## Package: RandomFieldsUtils (via r-universe)

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Version 1.2.5<br>Title Utilities for the Simulation and Analysis of Random Fields and Genetic Data<br>Author Martin Schlather [aut, cre], Alexander FreudenBerg [aut], Reinhard Furrer [ctb], Martin Kroll [ctb], Brian D. Ripley [ctb], John W. Ratcliff et al. (cph)<br>Maintainer Martin Schlather [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de)<br>Depends R (>=3.0)<br>Imports utils, methods, parallel<br>Suggests spam, RandomFields<br>Description Various utilities are provided that might be used in spatial statistics and elsewhere. It delivers a method for solving linear equations that checks the sparsity of the matrix before any algorithm is used.<br>Copyright MIT licence on sse2neon.H<br>License GPL (>= 3)<br>NeedsCompilation yes<br>Date/Publication 2022-04-19 11:32:32 UTC<br>Repository https://spatstat.r-universe.dev<br>RemoteUrl https://github.com/rubak/RandomFieldsUtils<br>RemoteRef HEAD<br>RemoteSha 8104bbc1fb4b83dfc49df5b8627d26a49539882b

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Cholesky Cholesky Decomposition of Positive Definite Matrices

## Description

This function calculates the Cholesky decomposition of a matrix.

## Usage

cholx (a)
chol2mv(C, n)
tcholRHS (C, RHS)

## Arguments

a
C
n
RHS
a square real-valued positive definite matrix
a (pivoted) Cholesky decomposition calculated by cholx
integer. Number of realisations of the multivariate normal distribution vector

## Details

If the matrix is diagonal direct calculations are performed.
Else the Cholesky decomposition is tried.

## Value

cholx returns a matrix containing the Cholesky decomposition (in its upper part).
chol2mv takes the Cholesky decomposition and returns a n realisations of a multivariate normal distribution with mean 0 and covariance function a
tcholRHS multiplies the vector RHS from the right to lower triangular matrix of the Cholesky decomposition. See examples below.

## Author(s)

Martin Schlather, [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de), https://www.wim.uni-mannheim.de/ schlather/

## References

Harbrecht, H., Peters, M., Schneider, R. (2012) On the low-rank approximation by the pivoted Cholesky decomposition. Appl. Num. Math. 62, 428-440.

## Examples

```
##########################
## Example showing the use of chol2mv and tcholRHS
n <- 10
M <- matrix(nc=n, runif(n^2))
M <- M %*% t(M) + diag(n)
C <- cholx(M)
set.seed(0)
v1 <- chol2mv(C, 1)
set.seed(0)
v2 <- tcholRHS(C, rnorm(n))
stopifnot(all(v1 == v2))
##########################
## The following example shows pivoted Cholesky can be used
## and the pivotation permutation can be transferred to
## subsequent Cholesky decompositions
set.seed(0)
n <- if (interactive()) 1000 else 100
x<- 1:n
y<- runif(n)
M<- x %*% t(x) + rev(x) %*% t(rev(x)) + y %*% t(y)
## do pivoting
RFoptions(pivot = PIVOT_DO, la_mode=LA_INTERN)
print(system.time(C <- cholx(M)))
print(range(crossprod(C) - M))
str(C)
## use the same pivoted decomposition as in the previous decomposition
M2 <- M + n * diag(1:n)
RFoptions(pivot = PIVOT_IDX, la_mode=LA_INTERN,
    pivot_idx = attr(C, "pivot_idx"),
    pivot_actual_size = attr(C, "pivot_actual_size"))
print(system.time(C2 <- cholx(M2)))
print(range(crossprod(C2) - M2))
range((crossprod(C2) - M2) / M2)
str(C2)
RFoptions(pivot = PIVOT_AUTO, la_mode = LA_AUTO)
```

```
confirm
Test if Two Objects are (Nearly) Equal
```


## Description

confirm( $x, y$ ) is a utility to compare $R$ objects $x$ and $y$ testing 'near equality' base on all.equal. It is written too allow different behaviour on different operating systems

## Usage

confirm(x, y, ...)

## Arguments

$x, y, \ldots$ see all. equal

## Value

Only TRUE or error in linux-gnu. Otherwise logical.

## Author(s)

Martin Schlather, [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de), https://www.wim.uni-mannheim.de/ schlather/

## Examples

$x<-3$
confirm(gauss(x), $\left.\exp \left(-x^{\wedge} 2\right)\right)$
dbinorm Density of a bivariate normal distribution

## Description

The function calculates the value of a bivariate normal distribution with mean 0 .

## Usage

dbinorm (x, S)

## Arguments

X
a matrix containing the $x$ values and the $y$ values in the first and second row respectively. Or it is a list of two vectors.
S the covariance matrix; currently only diagonal matrix possible

## Value

a vector according to the size of $x$

## Author(s)

Martin Schlather, [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de), https://www.wim.uni-mannheim.de/ schlather/

## Examples

```
x <- matrix(1:6, nc=2) + 0.0
C <- diag(c(1, 2))
dbinorm(x, C)
```

FileExists Files

## Description

The function FileExists checks whether a file or a lock-file exists
The function LockRemove removes a lock-file

## Usage

```
FileExists(file, printlevel=RFoptions()$basic$printlevel)
LockFile(file, printlevel=RFoptions()$basic$printlevel)
LockRemove(file)
WaitOthers(file, i, cores, ideal.processes=ceiling(cores * 1.25),
                max.processes=ceiling(cores * 1.5),
                distance=5, time=5, path="./")
```


## Arguments

| file | name of the data file |
| :--- | :--- |
| printlevel | if PrintLevel<=1 no messages are displayed <br> i |
| integer; current value of process, usually the number of a loop index |  |
| cores | the number of cores on the machine |
| ideal.processes, max. processes, distance |  |
| integer. See Details |  |
| time | in minutes a process waits until it rechecks its environment |
| path | the current path of file |

## Details

FileExists checks whether file or file.lock exists. If none of them exists file.lock is created and hostname and PID are written into file.lock. This is useful if several processes use the same directory. Further, it is checked whether another process has tried to create the same file in the same instance. In this case FileExists returns for at least one of the processes that file.lock has already been created.

LockFile is the same as FileExists except that it does not check whether file already exists.
WaitOthers waits for others if more than ideal. processes processes have their value is less than i or if more than cores processes have their value is less than i-distance. It also waits if there are alreay max. processes are active. Note that WaitOthers write a file with ending '.wait', which is also deleted be LockRemove.

## Value

FileExists returns

1 if file already exists
2 if file.lock already exists
3 if file.lock was tried to be created, but another process inferred and got priority
0 otherwise, file and file.lock did not exist and file.lock has been created

## Author(s)

Martin Schlather, [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de), https://www.wim.uni-mannheim.de/ schlather/

## Examples

```
## Not run:
## the next command checks whether the file 'data.rda'
## or the file 'data.rda.lock' exists. If so, a positive
## value is returned. If not, the file 'data.rda.lock'
## is created and the value 0 returned.
FileExists("data.rda")
## the next command deletes the file 'data.rda.lock'
LockRemove("data.rda")
## End(Not run)
```


## Description

gauss is a stationary isotropic covariance model. The corresponding covariance function only depends on the distance $r \geq 0$ between two points and is given by

$$
C(r)=e^{-r^{2}}
$$

## Usage

gauss(x, derivative=0)

## Arguments

| $x$ | numerical vector; for negative values the modulus is used |
| :--- | :--- |
| derivative | value in $0: 4$. |

## Value

If derivative=0, the function value is returned, otherwise the derivativeth derivative.
A vector of length( $x$ ) is returned; nu is recycled; scaling is recycled if numerical.

## Author(s)

Martin Schlather, [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de), https://www.wim.uni-mannheim.de/ schlather/

## References

Gelfand, A. E., Diggle, P., Fuentes, M. and Guttorp, P. (eds.) (2010) Handbook of Spatial Statistics. Boca Raton: Chapman \& Hall/CRL.

Stein, M. L. (1999) Interpolation of Spatial Data. New York: Springer-Verlag

## Examples

```
x <- 3
confirm(gauss(x), exp(-x^2))
```

host System calls

## Description

The functions hostname and pid return the host name and the PID, respectively.

## Usage

hostname()
pid()

## Details

If R runs on a unix platform the host name and the PID are returned, otherwise the empty string and naught, respectively.

## Value

| hostname | returns a string |
| :--- | :--- |
| pid | returns an unsigned integer |

## Author(s)

Martin Schlather, [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de), https://www.wim.uni-mannheim.de/ schlather/

## Examples

```
cat("The name of your computer is '", hostname(),
    "'. Your R program has current pid ", pid(), ".\n", sep="")
```

    Instruction Set CPU instruction set
    
## Description

The function checks whether a certain instruction is used (missed) under the current compilation of a package.

## Usage

uses.simd.instruction(which=NULL, pkgs=NULL)
misses.simd.instruction(which=NULL, pkgs=NULL)

## Arguments

$$
\begin{array}{ll}
\text { which character vector with values in "SSE2", "SSSE3", "AVX", "AVX2", "CUDA" } \\
\text { pkgs } & \text { character vector or missing. }
\end{array}
$$

## Value

logical vector of length which or matrix with number of rows equal to the length of which. An element is TRUE if the instruction set is used (missed) by the package.

If an arguments is NULL all available information is given.

## Author(s)

Martin Schlather, [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de), https://www.wim.uni-mannheim.de/ schlather/

## Examples

```
    uses.simd.instruction()
    misses.simd.instruction()
```

    Internal functions Internal functions
    
## Description

These functions are internal and should not be used.

## Usage

```
    checkExamples(exclude = NULL, include=1:length(.fct.list),
        ask=FALSE, echo=TRUE, halt=FALSE, ignore.all = FALSE,
        path=package, package = "RandomFields",
        read.rd.files=TRUE, local = FALSE, libpath \(=\) NULL,
        single.runs = FALSE, reset, catcherror=TRUE)
    Dependencies(pkgs = all.pkgs, dir = "Dependencies",
        install = FALSE, check=TRUE, reverse=FALSE,
        package="RandomFields")
    debugging_level()
```


## Arguments

exclude, include, ask, echo, halt, ignore.all, path, package, read.rd.files, local, libpath, single.runs, reset, catcherror
internal; ignore.all refers to the 'all' export statement in the namespace - whether this should be ignored. If read.rf.files is TRUE or a path to the Rd files, then the man pages are analysed to get all examples; ignore. all is then ignored. If FALSE only examples of functions (which are searched in the environments) are run.
pkgs, dir, install, check, reverse internal

## Author(s)

Martin Schlather, [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de), https://www.wim.uni-mannheim.de/ schlather/

## Examples

\#\# internal function: no examples given

## Description

matern calculates the Whittle-Matern covariance function (Soboloev kernel).
The Whittle model is given by

$$
C(r)=W_{\nu}(r)=2^{1-\nu} \Gamma(\nu)^{-1} r^{\nu} K_{\nu}(r)
$$

where $\nu>0$ and $K_{\nu}$ is the modified Bessel function of second kind.
The Matern model is given by

$$
C(r)=\frac{2^{1-\nu}}{\Gamma(\nu)}(\sqrt{2 \nu} r)^{\nu} K_{\nu}(\sqrt{2 \nu} r)
$$

The Handcock-Wallis parametrisation equals

$$
C(r)=\frac{2^{1-\nu}}{\Gamma(\nu)}(2 \sqrt{\nu} r)^{\nu} K_{\nu}(2 \sqrt{\nu} r)
$$

## Usage

```
whittle(x, nu, derivative=0,
        scaling=c("whittle", "matern", "handcockwallis"))
    matern(x, nu, derivative=0,
        scaling=c("matern", "whittle", "handcockwallis"))
```


## Arguments

X
nu
derivative
scaling
numerical vector; for negative values the modulus is used numerical vector with positive entries
value in $0: 4$.
numerical vector of positive values or character; see Details.

## Value

If derivative $=0$, the function value is returned, otherwise the derivativeth derivative.
A vector of length ( $x$ ) is returned; nu is recycled; scaling is recycled if numerical.
If scaling has a numerical values $s$, the covariance model equals

$$
C(r)=\frac{2^{1-\nu}}{\Gamma(\nu)}(s \sqrt{\nu} r)^{\nu} K_{\nu}(s \sqrt{\nu} r)
$$

The function values are rather precise even for large values of nu.

## Author(s)

Martin Schlather, [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de), https://www.wim.uni-mannheim.de/ schlather/

## References

Covariance function

- Chiles, J.-P. and Delfiner, P. (1999) Geostatistics. Modeling Spatial Uncertainty. New York: Wiley.
- Gelfand, A. E., Diggle, P., Fuentes, M. and Guttorp, P. (eds.) (2010) Handbook of Spatial Statistics. Boca Raton: Chapman \& Hall/CRL.
- Guttorp, P. and Gneiting, T. (2006) Studies in the history of probability and statistics. XLIX. On the Matern correlation family. Biometrika 93, 989-995.
- Handcock, M. S. and Wallis, J. R. (1994) An approach to statistical spatio-temporal modeling of meteorological fields. JASA 89, 368-378.
- Stein, M. L. (1999) Interpolation of Spatial Data - Some Theory for Kriging. New York: Springer.

See Also
nonstwm

## Examples

```
x <- 3
confirm(matern(x, 0.5), exp(-x))
confirm(matern(x, Inf), gauss(x/sqrt(2)))
confirm(matern(1:2, c(0.5, Inf)), exp(-(1:2)))
```

nonstwm nonstwm

## Description

The non-stationary Whittle-Matern model $C$ is given by

$$
C(x, y)=\Gamma(\mu) \Gamma(\nu(x))^{-1 / 2} \Gamma(\nu(y))^{-1 / 2} W_{\mu}(f(\mu)|x-y|)
$$

where $\mu=[\nu(x)+\nu(y)] / 2$, and $\nu$ must a positive function.
$W_{\mu}$ is the covariance function whittle.
The function $f$ takes the following values

$$
\begin{aligned}
& \text { scaling }=\text { "whittle" }: f(\mu)=1 \\
& \text { scaling }=\text { "matern" }: f(\mu)=\sqrt{2 \nu} \\
& \text { scaling }=\text { "handcockwallis" }: f(\mu)=2 \sqrt{\nu} \\
& \text { scaling }=\mathbf{s}, \text { numerical }: f(\mu)=s * \sqrt{n u}
\end{aligned}
$$

## Usage

nonstwm(x, y, nu, log=FALSE, scaling=c("whittle", "matern", "handcockwallis"))

## Arguments

$\mathrm{x}, \mathrm{y}$ nu positive value or a function with positive values and $x$ as argument log logical. If TRUE the logirithm of the covariance function is returned. scaling positive value or character; see Details.

## Value

A single value is returned.

## Author(s)

Martin Schlather, [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de), https://www.wim.uni-mannheim.de/ schlather/
orderx

## References

- Stein, M. (2005) Nonstationary Spatial Covariance Functions. Tech. Rep., 2005


## See Also

matern.

## Examples

```
nonstwm(2, 1, sin)
```


## orderx Ordering Permutation

## Description

orderx has the same functionality as order, except that $\operatorname{orderx}(\ldots$, from=from, to=to $)$ is the same as order[from:to]

## Usage

$\operatorname{prderx}(x$, from=1, to=length $(x)$, decreasing=FALSE, na.last $=N A)$

## Arguments

$x \quad$ an atomic vector
from, to $\quad \operatorname{order}(\ldots$, from=from, to=to) equals $\operatorname{order}(\ldots)$ [from:to]
decreasing logical. Should the sort order be increasing or decreasing?
na.last for controlling the treatment of NAs. If TRUE, missing values in the data are put last; if FALSE, they are put first; if NA, they are removed (see the Notes in order)

## Details

The smaller the difference to-from is compared to the length of $x$, the faster is order $x$ compared to order.
Particularly, orderx (. . . , from=k, to=k) is much faster than order (. . ) [k].
orderx is never really slower than order.
For further details see order.

## Value

integer vector of length to-from +1 .

## Author(s)

Martin Schlather, [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de), https://www.wim.uni-mannheim.de/ schlather/

## See Also

sortx

## Examples

```
x<- runif(10^6)
k <- 10
system.time(y<-order(x)[1:k])
system.time(z<-orderx(x, from=1, to=k)) ## much faster
stopifnot(all(x[y ]== x[z])) ## same result
```

Print Print method returning also the names automatically

## Description

prints variable names and the values

## Usage

Print(..., digits = 6, empty.lines = 2)

## Arguments

| $\ldots$. | any object that can be print-ed |
| :--- | :--- |
| digits | see print |
| empty.lines | number of leading empty lines |

## Value

prints the names and the values; for vectors cat is used and for lists str

## Author(s)

Martin Schlather, [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de), https://www.wim.uni-mannheim.de/ schlather/

## Examples

```
    a <- 4
    b <- list(c=5, g=7)
    m <- matrix(1:4, nc=2)
    Print(a, b, m)
```


## RFoptions Setting control arguments

## Description

RFoptions sets and returns control arguments for the analysis and the simulation of random fields

## Usage

```
RFoptions(..., no.class=FALSE, install.control=NULL)
```


## Arguments

```
... arguments in tag = value form, or a list of tagged values. See 'Details' for
    options in package RandomFieldsUtils.
no.class logical. If TRUE the list is returned without class specification.
install.control
    list. See Details, Part 2
```


## Details

The subsections below comment on

1. basic: Basic options
2. install.control
3. installNrun: Options for installation and running
4. solve: Options for solving linear systems
5. Reserved words

## 1. Basic options

asList logical. Lists of arguments are treated slightly different from non-lists. If asList=FALSE they are treated the same way as non-lists. This options being set to FALSE after calling RFoptions it should be set as first element of a list.
Default: TRUE
cores Number of cores for multicore algorithms; currently only used for the Cholesky decomposition.
Default : 1 if the package has been compiled with standard flags of CRAN and 0.75 * cores() $+0.25 *$ cpus() else.
Note that cores has not effect if set locally in this package or in package miraculix.
cPrintlevel cPrintlevel is automatically set to printlevel when printlevel is changed.
Standard users will never use a value higher than 3 .
0 : no messages
1 : messages and warnings when the user's input looks odd

2 : messages (and internal errors) documenting the choice of the simulation method
3 : further user relevant informations
4 : information on recursive function calls
5 : function flow information of central functions
6 : errors that are internally treated
7 : details on building up the covariance structure
8 : details on taking the square root of the covariance matrix
9 : details on intermediate calculations
10 : further details on intermediate calculations

Note that printlevel works on the R level whereas cPrintlevel works on the C level. cPrintlevel should be changed only globally.
Default: 1
efficient logical. If TRUE then always the most time efficient code is used.
Default: TRUE. It is strongly recommended to retain this value.
helpinfo logical. If TRUE then additional information is printed for more efficient programming in R.
Default: TRUE
printlevel If printlevel $\leq 0$ there is not any output on the screen. The higher the number the more tracing information is given. Standard users will never use a value higher than 3 .
0 : no messages
1 : important (error) messages and warnings
2 : less important messages
3 : details, but still for the user
4 : recursive call tracing
5 : function flow information of large functions
6 : errors that are internally treated
7 : details on intermediate calculations
8 : further details on intermediate calculations

## Default: 1

seed integer (currently only used by the package RandomFields). If NULL or NA set. seed is not called. Otherwise, set.seed(seed) is set before any simulations are performed.
If the argument is set locally, i.e., within a function, it has the usual local effect. If it is set globally, i.e. by RFoptions the seed is fixed for all subsequent calls.
If the number of simulations $n$ is greater than one and if RFoptions(seed=seed) is set, the $i$ th simulation is started with the seed 'seed $+i-1$ '.
skipchecks logical. If TRUE, several checks whether the given parameter values and the dimension are within the allowed range is skipped. Do not change the value of this variable except you really know what you do.
Default: FALSE
verbose logical. If FALSE it identical to printlevel $=1$ else to printlevel $=2$.
bigendian logical. Read only.
2. install.control: Details on argument install.control install.control may contain any argument of install. packages except type. This options is currently tailored for MS and Linux on Intel machines, only. The argument configure. args may not contain 'CXX_FLAGS' which should be passed as an extra argument with the list.
Note that if this argument is given (even with value NULL), an immediate installation takes place. In case the user tries to force to install 0 packages, an overview over the packages is given. If the user is asked whether re-installation shall take place, user can pass arguments to install.packages, e.g., "quiet=FALSE".
If install. control is given, no further argument may be passed to RFoptions.
Additional components of install. control and special behaviours are:
path the path to the locally saved tar balls
verbose, quiet They affect also the behaviour of RFoptions.
force TRUE reinstallation of all attached libraries based on and including RandomFieldsUtils. I.e., RFoptions(install. control=list(force=TRUE)) is the strongest form of forcing reinstallation.
FALSE In case some packages have to be re-installed the user will be asked.
not given reinstallation of the attached libraries based on and including RandomFieldsUtils that have not been tried yet in the current session.
pkgs=NULL brief overview over the installed packages based on RandomFieldsUtils
CROSS logical or character. CROSS is passed to 'configure.ac'.
"noflag" No extra compiler flag is set with respect to SIMD. This is the default.
TRUE each file is compiled with its specific SIMD/AVX compiler flags; this guarantees the compatiblity on a plattform with different sets of kernels. No SIND/AVX flag should be given by the user. Cross-compilation supported; no check is performed whether the code would run on the compiling CPU.
"nosimd" It is assumed that no SIMD is available and the flag "-no-sse2" is set (if possible).
"sse2" Same behaviour as TRUE, but all CPUs have at least "sse2" available.
"sse3", "ssse3", "sse41", "avx", "avx2" Alternatives to "sse2". Giving the highest guaranteed SIMD recognition leads to the most efficient code.
FALSE each file is compiled with all SIMD/AVX flags recognized by both the CPU and the compiler (no cross-compilation); users may add their own SIMD/AVX flags. This might lead to faster code, but which is not downwards compatible.
NA Same as FALSE except that the flag -mno-sse2 is set when no SIMD is needed or used.
This option can be set to "avx" interactively if install="ask".
CC_FLAGS character. Flags passed to 'configure. ac'.
SIMD_FLAGS character. A subset of "sse2", "sse3", "ssse3", "sse41", "avx", "avx2", "arch=xxx", etc. which will be tried instead of default flags. SIMD_FLAGS is passed to 'configure. ac'.
LOCAL_ONLY logical. If TRUE, the web is not searched for the latest version of the package.
MEM_IS_ALIGNED logical. If TRUE, then the memory is assumed to be aligned. If FALSE then the SIMD load commands _mm_*load_* are replaced by _mm_*loadu_*. If given, then force is set to TRUE.
USE_GPU logical. Force or hinder the compilation for the GPU

## 3. installNrun: Options for installing and for determining basic behaviour

install character. Only used by linux systems and alike including macOS The default by CRAN is that SIMD/AVX cannot be used at full extend. install determines what the action if the compiled version does not use the full CPU capacities. Since the use of GPU is heavily hardware dependent, its auto-recompilation is only performed in tow line of an AVX recompilation. The users usually use
"no" no re-installation
"ask" asks whether the library should be reinstalled, using the full capacity of the CPU according to the package.
"install" performs the auto-recompilation without asking. Note that only the command RFoptions(install.control=list(force=TRUE)) forces re-compilation of the currently loaded packages that are based on RandomFieldsUtils.
Note that, in each session, a package can be reinstalled only. This feature avoids trying to run jobs that cannot be done (e.g. 1 due to missing programs of the OS). See argument install. control for overwriting this behaviour.
Default: at starting point it is "ask" or "no", but the value may change during the session.
installPackages logical. Read only. Indicates whether packages are left to be re-installed. RFoptions(install="no") sets it to FALSE. RFoptions(install="no", install="ask") sets it to TRUE.
kahanCorrection obsolete. logical. If TRUE, the Kahan summation algorithm is used for calculating scalar products.
Default: false
la_mode determines
LA_AUTO, "auto" If a graphic card is recognized, LA_GPU is used. In all other cases the default is primarily LA_R. Only on linux systems, the package peforms a simple speed test and takes LA_INTERN if it is faster than LA_R; the time, hence the choice, depends also on the number of cores used.
LA_INTERN, "intern" mostly own algorithms, often based on SIMD/AVX. This option is of interest only if no advanced BLAS/LAPACK has been compiled into R
LA_R, "R" BLAS/LAPACK implementation used by R
LA_GPU, "GPU" This option is available when the package has been compiled with nvcc.
LA_QUERY, "query" Request on currently used set-up
Default: LA_AUTO
mem_is_alignedlogical. Read only. See MEM_IS_ALIGNED in install. control.
warn_parallel Logical. RandomFieldsUtils and packages using it, such as RandomFields and
miraculix, should now be prepared for parallelization using package parallel, for instance.
Internal OMP parallelization of RandomFieldsUtils is done, but only at a view points of the
subsequent packages.
As a few parts cannot be in parallel technically or from a logical point of view, a hint or a warning is given, if such a point is not accessed adequately. These messages can be turned off by warn_parallel = FALSE.
Default: TRUE.
warn_unknown_option integer.
$0,1,-1$ Unknown options are all ignored. If the value is positive, a hint is delivered whenever an unknown option is ignored.
-2,2 Unknown options that start with a capital letter are ignored. All others lead to an error. (Note that all RFoptions start with a minor letter.) If the value is positive, a hint is delivered whenever an unknown option is ignored.
3,-3 Unknown options that consists of a single capital letter are ignored. All others lead to an error. (Note that all RFoptions start with a minor letter.) If the value is positive, a hint is delivered whenever an unknown option is ignored.
4 (and other values) Any unknown option leads to an error.
Default for RandomFieldsUtils: 3
Default for RandomFields: 1

## 4. solve: Options for solving linear systems

det_as_log
eigen2zero When the svd or eigen decomposition is calculated, all values with modulus less than or equal to eigen2zero are set to zero.
Default: 1e-12
max_chol integer. Maximum number of rows of a matrix in a Cholesky decomposition
Default: 16384
max_svd integer. Maximum number of rows of a matrix in a svd decomposition
Default: 10000
pivot_partialdet logical. If TRUE then in case of low-rank matrices the determinant is calculated only in the part with positive eigenvalues
pivot Type of pivoting for the Cholesky decomposition. Possible values are
PIVOT_NONE, "no" No pivoting.
PIVOT_AUTO, "auto" If the matrix has a size greater than $3 \times 3$ and Choleskey fails without pivoting, privoting is done. For matrices of size less than $4 \times 4$, no pivoting and no checks are performed. See also PIVOT_DO
PIVOT_DO, "do" Do always pivoting. NOTE: privoted Cholesky decomposition yields only very approximately an upper triangular matrix $L$, but still $L^{\wedge} t L=M$ holds true.
PIVOT_IDX, "idx" uses the same pivoting as in the previous pivoted decomposition. This option becomes relevant only when simulations with different parameters or different models shall be performed with the same seed so that also the pivoting must be coupled.

Default: PIVOT_NONE
pivot_actual_size integer. Genuine dimension of the linear mapping given by a matrix in cholx. This is a very rarely used option when pivoting with pivot=PIVOT_IDX.
pivot_check logical. Only used in pivoted Cholesky decomposition. If TRUE and a numerically zero diagonal element is detected, it is checked whether the offdiagonal elements are numerically zero as well. (See also pivot_max_deviation and pivot_max_reldeviation.) If NA then only a warning is given.
Default: TRUE
pivot_idx vector of integer. Sequence of pivoting indices in pivoted Cholesky decomposition. Note that pivot_idx[1] gives the number of indices that will be used. The vector must have at least the length pivot_idx[1] +1 .
Default: NULL
pivot_relerror positive number. Tolerance for (numerically) negative eigenvalues and for (numerically) overdetermined systems appearing in the pivoted Cholesky decomposition.
Default: 1e-11
pivot_max_deviation positive number. Together with pivot_max_reldeviation it determines when the rest of the matrix (eigenvalues) in the pivoted Cholesky decomposition are considered as zero.
Default: 1e-10
pseudoinverse logical. In case of a singular matrix $M$, shall the pseudo inverse be returned for solvex(M)?
Default: FALSE
pivot_max_reldeviation positive number. Together with pivot_max_deviation it determines when the rest of the matrix (eigenvalues) in the pivoted Cholesky decomposition are considered as zero.
Default: 1e-10
solve_method vector of at most 3 integers that gives the sequence of methods in order to inverse a matrix or to calculate its square root: "cholesky", "svd", "lu", "eigen" "sparse", "method undefined". In the latter case, the algorithm decides which method might suit best.
Note that if use_spam is not false the algorithm checks whether a sparse matrix algorithm should be used and which is then tried first.
Default: "method undefined".
spam_factor integer. See argument spam_sample_n.
Default: 4294967
spam_min_n integer vector of size 2 . The minimal size for a matrix to apply a sparse matrix algorithms automatically. The second value is used in case the GPU is activated.
Default: c(400, 4000)
spam_min_p (spam_min_p) a numbers in ( 0,1 ) giving the proportion of zero above which an sparse matrix algorithm is used. The second value is used in case the GPU is activated.
Default: 0.8 (0.9)
spam_pivot integer. Pivoting algorithm for sparse matrices:
PIVOT_NONE No pivoting
PIVOTSPARSE_MMD
PIVOTSPARSE_RCM
See package spam for details.
Default: PIVOTSPARSE_MMD
spam_sample_n (spam_sample_n_GPU) Whether a matrix is sparse or not is tested by a 'random' sample of size spam_sample_n; The selection of the sample is iteratively obtained by multiplying the index by spam_factor modulo the size of the matrix.
Default: 500 (10000).
spam_tol largest absolute value being considered as zero. Default: DBL_EPSILON
svdtol Internal. When the svd decomposition is used for calculating the square root of a matrix then the absolute componentwise difference between this matrix and the square of the square root must be less than svdtol. No check is performed if svdtol is not positive.
Default: 0
use_spam Should the package spam (sparse matrices) be used for matrix calculations? If TRUE spam is always used. If FALSE, it is never used. If NA its use is determined by the size and the sparsity of the matrix.
Default: NA.

## 5. Reserved Words

list_ list_ usually equals the output of RFoptions(). This argument is used to reset the RFoptions. Some of the options behave differently if passed through list_. E.g. a warning counter is not reset. The argument list_ cannot be combined with any other arguments.
getoptions_ string vector of prefixes that indicate classes of options. In this package they can be "basic" and "solve". (E.g. package RandomFields has many more classes of options.) The given classes of options are then returned by RFoptions(). Note that the values are the previous values. getoptions_ must always be the very first argument.
saveoptions_ string vector of prefixes. Same as for getoptions_, except that important classes are always returned and thus should not be given. Hence saveoptions_ is often a convenient short cut for getoptions_. The class always included in this package is "basic", in package RandomFields these are the two classes "basic" and "general".
saveoptions_ must always be the very first argument. In particular, it may not given at the same time with getoptions_.
local_ logical. This options is allowed only when advanced packages are used, see RandomFields.
warnUnknown_ integer. Same as option warn_unknown_option, except that its value overwrites the value of warn_unknown_option in the current command RFoptions. This options must be placed between CODE and getoptions_, if the latter are used.

## Value

NULL if any argument is given, and the full list of arguments, otherwise.

## Author(s)

Martin Schlather, [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de), https://www.wim.uni-mannheim.de/ schlather/

## Examples

```
n <- 10
M <- matrix(1, ncol=n, nrow=n)
## Not run:
try(chol(M)) ## error, as M is not strictly positive definite
```

```
try(cholx(M)) ## also fails
## End(Not run)
RFoptions(la_mode=LA_INTERN, pivot=PIVOT_AUTO)
cholx(M) ## works
RFoptions(la_mode=LA_R)
RFoptions(solve_method="svd", pseudoinverse=TRUE)
solvex(M)
RFoptions(solve_method="method undefined", pseudoinverse=FALSE)
```

rowMeansx Some Further Row and Column Functions

## Description

The function rowMeansx returns weighted row means;
the function colMax returns column maxima;
the function rowProd returns the product of each row;
the function quadratic calculates a quadratic form
the function SelfDivByRow devides each column by a scalar;
the function dotXV calculates columnwise the dot product;
the function crossprodx calculates the cross product (using AVX);
the function scalarx calculates the scalar product (using AVX);

## Usage

```
rowMeansx(x, weight=NULL)
colMax(x)
rowProd(x)
SelfDivByRow(x, v)
quadratic(x, v)
dotXV(x, w)
crossprodx(x,y,mode=-1)
scalarx(x, y, mode=0)
```


## Arguments

| $x$ | numerical (or logical) matrix |
| :--- | :--- |
| $v$ | vector whose length equals the number of columns of $x$ |
| $w$ | vector whose length equals the number of rows of $x$ |
| weight | numerical or logical vector of length $\operatorname{nrow}(x)$ |

```
y numerical matrix
mode integer between 0 and 8 or negative, indicating that the default value should
    be used. Determine the algorithm how the scalar product is calculated. These
    values are experimental and may change their meaning.
```


## Details

quadratic $(\mathrm{x}, \mathrm{v})$ calculates the quadratic form $v^{\top} x v$; The matrix x must be squared.

## Value

rowMeansx returns a vector of lengthnrow(x).
colMax returns a vector of length ncol $(x)$.
rowProd returns a vector of length nrow ( $x$ ).
quadratic returns a scalar.
SelfDivByRow returns a matrix of same size as $x$.
$\operatorname{dot} \mathrm{XV}$ returns a matrix of same size as x .

## Author(s)

Martin Schlather, [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de), https://www.wim.uni-mannheim.de/ schlather/

## Examples

```
c <- if (interactive()) 10000 else 10
r <- if (interactive()) 20000 else 20
M <- matrix(nr=r, 1:(c * r))
## unweighted means, compare to rowMeans
print(system.time(m1 <- rowMeans(M)))
print(system.time(m2 <- rowMeansx(M)))
stopifnot(all.equal(m1, m2))
## weighted row means, compare to rowMeans
W <- 1 / (ncol(M) : 1)
print(system.time({M0 <- t(W * t(M)); m1 <- rowMeans(M0)}))
print(system.time(m2 <- rowMeansx(M, W)))
stopifnot(all.equal(m1, m2))
print(system.time(m1 <- apply(M, 2, max)))
print(system.time(m2 <- colMax(M)))
stopifnot(m1 == m2)
```

```
sleep.milli Sleep
```


## Description

Process sleeps for a given amount of time

## Usage

sleep.milli(n)
sleep.micro(n)

## Arguments

$\mathrm{n} \quad$ integer. sleeping time units

## Value

No value is returned.

## Author(s)

Martin Schlather, [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de), https://www.wim.uni-mannheim.de/ schlather/

## Examples

\#\# next command waits half a second before returning
sleep.milli(500)
solve Solve a System of Equations for Positive Definite Matrices

## Description

This function solves the equality $a x=b$ for $x$ where $a$ is a positive definite matrix and $b$ is a vector or a matrix. It is slightly faster than the inversion by the cholesky decomposition and clearly faster than solve. It also returns the logarithm of the determinant at no additional computational costs.

## Usage

solvex(a, b=NULL, logdeterminant=FALSE)

## Arguments

a
a square real-valued matrix containing the coefficients of the linear system. Logical matrices are coerced to numeric.
b a numeric or complex vector or matrix giving the right-hand side(s) of the linear system. If missing, $b$ is taken to be an identity matrix and solvex will return the inverse of a.
logdeterminant logical. whether the logarithm of the determinant should also be returned

## Details

If the matrix is diagonal direct calculations are performed.
Else if the matrix is sparse the package spam is used.
Else the Cholesky decomposition is tried. Note that with RFoptions(pivot=) pivoting can be enabled. Pivoting is about $30 \%$ slower.
If it fails, the eigen value decomposition is tried.

## Value

If logdeterminant=FALSE the function returns a vector or a matrix, depending on $b$ which is the solution to the linear equation. Else the function returns a list containing both the solution to the linear equation and the logarithm of the determinant of a.

## Author(s)

Martin Schlather, [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de), https://www.wim.uni-mannheim.de/ schlather/

## References

See chol.spam of the package spam.

## Examples

```
RFoptions(solve_method = "cholesky", printlevel=1)
set.seed(1)
n <- 1000
x <- 1:n
y <- runif(n)
## FIRST EXAMPLE: full rank matrix
M <- exp(-as.matrix(dist(x) / n))
b0 <- matrix(nr=n, runif(n * 5))
b <- M %*% b0 + runif(n)
## standard with 'solve'
print(system.time(z <- zR <- solve(M, b)))
print(range(b - M %*% z))
stopifnot(all(abs((b - M %*% z)) < 2e-11))
```

```
## using exactly the algorithm used in R
RFoptions(pivot=PIVOT_NONE, la_mode=LA_R) ## (default)
print(system.time(z <- solvex(M, b)))
print(range(b - M %*% z))
stopifnot(all(z == zR))
## Without pivoting, internal code:
RFoptions(pivot=PIVOT_NONE, la_mode=LA_INTERN) ## (default)
print(system.time(z <- solvex(M, b)))
print(range(b - M %*% z))
stopifnot(all(abs((b - M %*% z)) < 2e-11))
## Pivoting is slower here:
RFoptions(pivot=PIVOT_DO, la_mode=LA_INTERN)
print(system.time(z <- solvex(M, b)))
print(range(b - M %*% z))
stopifnot(all(abs((b - M %*% z)) < 2e-11))
## SECOND EXAMPLE: low rank matrix
M<- x %*% t(x) + rev(x) %*% t(rev(x)) + y %*% t(y)
b1 <- M %*% b0
## Without pivoting, it does not work
RFoptions(pivot=PIVOT_NONE, la_mode=LA_R)
## Not run: try(solve(M, b1))
RFoptions(pivot=PIVOT_NONE, la_mode=LA_INTERN)
## Not run: try(solvex(M, b1))
## Pivoting works -- the precision however is reduced :
RFoptions(pivot=PIVOT_DO, la_mode=LA_INTERN)
print(system.time(z1 <- solvex(M, b1)))
print(range(b1 - M %*% z1))
stopifnot(all(abs((b1 - M %*% z1)) < 2e-6))
## Pivoting fails, when the equation system is not solvable:
b2 <- M + runif(n)
## Not run: try(solvex(M, b2))
RFoptions(pivot = PIVOT_AUTO, la_mode = LA_AUTO)
```

sortx Sorting Vectors

## Description

sortx has the same functionality as sort, except that sortx (. . . from=from, to=to) is the same as sort[from:to]

Sort a vector or factor into ascending or descending order.

## Usage

sortx(x, from=1, to=length(x), decreasing=FALSE, na.last = NA)

## Arguments

x an atomic vector
from, to sort(..., from=from, to=to) equals sort(...)[from:to]
decreasing logical. Should the sort sort be increasing or decreasing?
na.last for controlling the treatment of NAs. If TRUE, missing values in the data are put last; if FALSE, they are put first; if NA, they are removed (see the Notes in sort)

## Details

The smaller the difference to-from is compared to the length of $x$, the faster is sortx compared to sort.

Particularly, sortx (. . . from=k, to=k) is much faster than sort (. . ) [k].
For further details see sort.

## Value

vector of length to-from +1 .

## Author(s)

Martin Schlather, [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de)

## See Also

## orderx

## Examples

```
x <- runif(10^6)
k <- 10
system.time(y<-sort(x)[1:k])
system.time(z<-sortx(x, from=1, to=k)) ## much faster
stopifnot(all(y == z)) ## same result
```


## Description

These functions return the values of the modified Struve functions and related functions

## Usage

```
struveH(x, nu)
struveL(x, nu, expon.scaled=FALSE)
I0L0(x)
```


## Arguments

$x \quad$ non-negative numeric vector
nu numeric vector
expon.scaled logical; if TRUE, the results are exponentially scaled in order to avoid overflow or underflow respectively.

## Details

I0L0 returns besselI (nu=0). minus struveL(nu=0).

## Value

Numeric vector with the (scaled, if expon. scaled = TRUE) values of the corresponding function.
The length of the result is the maximum of the lengths of the arguments $x$ and nu. The two arguments are recycled to that length.

## Author(s)

Martin Schlather, [schlather@math.uni-mannheim.de](mailto:schlather@math.uni-mannheim.de), https://www.wim.uni-mannheim.de/ schlather/

## References

- MacLeod, A.J. (1993) Chebyshev expansions for modified Struve and related functions, Mathematics of Computation, 60, 735-747
- Abramowitz, M., and Stegun, I.A. (1984) Pocketbook of Mathematical Functions, Verlag Harry Deutsch


## See Also

besselI

## Examples

if (FALSE) \{
$x<-\operatorname{seq}(1,2,0.1)$
struveH (x, 0)
struveH(x, 1)
IOL0(x) - (besselI(x, nu=0) - struveL(x, 0))
besselI(x, nu=1) - struveL(x, 1) \#\# cf. Abramovitz \& Stegun, table 12.1
\}

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