.272429321
      0.763593461
      -0.891921127
      1
      1 0.5605877

      0.5
      0.5
      0.414641434
      -0.799009249
      0.435683299
      1
      -1 1.1476226

# 3
# 4
# 5

      0.5 -1.539950042 -1.147657009 -1.237538422
      1

      0.5 -0.928567035 -0.289461574 -0.224267885
      1

# 6
       0.5
              0.5
                                                                                     1 0.4040279
# 7
                                                                                     1 2.2294766
\# use basepoint 0 on the boundary of the zonohedron
  note that only 2 directions point into the interior
raytrace( rp10, c(0,0,0), dir )
   base.1 base.2 base.3 direction.1 direction.2 direction.3 facetidx sign
                        0 1.262954285 -0.294720447 -0.299215118 NA NA
# 2
                        0 -0.326233361 -0.005767173 -0.411510833
                                                                                NA NA
                                                                                                NA
# 3
                       0 1.329799263 2.404653389 0.252223448
                                                                               6 -1 1.128580
                                                                               NA NA
                        0 1.272429321 0.763593461 -0.891921127
# 4
         0
             a
                                                                                                NA
             0
                       0 0.414641434 -0.799009249 0.435683299
# 5
         0
                                                                               1 -1 2.295245
             0 0 -1.539950042 -1.147657009 -1.237538422
0 0 -0.928567035 -0.289461574 -0.224267885
# 6
         0
                                                                               NA NA NA
# 7
                                                                                NA NA
                                                                                                NA
```

zonohedron-section

compute the intersection of a plane and the boundary of a zonohedron

Description

Generically, a plane intersects the boundary of a zonohedron in a convex polygon. Computing that polygon is the chief goal of this function.

For a supporting plane, the intersection is a face of the zonohedron, but in this function only one point of intersection is computed and returned.

Usage

```
## S3 method for class 'zonohedron'
section( x, normal, beta, tol=1.e-10, plot=FALSE, ... )
```

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Arguments

x	a zonohedron object as returned by the constructor zonohedron()	
normal	a non-zero numeric 3-vector - the normal of all the planes	
beta	a numeric M-vector of line-constants. The equation of the k'th plane k is: $< x, normal > = beta[k]$.	
tol	a small positive number, used as the tolerance for the plane being considered a supporting plane	
plot	if TRUE, the polygons formed by the the intersection of the planes and the boundary of the zonohedron are <i>added</i> to an existing 3D plot of the zonohedron x, see plot.zonohedron(). The polygons are drawn in red.	
	not used	

Details

Given a plane, the function finds all the facets of the zonohedron that intersect the plane. For each such facet it computes a single point of intersection on the boundary of the facet. For the parallelograms, the computation is done in a C function; and for zonogon facets with 3 or more generators, the computation is done in section.zonogon(). Orientation is handled carefully so that no point appears twice. The facets are not processed in order around the boundary, so these points are in no particular order. They are put in polygon order by sorting them by angle around a suitable "diameter" of the zonohedron.

Value

section.zonohedron() returns a list of length M (=length(beta)), and the i'th item in the list is a data frame with these columns:

a data frame with these columns.				
point	a Px3 matrix with the P points of the i'th polygon in the rows. If the plane does			

not intersect the zonohedron, then P=0 and the matrix has 0 rows. If the plane is a supporting plane, the polygon is degenerate and P=1 and the matrix has 1 row. The row names of section are the indexes of the facets that contain the vertices

of the polygon; see Details.

hyperidx index of a hyperplane that contains the given point

sign The sign specifying which of the 2 facets (selected or antipodal) contains the

given point. The value is +1 or -1.

The names of the list are readable strings that contain normal and beta[i].

In case of error, the function returns NULL.

Note

The package **rgl** is required for 3D plotting.

See Also

```
zonohedron(), plot.zonohedron(), section.zonogon()
```

zonoseg 53

Description

Construct a zonoseg from a numeric matrix with one row.

A zonoseg ("zonotope" + "segment") is my own personal term for a 1-dimensional zonotope. I could not find an alternative term. It is a linear image of the unit cube $[0,1]^n$ in the real numbers, and a compact segment of reals. The order of the generators has no effect on the zonoseg.

The image of the 2-transition subcomplex of $[0,1]^n$ is a compact subsegment of the zonoseg. The order of the generators affects this subsegment in a major way.

Usage

```
zonoseg( mat, e0=0, ground=NULL )
## S3 method for class 'zonoseg'
getsegment( x )
## S3 method for class 'zonoseg'
getsegment2trans( x )
## S3 method for class 'zonoseg'
print( x, ... )
```

Arguments

mat	a numeric matrix with 1 row whose entries determine the zonoseg. One or more entries must be non-zero. It is OK to have both positive and negative entries. mat can also be a numeric vector which is then converted to a matrix with 1 row.
e0	threshold for an entry of mat to be considered 0. Since the default is e0=0, by default an entry must be exactly 0 to become a loop in the associated matroid.
ground	The <i>ground set</i> of the associated matroid of rank 1 - an integer vector in strictly increasing order, or NULL. When ground is NULL, it is set to 1:ncol(mat). If ground is not NULL, length(ground) must be equal to ncol(mat). The point ground[i] corresponds to the <i>i'th</i> column of mat.
Χ	a zonoseg object as returned by zonoseg()
	not used

Details

A **zonoseg** object is a list with only 3 items: the associated matroid, the endpoints of the segment, and endpoints of the 2-transition subsegment.

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print.zonoseg() prints some information about the generators, and the endpoints of the segment plus the 2 vertices of the unit cube that map to these endpoints. It prints similar data for the 2-transition subsegment. Finally, it prints data on the associated matroid.

Value

zonoseg() returns a list with S3 class 'zonoseg'. In case of error, e.g. invalid mat, the function prints an error message and returns NULL.

getsegment() and getsegment2trans() return numeric 2-vectors - the min and max endpoints of the corresponding segments.

```
print.zonoseg() returns TRUE or FALSE.
```

Note

The ground set of positive integers should not be too sparse; otherwise performance may suffer.

References

Matroid - Wikipedia.

```
https://en.wikipedia.org/w/index.php?title=Matroid&oldid=1086234057
```

See Also

```
rank()
```

```
zono1 = zonoseg(c(1,-2,3,0,-3,-4))
zono1
                 6 -- 3 negative, 2 positive, and 1 loops.
# generators:
#
# segment:
                          [-9,4]
      value pcube.1 pcube.2 pcube.3 pcube.4 pcube.5 pcube.6
#
# zmin
       -9
            0
                     1
                             0
                                       0
                                             1
# zmax
                 1
                        0
# 2-transition subsegment: [-8,3]
            value source.1 source.2 source.3 source.4 source.5 source.6
                     1
# tmin-2trans
             -8
                                      0
                                                 0
                               1
                                                        1
                                                                  1
                                 0
                                                                  0
# tmax-2trans
                3
                        0
                                         1
                                                 1
# matroid:
                     6 points
                               {1 2 3 4 5 6}
# ground set:
# hyperplanes:
                     1
                           {4}
# rank:
                     1
# loops:
                     1
                         {4}
                           {1 2 3 5 6}
# multiple groups: 1
```

zonoseg-invert 55

```
# uniform:
                       FALSE
# paving:
                       TRUE
# simple:
                       FALSE
# This matroid is constructed from a 1x6 real matrix.
      1 2 3 4 5 6
# [1,] 1 -2 3 0 -3 -4
# The summary of the simplified matroid is:
     ground set:
                           1 points {1}
                 Point 1 corresponds to the multiple group {1 2 3 5 6} in the original ...
#
      hyperplanes:
                           1
                                  {}
#
      rank:
                            1
      loops:
                            0 {}
      multiple groups:
                            0 {}
      uniform:
                            TRUE
      paving:
                            TRUE
      simple:
                            TRUE
#
      This matroid is constructed from a 1x1 real matrix.
#
           1+...+6
      [1,]
               -13
```

so the 2-transition subsegment is a proper subset of the zonoseg

zonoseg-invert

invert points in a zonoseg

Description

For points in a zonoseg, find points in the unit cube that map to those points.

Usage

```
## S3 method for class 'zonoseg'
invert( x, z, tol=0, ... )
```

Arguments

```
x a zonoseg object as returned by the constructor zonoseg()
z a numeric M-vector
tol points that are within tol of a boundary point are taken to be that point
... not used
```

Details

For a point in the interior of the zonoseg, there are infinitely many points in the cube that map to it. This function tries to find one with the fewest number of non-zero components.

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Value

invert.zonoseg() returns a data.frame with M rows and these columns:

z the given point

pcube a point in the unit cube that maps to z. For the 2 boundary points, pcube is a

vertex. If z is outside the zonoseg, pcube is all NA, and a warning message is

printed

If the names of z are unique, they are copied to the row names of the output. The column names are copied from the ground set of the associated matroid.

In case of error, the function returns NULL.

See Also

```
zonoseg()
```

```
zono1 = zonoseg( c(1,-2,3,0,-3,-4) )
zono1
# generators:
                  6 -- 3 negative, 2 positive, and 1 loops.
                         [-9,4]
      value pcube.1 pcube.2 pcube.3 pcube.4 pcube.5 pcube.6
       -9
              0 1
                              0
                                      0
# zmin
                                            1
                1
                       0
                                      0
                                             0
                                                    0
# zmax
                              1
# 2-transition subsegment: [-8,3]
            value source.1 source.2 source.3 source.4 source.5 source.6
# tmin-2trans
                                       0
                                                0
              -8
                       1
                               1
                                                                1
                                                1
                                                        0
                                                                0
# tmax-2trans
               3
                       0
                                0
                                        1
z = c(0, -3*pi, pi, 2*pi, getsegment(zono1))
invert( zono1, z )
#
          z pcube.1
                       pcube.2
                                pcube.3
                                        pcube.4
                                                 pcube.5
# 2 -9.424778
                                    NA
                  NA
                           NA
                                             NA
                                                      NA
# 3 3.141593 1.0000000 0.0000000 0.7138642 0.0000000 0.0000000 0.0000000
# 4 6.283185
                           NA
                                    NA
                                             NA
# 5 -9.000000 0.0000000 1.0000000 0.0000000 0.0000000 1.0000000 1.0000000
# 6 4.00000 1.0000000 0.0000000 1.0000000 0.0000000 0.0000000 0.0000000
```

zonotope-getters 57

zonotope-getters

zonotope get functions

Description

get some important members of a zonotope

Usage

```
## S3 method for class 'zonotope'
getmatrix( x )

## S3 method for class 'zonotope'
getmatroid( x )

## S3 method for class 'zonotope'
getcenter( x )
```

Arguments

Х

a zonotope object - a zonohedron, a zonogon, or a zonoseg

Value

```
getmatrix() returns the matrix originally used to construct the zonotope x.
```

getmatroid() returns the matroid (possibly nonsimple) constructed from the matrix

getcenter() returns the center of the zonotope; which is also the center of radial symmetry. If x is an object-color solid, the center corresponds to the 50% graypoint. For the whitepoint multiply by 2.

See Also

```
zonohedron(), zonogon(), zonoseg()
```

zonotope-props

zonotope properties

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Description

Get some important boolean properties of a zonotope.

pointed means that 0 is a vertex of the zonotope. *salient* means that 0 is in the boundary of the zonotope. So pointed implies salient, but not the reverse.

A zonotope has an associated *convex cone* - allow the coefficients of the generators to be any non-negative numbers. For convex cones, *pointed* means that the cone is in an open linear halfspace (except for 0). And *salient* means that the cone is in a closed linear halfspace (the cone may contain a line).

In terms of generators (of both zonotopes and convex cones), *pointed* means that the generators are in an open linear halfspace (except for 0 generators). And *salient* means that the generators are in a closed linear halfspace.

Usage

```
## S3 method for class 'zonotope'
is_pointed( x )
## S3 method for class 'zonotope'
is_salient( x )
```

Arguments

Х

a zonotope object - a zonohedron, a zonogon, or a zonoseg

Details

For a **zonohedron**, if 0 is in the interior of an edge or a facet, then the zonohdron is salient but not pointed.

For a **zonogon**, if 0 is in the interior of an edge, then the zonogon is salient but not pointed. For a **zonoseg**, both *pointed* and *salient* are equivalent to 0 being a boundary point. And this is equivalent to all the non-zero generators having the same sign (all negative or all positive).

Value

TRUE or FALSE

References

```
Zonohedron - Wikipedia.
```

```
https://en.wikipedia.org/wiki/Zonohedron
```

See Also

```
zonohedron(), zonogon(), zonoseg()
```

zonotope-props 59

```
zono1 = zonoseg( c(1,-2,3,0,-3,-4) )
is_pointed( zono1 )
# [1] FALSE
is_salient( zono1 )
# [1] FALSE
```

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