# Package 'Rpdb'

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**License** GPL ( $\geq 3$ )

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Rpdb-package

Read, Write, Visualize and Manipulate PDB Files

## Description

Provides tools to read, write, visualize PDB files, and perform structural manipulations.

# Details

This package enables users, e.g. computational chemists, to manipulate molecular structures stored in PDB files. It enables users ro read, write and visualize PDB files. Various basic structural manipulations are also supported. Conversion of Cartesian coordinates into fractional coordinates. Spliting a molecular structure into fragments. Computation of centers-of-geometry and centers-ofmass. Wrapping molecular structure using periodical boundary conditions. Translation, rotation and reflection of atomic coordinates. Calculate atomic bond lengths, angles and dihedrals.

# Rpdb-package

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# References

More information on the PDB format can be found here: http://www.wwpdb.org/documentation/format33/v3.3.html

#### Examples

```
## Read a PDB file included in the package
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))</pre>
## Visualize the PDB file
visualize(x, mode = NULL)
## From Cartesian to fractional coordinates and vice versa
x \leftarrow xyz2abc(x)
basis(x)
natom(x, x$atoms$resid)
range(x)
centres(x)
x \le abc2xyz(x)
basis(x)
natom(x, x$atoms$resid)
range(x)
centres(x)
## Split and unsplit
F <- x$atoms$resid</pre>
x <- split(x, F)
x <- unsplit(x, F)</pre>
## Subset and merge
x.PCB.only <- subset(x, resname == "PCB")</pre>
x.DCB.only <- subset(x, resname == "DCB")</pre>
x <- merge(x.PCB.only, x.DCB.only)</pre>
## Duplicate and wrap
x <- replicate(x, a.ind = -1:1, b.ind = -1:1, c.ind = -1:1)</pre>
x <- wrap(x)
## Write the 'pdb' object 'x' to a temporary file.
write.pdb(x, file = tempfile())
```

addAxes

# Description

Add lattice vectors, Cartesian axes or PBC box to the current 'rgl' scene.

# Usage

```
addABC(x, lwd = 2, labels = TRUE, cex = 2)
addXYZ(lwd = 2, labels = TRUE, cex = 2)
addPBCBox(x, lwd = 2)
```

#### Arguments

х	an object of class 'cryst1' containing unit cell parameters.
lwd	a numeric value indicating the line width used to draw the axes or the PBC box.
labels	a logical value indicating whether the labels of the axes have to be drawn.
cex	a numeric value indicating the magnification used to draw the labels of the axes.

# Details

addABC: Add the lattice vectors a, b and c to the current rgl device. addXYZ: Add the Cartesian axes x, y and z to the current rgl device. addPBCBox: Add a box representing the Periodic Boundary Conditions of a molecular system.

## Value

Return (using invisible) a two-column data.frame containing the IDs and type indicators of the objects added to the scene.

#### See Also

visualize, rgl.open, par3d, addLabels

# Examples

```
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
visualize(x, type = "l", xyz = FALSE, abc = FALSE, pbc.box = FALSE, mode = NULL)
addXYZ()
addABC(x$cryst1)
addPBCBox(x$cryst1)</pre>
```

addLabels

# Description

Add Labels to the current 'rgl' scene.

#### Usage

```
addResLab(x, ...)
## S3 method for class 'atoms'
addResLab(x, at.centre = TRUE, col = "black", ...)
## S3 method for class 'pdb'
addResLab(x, at.centre = TRUE, col = "black", ...)
addEleLab(x, ...)
## S3 method for class 'atoms'
addEleLab(x, eleid = FALSE, col = "black", ...)
## S3 method for class 'pdb'
addEleLab(x, eleid = FALSE, col = "black", ...)
info3d(...)
## S3 method for class 'atoms'
info3d(x, id = rgl::rgl.ids(), col = "black", verbose = TRUE, adj = 0, ...)
## S3 method for class 'pdb'
```

# Arguments

х	an R object containing atomic coordinates.
	further arguments passed to or from other methods.
at.centre	a single element logical vector indicating if residue labels have to be added only at the position of the residue's centre-of-mass instead of at each atomic position.
col	the colors used to display the labels.
eleid	a single element logical vector indicating if the element ids have to be concate- nated with the element names to prepare the labels.
id	vector of ID numbers of 'rgl' items, as returned by rgl.ids. The vertexes of these items are used to display the labels.
verbose	a logical value specifying if information have to be printed to the terminal.

atoms

adj one value specifying the horizontal adjustment, or two, specifying horizontal and vertical adjustment respectively. See text3d

#### Details

addResLab add residue labels to the scene. If at.centre==TRUE only one label per residue is added at the centre of the residue. Otherwise, residue labels are added at each atomic positions. addEleLab add element labels to the scene at each atomic positions. info3d activate an interactive mode to add labels by selecting atoms by **right-clicking** on the current 'rgl' scene. To escape the interactive mode press the ESC key. The labels are as follow: "ResidResname:EleidElename"

## Value

addResLab and addEleLab return (using invisible) a two-column data.frame containing the IDs and type indicators of the objects added to the scene.

## See Also

pdb, visualize, measure

# Examples

```
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
visualize(x, type = "1", mode = NULL)
addResLab(x)
x <- read.pdb(system.file("examples/Pentacene.pdb", package="Rpdb"))
visualize(x, type = "1", mode = NULL)
addEleLab(x)</pre>
```

atoms

Create 'atoms' Object

# Description

Creates an object of class 'atoms' containing the data related to ATOM and HETATM records of a PDB file.

#### Usage

```
atoms(...)
## Default S3 method:
atoms(
   recname,
   eleid,
   elename,
   alt,
```

atoms

```
resname,
chainid,
resid,
insert,
x1,
x2,
x3,
occ,
temp,
segid,
basis = "xyz",
...
```

is.atoms(x)

# Arguments

)

	arguments passed to methods.
recname	a character vector containing the record name for each element.
eleid	a integer vector containing the element ID for each element.
elename	a character vector containing the element name for each element.
alt	a character vector containing the alternate location indicator for each element.
resname	a character vector containing the residue name for each element.
chainid	a character vector containing the chain ID for each element.
resid	a integer vector containing the residue ID for each element.
insert	a character vector containing the codes for insertion of residue of each element.
x1, x2, x3	a numeric vector containing the first, second and third coordinate for each ele- ment.
осс	a numeric vector containing the occupancie for each element.
temp	a numeric vector containing the temperature factor for each element.
segid	a character vector containing the segment ID for each element.
basis	a single element character vector indicating the type of basis vector used to express the atomic coordinates.
x	an R obecjt to be tested.

# Details

atoms is a generic function to create objects of class 'atoms'. The purpose of this class is to store ATOM and HETATM records from PDB files. The default method creates a atoms object from its different components, i.e.: recname, eleid, elename, alt, resname, chainid, resid, insert, x1, x2, x3, occ, temp, segid and basis. All the arguments have to be specified except basis which by default is set to "xyz" (Cartesian coordinates).

is. atoms tests if an object of class 'atoms', i.e. if it has a "class" attribute equal to atoms.

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atoms returns a data.frame of class 'atoms' with the following components:

**recname** a character vector containing the record name for each element.

eleid a integer vector containing the element ID for each element.

elename a character vector containing the element name for each element.

alt a character vector containing the alternate location indicator for each element.

resname a character vector containing the residue name for each element.

chainid a character vector containing the chain ID for each element.

resid a integer vector containing the residue ID for each element.

insert a character vector containing the codes for insertion of residue for each element.

x1, x2, x3 a numeric vector containing the first, second and third coordinate for each element.

occ a numeric vector containing the occupencie for each element.

temp a numeric vector containing the temperature factor for each element.

segid a character vector containing the segment ID for each element.

**basis** a single element character vector indicating the type of basis vector used to express the atomic coordinates.

is.atoms returns TRUE if x is an object of class 'atoms' and FALSE otherwise.

# See Also

basis, coords, pdb

# Examples

```
x <- atoms(recname = c("ATOM","ATOM"), eleid = 1:2, elename = c("H","H"), alt = "",
resname = c("H2","H2"), chainid = "", resid = c(1,1), insert = "",
x1 = c(0,0), x2 = c(0,0), x3 = c(0,1), occ = c(0.0,0.0), temp = c(1.0,1.0),
segid = c("H2","H2"))
print(x)
is.atoms(x)
```

basis

The Basis of an Object

## Description

Functions to get or set the basis of an object containing atomic coordinates.

basis

# Usage

```
basis(x)
## Default S3 method:
basis(x)
basis(x) <- value
## Default S3 replacement method:
basis(x) <- value
## S3 method for class 'pdb'
basis(x)
## S3 replacement method for class 'pdb'
basis(x) <- value</pre>
```

# Arguments

х	an R object containing atomic coordinates.
value	a single element character vector use to set the basis of x.

# Details

basis and basis<- are respectively generic accessor and replacement functions. The default methods get and set the basis attribute of an object containing atomic coordinates. This attribute indicate the type basis vector used to express atomic coordinates.

value must be equal to "xyz", for Cartesian, or "abc", for fractional coordinates. The methods for objects of class 'pdb' get and set the basis attribute of its atoms component.

## Value

For basis: NULL or a single element character vector. (NULL is given if the object has no basis attribute.)

For basis<-: the updated object. (Note that the value of basis(x) <- value is that of the assignment, value, not the return value from the left-hand side.)</pre>

# See Also

coords, atoms, pdb

#### Examples

```
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
basis(x)
x <- xyz2abc(x)
basis(x)</pre>
```

bond-angle-dihedral Atomic Bond Lengths, Angles and Dihedrals

#### Description

Compute bond lengths, angles and dihedral from atomic coordinates.

# Usage

```
bond(...)
## S3 method for class 'coords'
bond(x, sel1, sel2, ...)
## S3 method for class 'pdb'
bond(x, sel1, sel2, ...)
angle(...)
## S3 method for class 'coords'
angle(x, sel1, sel2, sel3, ...)
## S3 method for class 'pdb'
angle(x, sel1, sel2, sel3, ...)
dihedral(...)
## S3 method for class 'coords'
dihedral(x, sel1, sel2, sel3, sel4, ...)
## S3 method for class 'pdb'
dihedral(x, sel1, sel2, sel3, sel4, ...)
measure(...)
## Default S3 method:
measure(id = rgl::rgl.ids(), verbose = TRUE, ...)
## S3 method for class 'coords'
measure(x, id = rgl::rgl.ids(), verbose = TRUE, ...)
## S3 method for class 'pdb'
measure(x, id = rgl::rgl.ids(), verbose = TRUE, ...)
```

# Arguments

... further arguments passed to or from other methods.

Х	an R object containing atomic coordinates.
sel1, sel2, sel3,	sel4
	an integer or logical vector used to select atoms defining bonds, angles or dihe- drals. See details.
id	vector of ID numbers of 'rgl' items, as returned by rgl.ids. The vertexes of these items are used to compute the bond lengths, angles or dihedrals.
verbose	a logical value specifying if the information have to be printed to the terminal.

# Details

The number of selected atoms with sel1, sel2, sel3 and sel4 must be the same. sel1, sel2, sel3 and sel4 respectively select the first, second, third and fouth atoms defining bonds, angles or dihedrals.

measure activate an interactive mode to compute bond lengths, angles and dihedrals by selecting atoms by **right-clicing** on the current 'rgl' scene. To escape the active mode press the ESC key.

# Value

A numeric vector containing atomic bond lengths (in Angstrom), angles or dihedrals (in degrees)

# See Also

coords, pdb, info3d, visualize

# Examples

```
Pen <- read.pdb(system.file("examples/Pentacene.pdb", package="Rpdb"))
visualize(Pen, mode = NULL)
text3d(coords(Pen), texts = Pen$atoms$eleid)
bond(Pen, 3:4, 1:2)
angle(Pen, 3:4, 1:2, 5:6)
dihedral(Pen, 3:4, 1:2, 5:6, 6:5)</pre>
```

cellProperties Properties of a Unit Cell

# Description

Compute the Cartesian coordinates of lattice vectors, the volume or the density of a unit cell.

#### cellProperties

# Usage

```
cell.coords(...)
## Default S3 method:
cell.coords(abc, abg = c(90, 90, 90), digits = 3, ...)
## S3 method for class 'cryst1'
cell.coords(x, digits = 3, ...)
## S3 method for class 'pdb'
cell.coords(x, digits = 3, ...)
cell.volume(...)
## S3 method for class 'cryst1'
cell.volume(x, ...)
## S3 method for class 'pdb'
cell.volume(x, ...)
cell.density(...)
## Default S3 method:
cell.density(masses, volume, ...)
## S3 method for class 'pdb'
cell.density(x, ...)
```

#### Arguments

	further arguments passed to or from other methods.
abc	a length 3 numeric vector containing the length of the a, b and c lattice vectors.
abg	a length 3 numeric vector containing the angles (degrees) between the a, b and c lattice vectors (alpha, beta, gamma).
digits	an integer used to round the lattice vectors coordinates.
х	an R object containing lattice parameters.
masses	a numeric vector containing atomic masses.
volume	a single element numeric vector containing the volume of the unit cell in Angstrom cube.

# Details

cell.coords is a generic function which computes a 3x3 matrix whose columns contrain the Cartesian coordinates of lattice vectors. The 'a' and 'b' vectors are assumed to be respectively along the x-axis and in the xy-plane. The default method takes directly the lattice parameters as arguments. For objects of class cryst1 the lattice parameters are first extracted from the object and then the default method is called. For objects of class pdb the lattice parameters are extracted from their

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#### centres

cryst1 component and the default method is called.

cell.volume is a generic function to compute the volume of a unit cell. For objects of class 'cryst1', the unit cell parameters are directly used to compute the volume. For objects of class 'pdb', their cryst1 component is used.

cell.density is a generic function to compute the density of a unit cell. For objects of class 'pdb': First the volume of the unit cell is calculated by calling the cell.volume function on the cryst1 component of the 'pdb' object. Then the element names are converted into element symbols using the toSymbols function and their masses are taken from the elements data set. Finally the density is calculated using the sum of the atomic masses and the volume of the unit cell.

# Value

cell.coords returns a 3x3 matrix containing the Cartesian coordinates of lattice vectors arranged by columns.

cell.volume returns a single element numeric vector containing the volume of the unit cell in Angstrom cube.

cell.density returns a single element numeric vector containing the density of the unit cell in g.cm-3.

# See Also

cryst1, pdb, xyz2abc

# Examples

```
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
cell.volume(x)
cell.density(x)
cell.coords(x)</pre>
```

centres

Centres-of-Geometry and Centres-of-Mass

#### Description

Computes centres-of-geometry and centres-of-mass of groups of atoms.

# Usage

```
centres(...)
## S3 method for class 'coords'
centres(x, factor = NULL, weights = NULL, unsplit = FALSE, na.rm = FALSE, ...)
## S3 method for class 'atoms'
```

```
centres(x, factor = NULL, weights = NULL, unsplit = FALSE, na.rm = FALSE, ...)
```

```
## S3 method for class 'pdb'
centres(x, factor = NULL, weights = NULL, unsplit = FALSE, na.rm = FALSE, ...)
```

#### Arguments

	further arguments passed to or from other methods.
x	an R object containing atomic coordinates.
factor	a factor used to split the atomic coordinates by groups to compute multiple cen- tres.
weights	a numerical vector containing atomic weights used to compute centres-of-mass.
unsplit	a logical value indicating whether the coordinates of the centres have to be un- split to repeat their coordinates for each atom used for their calculation (used for wrapping by groups).
na.rm	a logical value indicating whether NA values should be stripped before the com- putation proceeds.

# Details

centres is a generic function to compute centres-of-geometry and centres-of-mass from an object containing atomic coordinates. For objects of class 'coords', 'atoms' and 'pdb', the coordinates of x are first splitted into groups defined by factor using the split function. For each group, the weighted mean of the x1, x2 and x3 components of x are calculated using weights. By default all atoms are assumed to have the same weight (calculation of centres-of-geometry). Finally, if unplit = TRUE the coordinates of the centres are unsplitted using the unsplit function to assign to each atom the coordinates of the centre to which they are attached (used for wrapping by groups).

For objects of class 'atoms' and 'pdb' by default factor is set to x\$resid and x\$coordinates\$resid, respectively, to compute the centre-of-geometry of the different resdiues. Notice that coordinates can be neglected for the calculation of the centres using NA values in factor.

# Value

Return an object of class 'coords' containing the coordinates of centres.

# See Also

```
coords, atoms, pdb, elements
```

and split, unsplit, factor for details about splitting data sets.

# Examples

```
# First lets read a pdb file
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))</pre>
```

# Centres-of-geometry of the residues

#### conect

```
centres(x)
# Centre-of-geometry of the whole structure
centres(x, factor = rep(1, natom(x)))
# or
centres(coords(x))
# Centres-of-geometry of the PCB and DCB residues
centres(x, factor = x$atoms$resname)
# Knowing the name of the elements forming
# the C60 of the PCBM molecules (PCB residues)
# we can compute the centres-of-geometry of
# the C60 by neglecting the other atoms of the
# PCB residues.
C60.elename <- paste0("C", sprintf("%0.3d", 1:60))</pre>
is.PCB <- x$atoms$resname == "PCB" # Produce a mask to select only the PCB residues
is.C60 <- is.PCB & x$atoms$elename %in% C60.elename # Produce a mask to keep only the C60
F <- x$atoms$resid # We use the residue IDs to split the coordinates
F[!is.C60] <- NA # We keep only the atoms of the C60
C60.centres <- centres(x, factor = F)
# Lets check the position of the C60 centres
visualize(x , mode = NULL)
spheres3d(C60.centres)
text3d(Ty(C60.centres, 2), text=paste0("PCB_", rownames(C60.centres)), cex=2)
# Centres-of-mass of the resdiues
symb <- toSymbols(x$atoms$elename) # Convert elename into elemental symbols</pre>
# Find the mass of the element in the periodic table
w <- elements[match(symb, elements[,"symb"]), "mass"]</pre>
centres(x, weights = w)
```

conect

Create 'conect' Object

# Description

Creates an object of class 'conect' containing the IDs of bonded atoms defining the connectivity of a molecular system.

#### Usage

conect(...)

## Default S3 method:

conect

```
conect(eleid.1, eleid.2, ...)
## S3 method for class 'coords'
conect(x, radii = 0.75, safety = 1.2, by.block = FALSE, ...)
## S3 method for class 'pdb'
conect(x, safety = 1.2, by.block = FALSE, ...)
```

is.conect(x)

#### Arguments

	arguments passed to methods.
eleid.1	a integer vector containing the IDs of bonded atoms.
eleid.2	a integer vector containing the IDs of bonded atoms.
х	an R object containing atomic coordinates.
radii	a numeric vector containing atomic radii used to find neigbours.
safety	a numeric value used to extend the atomic radii.
by.block	a logical value indicating whether the connectivity has to be determine by block (see details).

# Details

conect is a generic function to create objects of class 'conect'. The purpose of this class is to store CONECT records from PDB files, indicating the connectivity of a molecular system.

The default method creates a conect object from its different components, i.e.: eleid.1 and eleid.2. Both arguments have to be specified.

The S3 method for object of class 'coords' determine the connectivity from atomic coordinates. A distance matrix is computed, then, for each pair of atom the distance is compared to a bounding distance computed from atomic radii. If this distance is lower than the bounding distance then the atoms are assumed to be connected.

The S3 method for object of class 'pdb' first use element names to search for atomic radii in the elements data set. Then atomic coordinates and radii are passed to conect.coords.

If by.block == TRUE, a grid is defined to determined the connectivity by block. The method is slow but allow to deal with very large systems.

is.conect tests if an object of class 'conect', i.e. if it has a "class" attribute equal to conect.

#### Value

conect returns a two-column data.frame of class 'conect' whose rows contain the IDs of bonded atoms. The columns of this data.frame are described below:

- eleid.1 a integer vector containing the elements IDs defining the connectivity of the system.
- eleid.2 a integer vector containing the elements IDs defining the connectivity of the system.

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coords

is.conect returns TRUE if x is an object of class 'coords' and FALSE otherwise.

#### See Also

pdb

# Examples

```
# If atom 1 is connected to atom 2, 3, 4 and 5
# then we can prepare the following 'conect' object:
x <- conect(rep(1,4), 2:5)
print(x)
is.conect(x)
# Compute conectivity from coordinates
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"), CONECT = FALSE)
x$conect
x$conect <- conect(x)
x$conect</pre>
```

coords

The Atomic Coordinates of an Object

## Description

Get or set the atomic coordinates (either Cartesian or fractional coordinates) of an object.

#### Usage

```
coords(...)
coords(x) <- value
## Default S3 method:
coords(x1, x2, x3, basis = "xyz", ...)
## S3 method for class 'data.frame'
coords(x, basis = NULL, ...)
## S3 method for class 'matrix'
coords(x, basis = NULL, ...)
## S3 method for class 'atoms'
coords(x, ...)
## S3 replacement method for class 'atoms'
coords(x) <- value</pre>
```

coords

```
## S3 method for class 'pdb'
coords(x, ...)
## S3 replacement method for class 'pdb'
coords(x) <- value
is.coords(x)</pre>
```

#### Arguments

	further arguments passed to or from other methods.
x	an R object containing atomic coordinates.
value	an object of class 'coords' used for replacement
x1, x2, x3	numeric vectors containing the first, second and third coordinates.
basis	a single element character vector indicating the type of basis vector used to express the atomic coordinates.

# Details

The purpose of the 'coords' class is to store the coordinates of a molecular system and facilitate their manipulation when passing from the Cartesian to fractional coordinates and vice versa. coords and coords<- are generic accessor and replacement functions.

The default method of the coords function is actually a builder allowing to create a 'coords' object from its different components, i.e.: x1, x2, x3, and basis. All the arguments have to be specified except 'basis' which by default is set to "xyz" (Cartesian coordinates).

For an object of class 'atoms', the accessor function extracts its x1, x2 and x3 components as well as its basis attribute to create a 'coords' object. The replacement function set its x1, x2 and x3 components as well as its basis attribute.

For an object of class 'pdb', the accessor function extracts the x1, x2 and x3 components as well as the basis attribute of its atoms component to create a 'coords' object. The replacement function set the x1, x2 and x3 components as well as the basis attribute of its atoms component.

For 'matrix' and 'data.frame' objects, when basis==NULL this function search x, y, z or a, b, c columns in x.

If x, y, z columns are found they are used to a set the first, second and third coordinates of the returned 'coords' object. In that case the basis set of x is set to "xyz".

If a, b, c columns are found they are used to a set the first, second and third coordinates of the returned 'coords' object. In that case the basis set of x is set to "abc".

If the function doesn't found neither the x, y, z nor the a, b, c columns an error is returned.

When basis!=NULL it has to be equal to "xyz" or "abc" and x must have exactly 3 columns.

is.coords tests if x is an object of class 'coords', i.e. if x has a "class" attribute equal to coords.

#### cryst1

# Value

The accessor function returns a data.frame of class 'coords' whose columns contain the three coordinates of the atoms of a molecular system. The coordinates can either be Cartesian (basis attribute equal to "xyz") or fractional coordinates (basis attribute equal to "abc").

The replacement function returns an object of the same class as x with updated coordinates.

is.coords returns TRUE if x is an object of class 'coords' and FALSE otherwise

## See Also

basis

# Examples

```
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
is.coords(x)
is.coords(x$atoms)
## Replace the coordinates of x by translated coordinates
coords(x) <- coords(Tz(x, 10))</pre>
```

cryst1
--------

coords(x)

Create 'cryst1' Object

# Description

Create an object of class 'cryst1' containing the unit cell parameters and the name of the space group to associate with an object of class 'pdb'.

# Usage

```
cryst1(...)
## Default S3 method:
cryst1(abc, abg = c(90, 90, 90), sgroup = "P1", ...)
```

```
is.cryst1(x)
```

# Arguments

	further arguments passed to or from other methods.
abc	a numeric vector of length 3 containing the norms of the lattice vectors a, b and
	с.

distances

abg	a numeric vector of length 3 containing the angles between the lattice vectors $\alpha$ , $\beta$ and $\gamma$ .
sgroup	a character string giving the Hermann-Mauguin symbol of the space group.
x	an R object to be tested.

# Details

cryst1 is a generic function to create objects of class 'cryst1'. The purpose of this class is to store CRYST1 records from PDB files which contain the unit cell parameters and the name of the space group of a molecular system stored in a PDB file. The default method of the cryst1 function creates an object of class 'cryst1' from its different components, i.e.: abc, abg and sgroup. At least abc has to be specified.

is.cryst1 tests if an object is of class 'cryst1', i.e. if it has a "class" attribute equal to cryst1.

# Value

Function cryst1 returns a list of class 'cryst1' with the following components:

abc	a numeric vector of length 3 containing the norms of the lattice vectors a, b and c.
abg	a numeric vector of length 3 containing the angles between the lattice vectors $\alpha,$ $\beta$ and $\gamma.$
sgroup	a character string giving the Hermann-Mauguin symbol of the space group.
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Function is.cryst1 returns TRUE if x is an object of class 'cryst1' and FALSE otherwise.

# See Also

cell.coords,pdb

# Examples

```
x <- cryst1(abc = c(10, 10, 10), abg = c(90,90,90), sgroup = "P1")
is.cryst1(x)</pre>
```

distances

Inter-Atomic Distances

## Description

Computes inter-atomic distance vectors.

# distances

# Usage

```
distances(...)
## Default S3 method:
distances(
  dx1 = numeric(0),
  dx2 = numeric(0),
  dx3 = numeric(0),
  basis = "xyz",
  . . .
)
## S3 method for class 'coords'
distances(x, sel1, sel2, ...)
## S3 method for class 'atoms'
distances(x, sel1, sel2, ...)
## S3 method for class 'pdb'
distances(x, sel1, sel2, ...)
is.distances(x)
norm(...)
## S3 method for class 'distances'
norm(x, type = "xyz", ...)
```

#### Arguments

	further arguments passed to or from other methods.
dx1, dx2, dx3	numeric arrays containing the first, second and third components of the distance vectors.
basis	a single element character vector indicating the type of basis vector used to express the coordinates.
x	an R object containing atomic coordinates.
sel1, sel2	integer or logical vectors defining two atomic selections between which the distance vectors are computed.
type	a single element character vector indicating how to project the distances vectors before computing the norms. See details.

# Details

The purpose of the 'distances' class is to store the inter-atomic distance vectors and facilitate their manipulation when passing from the Cartesian to fractional references and vice versa. The default method of the distances function is actually a builder allowing to create a 'distances'

object from its different components, i.e.: dx1, dx2, dx3, and basis. All the arguments have to be specified except 'basis' which by default is set to "xyz" (Cartesian reference).

For objects of class 'coords', 'atoms', 'pdb', two sets of atomic coordinates, defined by sel1 and sel2, are extracted and inter-atomic distance vectors are computed between these two sets.

The method of the norm function for objects of class 'distances' computes the norm of the distances vectors. type specify how to project the distance vectors before computing the norms. By default no projection is perform. The three dx, dy, and dz components of the distance vectors are used to compute the norm. type can take the following values:

- x: The distance vectors are projected over x.
- y: The distance vectors are projected over y.
- z: The distance vectors are projected over z.
- xy: The distance vectors are projected in the xy-plan.
- yz: The distance vectors are projected in the yz-plan.
- zx: The distance vectors are projected in the zx-plan.
- xyz: The distance vectors are not projected (The three components of the distance vectors are used to compute the norm).

is.distances tests if x is an object of class 'distances', i.e. if x has a "class" attribute equal to distances.

# Value

The distance function return an object of class 'distances' containing inter-atomic distance vectors. The norm function return an array, with the same dimensions as the dx1, dx2, dx3 components of the 'distances' object for which the norms have to be computed, containing the norm of the distance vectors.

is. distances returns TRUE if x is an object of class 'distances' and FALSE otherwise

# See Also

coords, basis, xyz2abc, abc2xyz

# Examples

```
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
is.DCB7 <- x$atoms$resname == "DCB" & x$atoms$resid == 7
is.DCB8 <- x$atoms$resname == "DCB" & x$atoms$resid == 8
d <- distances(x, is.DCB7, is.DCB8)
norm(d, type = "xyz")
norm(d, type = "xy")
norm(d, type = "x")</pre>
```

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elements

# Description

This data set gives various information on chemical elements

#### Format

A data frame containing for each chemical element the following information.

num atomic number

symb elemental symbol

areneg Allred and Rochow electronegativity (0.0 if unknown)

rcov covalent radii (in Angstrom) (1.6 if unknown)

rbo "bond order" radii

rvdw van der Waals radii (in Angstrom) (2.0 if unknown)

maxbnd maximum bond valence (6 if unknown)

mass IUPAC recommended atomic masses (in amu)

elneg Pauling electronegativity (0.0 if unknown)

ionization ionization potential (in eV) (0.0 if unknown)

elaffinity electron affinity (in eV) (0.0 if unknown)

red red value for visualization

green green value for visualization

blue blue value for visualization

name element name

#### Source

Open Babel (2.3.1) file: element.txt

Created from the Blue Obelisk Cheminformatics Data Repository Direct Source: http://www.blueobelisk.org/ http://www.blueobelisk.org/repos/blueobelisk/elements.xml includes further bibliographic citation information

- Allred and Rochow Electronegativity from http://www.hull.ac.uk/chemistry/electroneg.php?type=Allred-Rochow

- Covalent radii from http://dx.doi.org/10.1039/b801115j

- Van der Waals radii from http://dx.doi.org/10.1021/jp8111556

inertia

# Examples

```
data(elements)
elements
# Get the mass of some elements
symb <- c("C","O","H")
elements[match(symb, elements[,"symb"]),"mass"]
# Get the van der Waals radii of some elements
symb <- c("C","O","H")
elements[match(symb, elements[,"symb"]),"rvdw"]</pre>
```

inertia

#### Moment of Inertia of a Molecular System

#### Description

Computes the inertia tensor of a molecular system from atomic coordinates and masses.

#### Usage

```
inertia(...)
## S3 method for class 'coords'
inertia(x, m = NULL, ...)
## S3 method for class 'atoms'
inertia(x, m = NULL, ...)
## S3 method for class 'pdb'
inertia(x, m = NULL, ...)
```

# Arguments

	further arguments passed to or from other methods.
x	an R object containing atomic coordinates.
m	a numeric vector containing atomic masses.

# Details

inertia is a generic function to compute the inertia tensor of a molecular system. For object of class 'coords' both atomic coordinates and masses have to be speifyed. For object of class 'atoms' the masses are determined from the elename component of the object (see toSymbols and masses). For object of class 'pdb' the atoms component is used.

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#### masses

# Value

Return the inertia tensor in a 3x3 matrix.

# See Also

toSymbols, masses, viewInertia

# Examples

```
C70 <- read.pdb(system.file("examples/C70.pdb", package="Rpdb"))
inertia(C70)
visualize(C70, mode = NULL)
viewXY()
viewInertia(C70)</pre>
```

masses

# Mass of Chemical Elements

# Description

masses(...)

masses(x, ...)

Determine the mass of chemical elements

# Usage

```
## Default S3 method:
masses(x, ...)
## S3 method for class 'pdb'
```

#### Arguments

	further arguments passed to or from other methods.
х	either a character or an integer vector containing element symbols or atomic
	numbers, or an object of class 'pdb' from which element symbols are determined
	(see details).

#### Details

masses is a generic function to determine the mass of chemical elements.

For objects of class 'pdb':

- First the element names are converted into element symbols using the toSymbols function.
- Then their masses are taken from the elements data set.

NA values are returned for unrecognized elements.

#### Value

Return a numeric vector containing the mass of chemical elements.

#### See Also

toSymbols

# Examples

```
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb",package="Rpdb"))
masses(x)</pre>
```

```
masses(c("C","Cl",NA,"AA","N"))
```

merge.coords Merging Molecular Systems

# Description

Merge two objects contaning atomic coordinates

#### Usage

```
## S3 method for class 'coords'
merge(x, y, ...)
## S3 method for class 'atoms'
merge(x, y, reindex = TRUE, ...)
## S3 method for class 'pdb'
merge(x, y, reindex = TRUE, ...)
```

## Arguments

х, у	objects of class 'coords' to be merged.
	further arguments passed to or from other methods.
reindex	a single element logical vector indicating if residue and element IDs have to be reindexed after merging.

# Details

To merge x and y they must have the same basis attributes (see basis).

For objects of class 'coords' and 'atoms' the atomic coordinates are directly merged by row.

For objects of class 'pdb', the atoms and conect components of the two pdb objects are merged by

#### mirror

row and the cryst1 components of x is used to build the returned object.

For objects of class 'atoms' and 'pdb' the residue and element IDs of y are shifted to avoid any confusion with those of x. If reindex == TRUE the reindex function is called to reinitialize the indexing of the returned object.

#### Value

Return an object of the same class as x and y merging x and y. If x and y have different basis attributes an error is returned.

# See Also

coords, atoms, pdb, basis, merge, merge.data.frame

# Examples

```
c1 <- coords( 1:3 , 4:6 , 7:9 , basis = "xyz")
c2 <- coords(10:12, 13:15, 16:18, basis = "xyz")
merge(c1,c2)</pre>
```

```
## Merging objects with different basis sets returns an error.
c2 <- coords(9:11, 12:14, 15:17, basis = "abc")
try(merge(c1,c2))
```

```
## Prepare a Pentacene/C70 dimer
C70 <- read.pdb(system.file("examples/C70.pdb", package="Rpdb"))
Pen <- read.pdb(system.file("examples/Pentacene.pdb", package="Rpdb"))
x <- merge(Tz(C70, 3.5, thickness=0.5), Pen)</pre>
```

mirror

Reflexion of Atomic Coordinates

#### Description

Perform a reflexion (or mirror) operation on atomic coordinates with respect to a given reflexion plane.

#### Usage

mirror(...)
## S3 method for class 'coords'
mirror(x, p1, p2 = NULL, p3 = NULL, mask = TRUE, cryst1 = NULL, ...)
## S3 method for class 'pdb'
mirror(x, p1, p2 = NULL, p3 = NULL, mask = TRUE, cryst1 = x\$cryst1, ...)

## Arguments

	further arguments passed to or from other methods.
х	an R object containing atomic coordinates.
p1	a numeric vector of length 3 containing the coordinates of the first point defining the reflexion plan. Can also be a 3x3 matrix or data.frame containing by row p1, p2 and p3.
p2	a numeric vector of length 3 containing the coordinates of the second point defin- ing the reflexion plane.
р3	a numeric vector of length 3 containing the coordinates of the third point defin- ing the reflexion plane.
mask	a logical vector indicating the set of coordinates to which to apply the reflexion.
cryst1	an object of class 'cryst1' use to convert fractional into Cartesian coordinates when need.

# Details

mirror is a generic function. Method for objects of class 'coords' first convert the coordinates into Cartesian coordinates using cryst1 if needed. Once reflected, the coordinates are reconverted back to the orginal basis set using again cryst1. Method for objects of class 'pdb' first extract coordinates from the object using the function coords, perform the reflection, and update the coordinates of the 'pdb' object using the function coords<-.

#### Value

An object of the same class as x with reflected coordinates.

# See Also

Helper functions for reflection with respect to a given Cartesian plane or a plane defined by two lattice vectors:

Mxy, Myz, Mzx, Mab, Mbc, Mca Passing from Cartesian to fractional coordinates (or Vis Versa): xyz2abc, abc2xyz

#### Examples

```
# First lets read a pdb file
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
cell <- cell.coords(x)
visualize(x, mode = NULL)</pre>
```

```
# Mirror operation with respect to the ab-plane
visualize(mirror(x, rep(0,3), p1=cell[, "a"], p2=cell[, "b"]), mode = NULL)
# Mirror operation with respect to the ab-plane for residue 1
visualize(mirror(x, rep(0,3), p1=cell[, "a"], p2=cell[, "b"],
    mask = x$atoms$resid == 1), mode = NULL)
```

mirrorHelpers

#### Description

Reflection of atomic coordinates with respect to a specific Cartesian plan or a plan defined by two lattice vectors.

# Usage

```
Mxy(...)
## S3 method for class 'coords'
Mxy(x, mask = TRUE, cryst1 = NULL, ...)
## S3 method for class 'pdb'
Mxy(x, mask = TRUE, cryst1 = x$cryst1, ...)
Myz(...)
## S3 method for class 'coords'
Myz(x, mask = TRUE, cryst1 = NULL, ...)
## S3 method for class 'pdb'
Myz(x, mask = TRUE, cryst1 = x$cryst1, ...)
Mzx(...)
## S3 method for class 'coords'
Mzx(x, mask = TRUE, cryst1 = NULL, ...)
## S3 method for class 'pdb'
Mzx(x, mask = TRUE, cryst1 = x$cryst1, ...)
Mab(...)
## S3 method for class 'coords'
Mab(x, cryst1, mask = TRUE, ...)
## S3 method for class 'pdb'
Mab(x, cryst1 = x$cryst1, mask = TRUE, ...)
Mbc(...)
## S3 method for class 'coords'
Mbc(x, cryst1, mask = TRUE, ...)
```

```
## S3 method for class 'pdb'
Mbc(x, cryst1 = x$cryst1, mask = TRUE, ...)
Mca(...)
## S3 method for class 'coords'
Mca(x, cryst1, mask = TRUE, ...)
## S3 method for class 'pdb'
Mca(x, cryst1 = x$cryst1, mask = TRUE, ...)
```

#### Arguments

	further arguments passed to or from other methods.
x	an R object containing atomic coordinates.
mask	a logical vector indicating the set of coordinates to which to apply the reflection.
cryst1	an object of class 'cryst1' use to convert fractional into Cartesian coordinates when need.

# Details

These functions are helper functions to perform a reflection with respect to a specific Cartesian plan or a plan defined by two lattice vectors. All of them call the mirror function.

# Value

An object of the same class as x with reflected coordinates.

# See Also

mirror and xyz2abc, abc2xyz for passing from Cartesian to fractional coordinates (or Vis Versa).

# Examples

```
# First lets read a pdb file
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
visualize(x, mode = NULL)
# Mirror operation with respect to the ab-plane
visualize(Mab(x), mode = NULL)
# Mirror operation with respect to the ab-plane for residue 1
visualize(Mab(x, mask = x$atoms$resid == 1), mode = NULL)</pre>
```

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natom

#### Description

Evaluates the number of atoms in an object containing atomic coordinates.

# Usage

```
natom(x, ...)
## S3 method for class 'coords'
natom(x, factor = NULL, ...)
## S3 method for class 'atoms'
natom(x, factor = NULL, ATOM = TRUE, HETATM = TRUE, ...)
## S3 method for class 'pdb'
natom(x, factor = NULL, ATOM = TRUE, HETATM = TRUE, ...)
```

#### Arguments

х	an R object containing atomic coordinates.
	further arguments passed to or from other methods.
factor	a factor used to split the object and evaluate the number of atoms in each group.
АТОМ	a single element logical vector indicating if ATOM records have to be considered or not.
HETATM	a single element logical vector indicating if HETATM records have to be con- sidered or not.

#### Details

natom is a generic function to evalute the number of atom in an object containing atomic coordinates. The atomic coordinates of the object are first filtered to keep ATOM and/or HETATM records as indicated by the 'ATOM' and 'HETATM' arguments. Then, if factor is specify, the object is splitted to evalute the number of atoms in each group defined by factor. If factor is not specify then the total number of atoms in the object is return.

# Value

Return an integer or a vector of integer of lenght equal to nlevels(factor) (if factor is specify) indication the number of atoms in the object or in the groups defined by factor.

# See Also

coords, atoms, pdb, factor, split

# Examples

```
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))</pre>
```

```
natom(x)
natom(x, x$atoms$resid)
natom(x, x$atoms$resname)
natom(x, HETATM = FALSE)
```

pdb

Create an Object of Class 'pdb'

#### Description

Creates an object of class 'pdb'.

# Usage

pdb(...)

```
## Default S3 method:
pdb(atoms, cryst1 = NULL, conect = NULL, remark = NULL, title = NULL, ...)
```

is.pdb(x)

#### Arguments

	further arguments passed to or from other methods.
atoms	a data.frame of class atoms containing ATOM and HETATM records use to create the pdb object.
cryst1	a list of class cryst1 containing the periodical bondary conditions and space group use to create the pdb object.
conect	a data.frame of class conect containing CONECT records use to create the pdb object.
remark	a character vector containing some REMARK records to be added to the pdb object.
title	a character vector containing some TITLE records to be added to the pdb object.
x	an R object to be tested

# Details

This function is the generic function to create objects of class 'pdb'. The purpose of this class is to store the data of molecular systems contained in PDB files. The default method of the pdb function creates an object of class 'pdb' from its different components, i.e.: title, remark, cryst1, atoms and conect. At least an object of class 'atoms' has to be specified.

is.pdb tests if x is an object of class 'pdb', i.e. if x has a "class" attribute equal to pdb.

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#### range.coords

#### Value

pdb returns a list of class 'pdb' with the following components:

title	a character vector containing the TITLE records found in a PDB file.
remark	a character vector containing the REMARK records found in a PDB file.
cryst1	a list of class 'cryst1' containing the first CRYST1 record found in a PDB file. All others are ignored.
atoms	a data.frame of class 'atoms' containing the ATOM and HETATM records found in a PDB file.
conect	a data.frame of class 'conect' containing the CONECT records found in a PDB file.

is.pdb returns TRUE if x is an object of class 'pdb' and FALSE otherwise.

# See Also

atoms, coords, cryst1, conect and read.pdb

# Examples

range.coords Ran

# Range of Atomic Coordinates

# Description

Determines the range of atomic coordinates.

# Usage

```
## S3 method for class 'coords'
range(x, na.rm = FALSE, finite = FALSE, ...)
## S3 method for class 'atoms'
range(x, na.rm = FALSE, finite = FALSE, ...)
```

# read.pdb

```
## S3 method for class 'pdb'
range(x, na.rm = FALSE, finite = FALSE, ...)
```

# Arguments

х	an R object containing atomic coordinates.
na.rm	logical, indicating if NA's should be omitted.
finite	logical, indicating if all non-finite elements should be omitted.
	further arguments passed to or from other methods.

# Value

Return a data.frame whose columns contain the range of the first, second and third coordinates of x.

# See Also

range, coords, atoms, pdb

# Examples

```
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
range(x)
range(range(x))</pre>
```

read.pdb

PDB File Reader

# Description

Reads a Protein Data Bank (PDB) coordinate file.

# Usage

```
read.pdb(
  file,
  ATOM = TRUE,
  HETATM = TRUE,
  CRYST1 = TRUE,
  CONECT = TRUE,
  TITLE = TRUE,
  REMARK = TRUE,
  MODEL = 1
)
```

# read.pdb

# Arguments

file	a single element character vector containing the name of the PDB file to be read.
АТОМ	a single element logical vector indicating whether ATOM records have to be read.
HETATM	a single element logical vector indicating whether HETATM records have to be read.
CRYST1	a single element logical vector indicating whether CRYST1 records have to be read.
CONECT	a single element logical vector indicating whether CONECT records have to be read.
TITLE	a single element logical vector indicating whether TITLE records have to be read.
REMARK	a single element logical vector indicating whether REMARK records have to be read.
MODEL	an integer vector containing the serial number of the MODEL sections to be read. Can also be equal to NULL to read all the MODEL sections or to NA to ignore MODEL records (see details).

# Details

The read.pdb function reads the TITLE, REMARK, ATOM, HETATM, CRYST1 and CONECT records from a PDB file. Three different reading modes can be used depending on the value of MODEL:

- When MODEL is a vector of integers, MODEL sections whose serial numbers match these integers are read.
- When MODEL == NULL, all MODEL sections are read.
- When MODEL == NA, MODEL records are ignored to read all ATOM and/or HETATM records together to return a single object.

# Value

When a single MODEL section is read, this function returns an object of class 'pdb' (a list with a class attribute equal to pdb) with the following components:

title	a character vector containing the TITLE records found in the PDB file.
remark	a character vector containing the REMARK records found in the PDB file.
cryst1	a list of class 'cryst1' containing the first CRYST1 record found in the PDB file. All others are ignored.
atoms	a data.frame of class 'atoms' containing the ATOM and HETATM records found in the PDB file.
conect	a data.frame of class 'conect' containing the CONECT records found in the PDB file.

When multiple MODEL sections are read, a list of object of class 'pdb' is returned.

## References

PDB format has been taken from: http://www.wwpdb.org/documentation/format33/v3.3.html

## See Also

write.pdb, pdb, cryst1, atoms, conect

# Examples

```
# Read a PDB file included with the package
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
# Visualize the PDB file
visualize(x, mode = NULL)
# Write the 'pdb' object 'x' in file "Rpdb.pdb" into the current directory
write.pdb(x, file = "Rpdb.pdb")
# Cleanup
unlink("Rpdb.pdb")</pre>
```

reindex

#### Reinitialize Object Indexing

# Description

Reinitialize the indexing of an object.

#### Usage

```
reindex(...)
## S3 method for class 'atoms'
reindex(x, eleid = TRUE, resid = TRUE, ...)
## S3 method for class 'pdb'
reindex(x, eleid = TRUE, resid = TRUE, ...)
```

#### Arguments

	further arguments passed to or from other methods.
x	an R object.
eleid	a single element logical vector indicating if elements IDs have to reindexed.
resid	a single element logical vector indicating if residues IDs have to reindexed.
#### replicate

#### Details

reindex is a generic function to reinitialize the indexing of an object or its components. The methods for objects of class 'atoms' reinitialize the residue and element IDs starting from 1 and avoiding gaps in the indexes. For objects of class 'pdb' their atoms and conect components are reindexed consistently.

## Value

Return an object of the same class as x with updated indexes.

#### See Also

pdb, atoms, subset.atoms, subset.pdb

## Examples

```
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
x <- subset(x, x$atoms$eleid %in% sample(x$atoms$eleid, 10))
print(x)
x <- reindex(x)
print(x)</pre>
```

replicate

Replicate Atomic Coordinates

## Description

Replicate atomic coordinates using periodic boundary conditions.

```
replicate(x, ...)
## S3 method for class 'coords'
replicate(x, cryst1 = NULL, a.ind = 0, b.ind = 0, c.ind = 0, ...)
## S3 method for class 'atoms'
replicate(x, cryst1 = NULL, a.ind = 0, b.ind = 0, c.ind = 0, ...)
## S3 method for class 'pdb'
replicate(x, a.ind = 0, b.ind = 0, c.ind = 0, cryst1 = NULL, ...)
```

## Arguments

x	an R object containing atomic coordinates to be replicated.
	further arguments passed to or from other methods.
cryst1	an object of class 'cryst1' containing periodical boundary conditions used for replicating.
a.ind	a vector of integers indicating the positions of the replicated cells along the a- axis.
b.ind	a vector of integers indicating the positions of the replicated cells along the b- axis.
c.ind	a vector of integers indicating the positions of the replicated cells along the c-axis.

#### Details

The replicate function replicate a unit cell along the lattice vectors a, b and c as as many times as indicated by the a.ind, b.ind and c.ind arguments. Discontinuous integer vectors can be used for a.ind, b.ind and c.ind to create layered supercells (See examples).

## Value

Return an object of class 'pdb' with replicated atomic coordinates.

## See Also

coords, atoms, pdb, cryst1

#### Examples

```
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))</pre>
```

```
# Create a 3x3 supercell
y <- replicate(x, a.ind = 0:2, b.ind = 0:2, c.ind = 0:2)</pre>
```

```
# Create a 3x3 supercell which might need to be wrapped (if some molecules are outside the cell) y <- replicate(x, a.ind = -1:1, b.ind = -1:1, c.ind = -1:1)
```

```
# Create a layered supercell with a vacuum layer in the bc-plan
y <- replicate(x, a.ind = c(0,2), b.ind = 0:2, c.ind = 0:2)</pre>
```

rotation

## Description

Rotation of atomic coordinates around a given vector.

#### Usage

```
R(...)
## S3 method for class 'coords'
R(obj, angle = 0, x = 0, y = 0, z = 1, mask = TRUE, cryst1 = NULL, ...)
## S3 method for class 'pdb'
R(obj, angle = 0, x = 0, y = 0, z = 1, mask = TRUE, cryst1 = obj$cryst1, ...)
```

## Arguments

	further arguments passed to or from other methods.
obj	an R object containing atomic coordinates.
angle	the angle of the rotation in degrees.
x	the x-component of the rotation vector.
У	the y-component of the rotation vector.
z	the z-component of the rotation vector.
mask	a logical vector indicating the set of coordinates to which the rotation has to be applyed.
cryst1	an object of class 'cryst1' use to convert fractional into Cartesian coordinates when need.

#### Details

R is generic functions. Method for objects of class 'coords' first convert the coordinates into Cartesian coordinates using cryst1 if needed. Once rotated, the coordinates are reconverted back to the orginal basis set using again cryst1. Method for objects of class 'pdb' first extract coordinates from the object using the function coords, perform the rotation, and update the coordinates of the 'pdb' object using the function coords<-.

#### Value

An object of the same class as x with rotated coordinates.

# See Also

Helper functions for rotation around a given Cartesian vector: Rx, Ry, Rz Passing from Cartesian to fractional coordinates (or Vis Versa): xyz2abc, abc2xyz

## Examples

```
# First lets read a pdb file
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
cell <- cell.coords(x)
visualize(x, mode = NULL)
# Rotation of the structure around the c-axis
visualize(R(x, 90, x=cell["x","c"], y=cell["y","c"], z=cell["z","c"]),
            mode = NULL)
# Rotation of the residue 1 around the c-axis
visualize(R(x, 90, x=cell["x","c"], y=cell["y","c"], z=cell["z","c"], mask = x$atoms$resid == 1),
            mode = NULL)</pre>
```

rotationHelpers Helper Functions for Rotation of Atomic Coordinates

#### Description

Rotation of atomic coordinates along a specific Cartesian vector.

#### Usage

```
Rx(...)
## S3 method for class 'coords'
Rx(x, angle = 0, mask = TRUE, cryst1 = NULL, ...)
## S3 method for class 'pdb'
Rx(x, angle = 0, mask = TRUE, cryst1 = x$cryst1, ...)
Ry(...)
## S3 method for class 'coords'
Ry(x, angle = 0, mask = TRUE, cryst1 = NULL, ...)
## S3 method for class 'pdb'
Ry(x, angle = 0, mask = TRUE, cryst1 = x$cryst1, ...)
Rz(...)
```

# rotationHelpers

```
## S3 method for class 'coords'
Rz(x, angle = 0, mask = TRUE, cryst1 = NULL, ...)
## S3 method for class 'pdb'
Rz(x, angle = 0, mask = TRUE, cryst1 = x$cryst1, ...)
```

## Arguments

	further arguments passed to or from other methods.
x	an R object containing atomic coordinates.
angle	the angle of the rotation in degrees.
mask	a logical vector indicating the set of coordinates to which the rotation has to be applyed.
cryst1	an object of class 'cryst1' use to convert fractional into Cartesian coordinates when need.

## Details

These functions are helper functions to perform a rotation around a specific Cartesian vector. All of them call the R function.

#### Value

An object of the same class as x with rotated coordinates.

## See Also

R and xyz2abc, abc2xyz for passing from Cartesian to fractional coordinates (or Vis Versa).

## Examples

```
# First lets read a pdb file
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
cell <- cell.coords(x)
visualize(x, mode = NULL)
# Rotation of the structure around the z-axis
visualize(Rz(x, 90), mode = NULL)</pre>
```

```
# Rotation of the residue 1 around the c-axis
visualize(Rz(x, 90, mask = x$atoms$resid == 1), mode = NULL)
```

split.pdb

#### Description

split divides a 'pdb' object by groups of atoms defined by f. unsplit reverses the effect of split.

## Usage

```
## S3 method for class 'pdb'
split(x, f, drop = FALSE, ...)
```

```
## S3 method for class 'pdb'
unsplit(value, f, drop = FALSE, ...)
```

#### Arguments

х	an object of class 'pdb' to be divided into groups.
f	a 'factor' in the sense that as.factor(f) defines the grouping, or a list of such factors in which case their interaction is used for the grouping.
drop	logical indicating if levels that do not occur should be dropped (if f is a factor or a list).
	further potential arguments passed to methods.
value	a list of 'pdb' objects compatible with a splitting of x. Recycling applies if the lengths do not match.

# Details

split produce a list of 'pdb' objects with the same cryst1, title and remark components as x. Only its atoms component is splitted while its conect component is cleaned to keep only the meaningful connectivity for each 'pdb' object of the list returned by the function. unlist produce a 'pdb' object with the same cryst1, title and remark components as the first element of value. The atoms and conect components of all the elements of value are combined by row.

## Value

The value returned from split is a list of 'pdb' objects containing the data for the groups of atoms. The components of the list are named by the levels of f (after converting to a factor, or if already a factor and drop=TRUE, dropping unused levels).

unsplit returns a 'pdb' object for which split(x, f) equals value.

## See Also

split, unsplit, pdb

#### subset.atoms

#### Examples

```
### Split a pdb file by residue IDs and write them into separate files
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
file.names <- paste0(x$atoms$resname, "_", x$atoms$resid, ".pdb")
file.names <- unique(file.names)
pdb.resid <- split(x, x$atoms$resid)
length(pdb.resid)
useless <- mapply(write.pdb, pdb.resid, file.names)
# Cleanup
unlink(file.names)
```

subset.atoms Subsetting 'atoms' and 'pdb' Objects

# Description

Return subsets of 'atoms' or 'pdb' objects which meet conditions.

#### Usage

```
## S3 method for class 'atoms'
subset(x, subset, drop = FALSE, reindex.all = TRUE, ...)
## S3 method for class 'pdb'
```

subset(x, subset, drop = FALSE, reindex.all = TRUE, ...)

## Arguments

х	object to be subsetted.
subset	logical expression indicating elements or rows to keep: missing values are taken as false.
drop	passed on to [ indexing operator.
reindex.all	a single element logical vector indicating if residues and elements IDs have to be reindexed after subsetting.
	further arguments to be passed to or from other methods.

#### Details

For an 'atoms' object the method is similar to the data.frame method (see subset) but allow to directly reindex the elements and residues IDs. For a 'pdb' object subsetting is applied on the atoms and conect components of the object in a consistent way. First the atoms component is subsetted and then the conect component is filtered to keep only the conectivity for the subset.

toSymbols

#### Value

Return a subsetted object of the same class as x.

# See Also

subset, pdb, atoms, reindex

# Examples

```
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
y <- subset(x, x$atoms$eleid %in% sample(x$atoms$eleid, 10))
is(y)
y <- subset(x$atoms, x$atoms$eleid %in% sample(x$atoms$eleid, 10))
is(y)
x <- coords(x)
y <- subset(x, x < 0)
is(y)</pre>
```

toSymbols

Atomic Symbols Converter

#### Description

Converts character strings or atomic numbers into atomic symbols.

# Usage

```
toSymbols(x, ...)
## S3 method for class 'integer'
toSymbols(x, ...)
## S3 method for class 'numeric'
toSymbols(x, ...)
## S3 method for class 'character'
toSymbols(x, nletters = 3, ...)
```

## Arguments

х	a vector to be converted into atomic symbols.
	further arguments passed to or from other methods.
nletters	an integer used to truncate the character strings before convertion.

## translation

#### Details

Each elements of x are converted into atomic symbols.

When x is an integer (or numeric) vector, atomic number are search into the elements data set to find associated atomic symbols.

When x is a character vector, toSymbols first removes all leading and trailing white spaces and numbers. Then translates the first character of the character strings to uppercase and all the others to lowercase. Finally, the character strings are tested for matching with element symbols provided by the elements data set. NA are produced for no matching.

#### Value

a character vector containing atomic symbols

#### See Also

elements

# Examples

```
x <- c(1:10)
toSymbols(x)
x <- c("C "," o","h1","1h","UU","SI0","cR")
toSymbols(x)
# 'nletters' can be used to truncate the character strings before
# conversion, if needed:
toSymbols("SIL", nletters=3) # return NA
toSymbols("SIL", nletters=2) # return "Si"
toSymbols("SIL", nletters=1) # return "S"
```

translation

Translation of Atomic Coordinates

#### Description

Translation of Cartesian or fractional coordinates.

```
Txyz(...)
## S3 method for class 'coords'
Txyz(
    obj,
    x = 0,
    y = 0,
```

translation

```
z = 0,
 mask = TRUE,
  thickness = NULL,
  cryst1 = NULL,
  • • •
)
## S3 method for class 'pdb'
Txyz(
 obj,
 x = 0,
 y = 0,
 z = 0,
 mask = TRUE,
  thickness = NULL,
 cryst1 = obj$cryst1,
  . . .
)
Tabc(...)
## S3 method for class 'coords'
Tabc(
 obj,
 a = 0,
 b = 0,
 c = 0,
 mask = TRUE,
 thickness = NULL,
 cryst1 = NULL,
  . . .
)
## S3 method for class 'pdb'
Tabc(
 obj,
 a = 0,
 b = 0,
 c = 0,
 mask = TRUE,
 thickness = NULL,
 cryst1 = obj$cryst1,
  . . .
)
```

# Arguments

... further arguments passed to or from other methods.

## translation

obj	an R object containing atomic coordinates.
х	the x-component of the translation vector.
У	the y-component of the translation vector.
z	the z-component of the translation vector.
mask	a logical vector indicating the set of coordinates to which to apply the transla- tion.
thickness	a numeric value indicating the fraction of the thicknees of the selected atom to be added to the translation vector (Usually 0, 0.5 or 1. See details).
cryst1	an object of class 'cryst1' use to convert Cartesian into fraction coordinates (or Vis Versa) when need.
а	the a-component of the translation vector.
b	the b-component of the translation vector.
С	the c-component of the translation vector.

#### Details

Txyz and Tabc are generic functions. Method for objects of class 'coords' first convert the coordinates into Cartesian or fractional coordinates using cryst1 if needed to performed the translation. Once translated, the coordinates are reconverted back to the orginal basis set using again cryst1. Method for objects of class 'pdb' first extract coordinates from the object using the function coords, perform the translation, and update the coordinates of the 'pdb' object using the function coords<-. The thickness argument can be use to translate selected atoms by a fraction of its thickness along the translation direction. This can be use when merging two fragments centered at the origin to build a dimer to avoid atomic overlap and set the inter-fragment distance (see examples).

#### Value

An object of the same class as x with translated coordinates.

# See Also

Helper functions for translation along given Cartesian or lattice vector: Tx, Ty, Tz, Ta, Tb, Tc Passing from Cartesian to fractional coordinates (or Vis Versa): xyz2abc, abc2xyz

#### Examples

```
# First lets read a pdb file
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
visualize(x, mode = NULL)
visualize(Txyz(x, y=10), mode = NULL)
visualize(Txyz(x, y=10, mask = x$atoms$resid==1), mode = NULL)
visualize(Tabc(x, b=1), mode = NULL)
visualize(Tabc(x, b=1, mask = x$atoms$resid==1), mode = NULL)</pre>
```

# Lets build a C70/Pentacene dimer with an inter-molecular distance equal to 3.5

```
C70 <- read.pdb(system.file("examples/C70.pdb", package="Rpdb"))
Pen <- read.pdb(system.file("examples/Pentacene.pdb", package="Rpdb"))
x <- merge(C70, Pen)
visualize(x, mode = NULL)
viewXY()
visualize(Txyz(x, x=0, y=0, z=3.5, mask = x$atoms$resname == "C70", thickness=0.5), mode = NULL)
viewXY()</pre>
```

translationHelpers Helper Functions for Translation of Atomic Coordinates

#### Description

Translation of atomic coordinates along a specific Cartesian or lattice vector.

```
Tx(...)
## S3 method for class 'coords'
Tx(obj, x = 0, mask = TRUE, thickness = NULL, cryst1 = NULL, ...)
## S3 method for class 'pdb'
Tx(obj, x = 0, mask = TRUE, thickness = NULL, cryst1 = obj$cryst1, ...)
Ty(...)
## S3 method for class 'coords'
Ty(obj, y = 0, mask = TRUE, thickness = NULL, cryst1 = NULL, ...)
## S3 method for class 'pdb'
Ty(obj, y = 0, mask = TRUE, thickness = NULL, cryst1 = obj$cryst1, ...)
Tz(...)
## S3 method for class 'coords'
Tz(obj, z = 0, mask = TRUE, thickness = NULL, cryst1 = NULL, ...)
## S3 method for class 'pdb'
Tz(obj, z = 0, mask = TRUE, thickness = NULL, cryst1 = obj$cryst1, ...)
Ta(...)
## S3 method for class 'coords'
Ta(obj, a = 0, mask = TRUE, cryst1 = NULL, ...)
## S3 method for class 'pdb'
```

```
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```

```
Ta(obj, a = 0, mask = TRUE, cryst1 = obj$cryst1, ...)
Tb(...)
## S3 method for class 'coords'
Tb(obj, b = 0, mask = TRUE, cryst1 = NULL, ...)
## S3 method for class 'pdb'
Tb(obj, b = 0, mask = TRUE, cryst1 = obj$cryst1, ...)
Tc(...)
## S3 method for class 'coords'
Tc(obj, c = 0, mask = TRUE, cryst1 = NULL, ...)
## S3 method for class 'pdb'
Tc(obj, c = 0, mask = TRUE, cryst1 = obj$cryst1, ...)
```

## Arguments

•••	further arguments passed to or from other methods.
obj	an R object containing atomic coordinates.
x	the x-component of the translation vector.
mask	a logical vector indicating the set of coordinates to which to apply the transla- tion.
thickness	a numeric value indicating the fraction of the thicknees of the selected atom to be added to the translation vector (Usually 0, 0.5 or 1. See details).
cryst1	an object of class 'cryst1' use to convert Cartesian into fraction coordinates (or Vis Versa) when need.
У	the y-component of the translation vector.
Z	the z-component of the translation vector.
а	the a-component of the translation vector.
b	the b-component of the translation vector.
с	the c-component of the translation vector.

# Details

These functions are helper functions to perform a translation along a specific Cartesian or lattice vector. All of them call either the Txyz or Tabc function.

#### Value

An object of the same class as x with translated coordinates.

# See Also

Txyz, Tabc Passing from Cartesian to fractional coordinates (or Vis Versa): xyz2abc, abc2xyz

#### Examples

```
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
visualize(x, mode = NULL)
visualize(Ty(x, 10), mode = NULL)
visualize(Tb(x, 1 ), mode = NULL)
visualize(Tb(x, 1 ), mode = NULL)
# Lets build a C70/Pentacene dimer with an inter-molecular distance equal to 3.5
C70 <- read.pdb(system.file("examples/C70.pdb", package="Rpdb"))
Pen <- read.pdb(system.file("examples/Pentacene.pdb", package="Rpdb"))
x <- merge(C70, Pen)
visualize(x, mode = NULL)
visualize(x, mode = NULL)
visualize(Tz(x, z=3.5, mask = x$atoms$resname == "C70", thickness=0.5), mode = NULL)
viewXY()</pre>
```

universalConstants Universal Constants

#### Description

This data set provides various universal constants

## Format

A data frame containing for each universal constant the following information.

Quantity a character vector containing a short description of the constants.

Value a numeric vector containing the value of the constants.

Unit a character vector indicating the unit of the constants.

#### Source

http://www.ebyte.it/library/educards/constants/ConstantsOfPhysicsAndMath.html

#### unsplit

## Examples

```
# Data for the speed of light
universalConstants["c",]
# Return the speed of light in m.s-1
universalConstants["c","Value"]
# Return the Planck constant in J.s
universalConstants["h","Value"]
```

unsplit

#### Reassemble Groups

# Description

unsplit reverses the effect of split.

## Usage

```
unsplit(value, f, drop = FALSE, ...)
## Default S3 method:
unsplit(value, f, drop = FALSE, ...)
```

#### Arguments

value	a list of vectors or data frames compatible with a splitting of x. Recycling applies if the lengths do not match.
f	a 'factor' in the sense that as.factor(f) defines the grouping, or a list of such factors in which case their interaction is used for the grouping.
drop	logical indicating if levels that do not occur should be dropped (if f is a factor or a list).
	further potential arguments passed to methods.

## Details

unsplit is a generic functions with a default method (Method dispatch takes place based on the class of the first element of value) working with lists of vectors or data frames (assumed to have compatible structure, as if created by split). It puts elements or rows back in the positions given by f. In the data frame case, row names are obtained by unsplitting the row name vectors from the elements of value.

f is recycled as necessary and if the length of x is not a multiple of the length of f a warning is printed.

Any missing values in f are dropped together with the corresponding values of x.

#### Value

Returns a vector or data frame for which split(x, f) equals value

#### References

Becker, R. A., Chambers, J. M. and Wilks, A. R. (1988) *The New S Language*. Wadsworth & Brooks/Cole.

## See Also

cut to categorize numeric values. strsplit to split strings.

# Examples

```
require(stats); require(graphics)
n <- 10; nn <- 100
g <- factor(round(n * runif(n * nn)))</pre>
x <- rnorm(n * nn) + sqrt(as.numeric(g))</pre>
xg <- split(x, g)</pre>
boxplot(xg, col = "lavender", notch = TRUE, varwidth = TRUE)
sapply(xg, length)
sapply(xg, mean)
### Calculate 'z-scores' by group (standardize to mean zero, variance one)
z <- unsplit(lapply(split(x, g), scale), g)</pre>
# or
zz <- x
split(zz, g) <- lapply(split(x, g), scale)</pre>
# and check that the within-group std dev is indeed one
tapply(z, g, sd)
tapply(zz, g, sd)
### Data frame variation
# Notice that assignment form is not used since a variable is being added
g <- airquality$Month
l <- split(airquality, g)</pre>
1 <- lapply(1, transform, 0z.Z = scale(0zone))</pre>
aq2 <- unsplit(l, g)</pre>
head(aq2)
with(aq2, tapply(Oz.Z, Month, sd, na.rm=TRUE))
### Split a matrix into a list by columns
ma \le cbind(x = 1:10, y = (-4:5)^2)
split(ma, col(ma))
```

split(1:10, 1:2)

vectorialOperations Basic Vectorial Operations

# Description

Basic vectorial operations such as scalar product and vectorial product

#### Usage

```
dotProd(U, V)
vectNorm(U)
rotVect(U, n = 1)
```

vectProd(U, V)

#### Arguments

U	a numeric vector of length 3.
V	a numeric vector of length 3.
n	an integer.

# Value

- dotProct return a single element numeric vector.
- vectNorm return a single element numeric vector.
- rotVect return a numeric vector of length 3.
- vectProct return a numeric vector of length 3.

## See Also

matmult

## Examples

```
Vx <- c(3,0,0)
vectNorm(Vx)
Vx <- Vx / vectNorm(Vx)
Vy <- c(0,1,0)
Vz <- vectProd(Vx, Vy)
print(Vz)</pre>
```

viewAxis

#### Description

Set the view of the current 'rgl' scene aligning one vector perpendicularly to the screen and placing another one in the horizontal plane.

## Usage

```
viewAxis(V1, V2)
viewXY()
viewYZ()
viewZX()
viewAB(cryst1)
viewBC(cryst1)
viewCA(cryst1)
viewInertia(x, m = NULL)
```

## Arguments

V1	a length 3 numeric vector.
V2	a length 3 numeric vector.
cryst1	an object of class 'cryst1'.
x	an R object containing atomic coordinates.
m	a numeric vector containing atomic masses.

# Details

viewAxis set the view of the current rgl scene (by setting UserMatrix; see par3d for more details) so that V1 is perpendicular to the screen and V2 is in the horizontal plane. The other functions documented here are helper functions calling viewAxis to set the view using particular Cartesian or lattice vectors. For functions viewAB, viewBC and viewCA a 'cryst1' object has to be specifyed to define the lattice vectors used to set the view. The function viewInertia computes the inertia tensor from the atomic coordinates and masses (see inertia) and sets the view to its eigen vectors basis set.

## Value

No return value, called for side effects.

## visualize

# See Also

visualize, cell.coords, par3d, rgl.open

#### Examples

```
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
visualize(x, mode = NULL)
viewAB(x$cryst1)
C70 <- read.pdb(system.file("examples/C70.pdb", package="Rpdb"))
visualize(C70, mode = NULL)
viewXY()
viewInertia(C70)</pre>
```

visualize

#### Visualize a Molecular Structure

# Description

Use the rgl library to visualize in 3D a molecular structure.

```
visualize(...)
## S3 method for class 'coords'
visualize(
  х,
  elename = NULL,
  cryst1 = NULL,
  conect = NULL,
 mode = NULL,
  type = "1",
  xyz = NULL,
  abc = NULL,
  pbc.box = NULL,
  1wd = 2,
  lwd.xyz = lwd,
  lwd.abc = lwd,
  lwd.pbc.box = lwd,
  cex.xyz = 2,
  cex.abc = 2,
  col = NULL,
  bg = "#FAFAD2",
  radii = "rvdw",
  add = FALSE,
```

```
windowRect = c(0, 0, 800, 600),
 FOV = 0,
  userMatrix = diag(4),
  • • •
)
## S3 method for class 'data.frame'
visualize(
  х,
  elename = NULL,
  cryst1 = NULL,
  conect = NULL,
 mode = NULL,
  type = "1",
  xyz = NULL,
  abc = NULL,
  pbc.box = NULL,
  1wd = 2,
  lwd.xyz = lwd,
  lwd.abc = lwd,
  lwd.pbc.box = lwd,
  cex.xyz = 2,
  cex.abc = 2,
  col = NULL,
  bg = "#FAFAD2",
  radii = "rvdw",
  add = FALSE,
 windowRect = c(0, 0, 800, 600),
  FOV = 0,
  userMatrix = diag(4),
  . . .
)
## S3 method for class 'matrix'
visualize(
  х,
  elename = NULL,
  cryst1 = NULL,
  conect = NULL,
 mode = NULL,
  type = "1",
  xyz = NULL,
  abc = NULL,
  pbc.box = NULL,
  1wd = 2,
  lwd.xyz = lwd,
  lwd.abc = lwd,
  lwd.pbc.box = lwd,
```

## visualize

```
cex.xyz = 2,
  cex.abc = 2,
  col = NULL,
  bg = "#FAFAD2",
  radii = "rvdw",
  add = FALSE,
 windowRect = c(0, 0, 800, 600),
 FOV = 0,
 userMatrix = diag(4),
  . . .
)
## S3 method for class 'atoms'
visualize(
  х,
  cryst1 = NULL,
  conect = NULL,
 mode = NULL,
  type = "1",
  xyz = NULL,
  abc = NULL,
  pbc.box = NULL,
  1wd = 2,
  lwd.xyz = lwd,
  lwd.abc = lwd,
  lwd.pbc.box = lwd,
  cex.xyz = 2,
  cex.abc = 2,
  col = NULL,
  bg = "#FAFAD2",
  radii = "rvdw",
  add = FALSE,
 windowRect = c(0, 0, 800, 600),
  FOV = 0,
  userMatrix = diag(4),
  . . .
)
## S3 method for class 'pdb'
visualize(
  х,
 mode = NULL,
  type = "1",
  xyz = NULL,
  abc = NULL,
  pbc.box = NULL,
  1wd = 2,
  lwd.xyz = lwd,
```

visualize

```
lwd.abc = lwd,
  lwd.pbc.box = lwd,
  cex.xyz = 2,
  cex.abc = 2,
  col = NULL,
 bg = "#FAFAD2",
  radii = "rvdw",
 add = FALSE,
 windowRect = c(0, 0, 800, 600),
 FOV = 0,
  userMatrix = diag(4),
  . . .
)
## S3 method for class 'character'
visualize(
 х,
 mode = NULL,
  type = "1",
  xyz = NULL,
  abc = NULL,
  pbc.box = NULL,
  1wd = 2,
  lwd.xyz = lwd,
  lwd.abc = lwd,
  lwd.pbc.box = lwd,
  cex.xyz = 2,
  cex.abc = 2,
  col = NULL,
  bg = "#FAFAD2",
  radii = "rvdw",
  add = FALSE,
 windowRect = c(0, 0, 800, 600),
 FOV = 0,
  userMatrix = diag(4),
  . . .
)
```

# Arguments

	further arguments passed to or from other methods.
x	an object or the name of a PDB file containing the molecular structure to visualize.
elename	a character vector containing the atomic names used to chose atom colors and radii.
cryst1	an object of class 'cryst1'. See cryst1
conect	an object of class 'conect'. See conect

mode	a single element character vector indicating the visualization mode (See details).
type	a character string indicating the visualization style (See details).
xyz	a logical value indicating whether the x, y and z axes have to be added to the scene. See details
abc	a logical value indicating whether the a, b and c axes have to be added to the scene. See details
pbc.box	a logical value indicating whether the pbc box has to be added to the scene. See details
lwd	a numeric value indication the line width used to plot the axes, the pbc box and atomic bonds when type = "1" (see details).
lwd.xyz	a numeric value indicating the line width used to plot the x, y and z axes.
lwd.abc	a numeric value indicating the line width used to plot the a, b and c axes.
lwd.pbc.box	a numeric value indicating the line width used to plot the pbc box.
cex.xyz	a numeric value indicating the magnification used to plot the labels of the x, y and z axes.
cex.abc	a numeric value indicating the magnification used to plot the labels of the a, b and c axes.
col	a vector indicating the colors to use to plot each atom.
bg	the color of the background
radii	either a character string indicating the type of radii or a numeric vector specify- ing the radii of each atom to use to plot atoms as spheres (see details).
add	a logical value indicating whether the plot has be to added to a existing scene (see rgl.cur and open3d).
windowRect	a vector of four integers indicating the left, top, right and bottom of the displayed window in pixels (see par3d).
FOV	the field of view. This controls the degree of parallax in the perspective view (see par3d).
userMatrix	a 4 by 4 matrix describing user actions to display the scene (see par3d).

#### Details

Three different visualization styles are allowed.

- When type="p": Points are drawn at each atomic positions (very light visualization mode).
- When type="1": Lines are drawn between bonded atoms. The connectivity of the system has to be specifyed.
- When type="s": Spheres are drawn at each atomic positions (heavy visualization mode). The radii of the spheres are given by radii.
  - When radii="rcov": Covalent radii, taken from the elements data set, are used.
  - When radii="rvdw": Van der Waals radii, taken from the elements data set, are used.
  - When radii is a numeric vector: The numeric values are used to assign to each atom a radius. If length(radii) != natom(pdb) radii is recycled.

When xyz, abc or pbc.box are NULL, the axis or pbc box are are added depending if a 'cryst1' object can be found.

Two different interactive visualization modes are avalable:

- When mode="measure": bond lengths, angles and dihedrals can be measured by **right-clicing** on the atoms.
- When mode="info": atomic labels can be added to the scene by **right-clicing** on the atoms. The labels are as follow: "ResidResname:EleidElename"

When mode=NULL the interactive mode is disabled. To escape the interactive mode press the ESC key.

#### Value

Return (using invisible) a two-column data.frame containing the IDs and type indicators of the objects added to the scene.

#### See Also

addXYZ, addABC, addPBCBox, par3d, select3d, measure, info3d

# Examples

```
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
visualize(x, type = "1", mode = NULL)
visualize(x, type = "s", radii = "rcov", mode = NULL)
visualize(x, type = "s", radii = "rvdw", mode = NULL)
visualize(x, type = "p", mode = NULL)
visualize(subset(x, resid != 1), type = "1", mode = NULL)
visualize(subset(x, resid == 1), type = "s", add = TRUE, mode = NULL)</pre>
```

```
wrap
```

#### Wrap Atomic Coordinates

#### Description

Wraps atomic coordinates using periodic boundary conditions.

```
wrap(x, ...)
## S3 method for class 'coords'
wrap(x, cryst1 = NULL, factor = NULL, ...)
## S3 method for class 'atoms'
wrap(x, cryst1 = NULL, factor = NULL, ...)
## S3 method for class 'pdb'
wrap(x, cryst1 = x$cryst1, factor = NULL, ...)
```

#### write.pdb

#### Arguments

x	an R object containing atomic coordinates to be wrapped.
	further arguments passed to or from other methods.
cryst1	an object of class 'cryst1' containing periodic boundary conditions used for wrapping.
factor	a factor used to wrap the atoms by groups.

#### Details

The wrap function translates all atoms out of the unit cell back into the unit cell using periodic boundary conditions. To do so, the wrap function first converts Cartesian into fractional coordinates. Then atoms with fractional coordinates greater than 1 or lower than 0 are respectively translated by -1 or +1. Finally, if the original atomic coordinates were Cartesian coordinates their are reconverted into Cartesian coordinates.

## Value

Return an object of class 'pdb' with wrapped atomic coordinates.

#### See Also

coords, atoms, pdb, cryst1, centres.pdb, xyz2abc

#### Examples

```
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))</pre>
```

```
# Translation of the atoms along x-axis
x$atoms$x1 <- x$atoms$x1 + 10</pre>
```

```
# Wrapping the structure
y <- wrap(x)</pre>
```

write.pdb PDB File Writer

#### Description

Writes a Protein Data Bank (PDB) coordinate file from an object of class 'pdb'.

```
write.pdb(x, file = "Rpdb.pdb")
```

xyz2abc

#### Arguments

х	an object, or a list of objects, of class 'pdb'.
file	a single element character vector containing the name of the PDB file to be created.

# Details

All data stored in the 'pdb' object are written to a PDB file. A list of objects of class 'pdb' can be provided to write multiple MODEL data into a single file. In this case, each 'pdb' object of the list must have the same cryst1 and conect components.

To write only a subset of a 'pdb' object see function subset.pdb.

## Value

No return value, called for side effects.

## References

PDB format is described at: http://www.wwpdb.org/documentation/format33/v3.3.html

#### See Also

read.pdb, pdb, cryst1, atoms, conect, subset.pdb

#### Examples

```
# Read a PDB file included with the package
pdb <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
# Write the pdb object to file "Rpdb.pdb" in the current directory</pre>
```

# Cleanup
unlink("Rpdb.pdb")

write.pdb(pdb, file = "Rpdb.pdb")

xyz2abc

From Cartesian to Fractional Coordinates and Vice Versa

## Description

Converts Cartesian coordinates into fractional coordinates and vice versa.

# xyz2abc

## Usage

```
xyz2abc(...)
## S3 method for class 'coords'
xyz2abc(x, cryst1, ...)
## S3 method for class 'atoms'
xyz2abc(x, cryst1, ...)
## S3 method for class 'pdb'
xyz2abc(x, cryst1 = x$cryst1, ...)
## S3 method for class 'distances'
xyz2abc(x, cryst1, ...)
abc2xyz(...)
## S3 method for class 'coords'
abc2xyz(x, cryst1, ...)
## S3 method for class 'atoms'
abc2xyz(x, cryst1, ...)
## S3 method for class 'pdb'
abc2xyz(x, cryst1 = x$cryst1, ...)
## S3 method for class 'distances'
abc2xyz(x, cryst1, ...)
```

# Arguments

	arguments passed to methods.
x	an R object containing atomic coordinates.
cryst1	an object of class cryst1.

# Details

For atoms and pdb objects, the atomic coordinates are first extracted from x using the coords function. Then, using the periodic boundary conditions stored into cryst1, the coordinates are converted from Cartesian to fractional (for the xyz2abc functions) or from fractional to Cartesian (for the abc2xyz functions) coordinates. Finally, for atoms and pdb objects, the new atomic coordinates are reassigned to the original x object using the coords<- function and x is returned.

# Value

Return an object of the same class as x, with atomic coordinates expressed in a different basis set.

# See Also

basis, coords, atoms, pdb, cryst1

# Examples

```
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
basis(x)
x <- xyz2abc(x)
basis(x)
x <- abc2xyz(x)
basis(x)</pre>
```

```
# This example returns an error because the coordinates stored
# into the PDB file are already Cartesian coordinates.
x <- read.pdb(system.file("examples/PCBM_ODCB.pdb", package="Rpdb"))
try(x <- abc2xyz(x))</pre>
```

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