

Package ‘pnd’

May 20, 2025

Type Package

Title Parallel Numerical Derivatives, Gradients, Jacobians, and Hessians of Arbitrary Accuracy Order

Version 0.1.0

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Description Numerical derivatives through finite-difference approximations can be calculated using the 'pnd' package with parallel capabilities and optimal step-size selection to improve accuracy. These functions facilitate efficient computation of derivatives, gradients, Jacobians, and Hessians, allowing for more evaluations to reduce the mathematical and machine errors. Designed for compatibility with the 'numDeriv' package, which has not received updates in several years, it introduces advanced features such as computing derivatives of arbitrary order, improving the accuracy of Hessian approximations by avoiding repeated differencing, and parallelising slow functions on Windows, Mac, and Linux.

License EUPL

Encoding UTF-8

URL <https://github.com/Fifis/pnd>

BugReports <https://github.com/Fifis/pnd/issues>

Depends R (>= 3.4.0)

Imports parallel, Rdpack

Suggests numDeriv, knitr, rmarkdown, testthat (>= 3.0.0)

RdMacros Rdpack

RoxygenNote 7.3.2

VignetteBuilder knitr

Config/testthat/edition 3

NeedsCompilation no

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Repository CRAN

Date/Publication 2025-05-20 07:40:02 UTC

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alignStrings	<i>Align printed output to the longest argument</i>
--------------	---

Description

Align printed output to the longest argument

Usage

```
alignStrings(x, names = NULL, pad = c("l", "c", "r"))
```

Arguments

x	A numeric vector or matrix to be aligned with a vector of column names.
names	Optional: if x does not have (column) names, a character vector of element or column names to be output first. Ignored if x is named. Numeric inputs are converted to character automatically.
pad	A single character: "l" for left padding (flush-right justification), "c" for centre, and "r" for right padding (flush-left justification).

Value

A character matrix with the first row of names and the rest aligned content

Examples

```
x <- structure(1:4, names = month.name[1:4])
print(alignStrings(x, names(x)), quote = FALSE)
print(alignStrings(x, names(x), pad = "c"), quote = FALSE) # Centring
print(alignStrings(x, names(x), pad = "r"), quote = FALSE) # Left alignment

x <- matrix(c(1, 2.3, 4.567, 8, 9, 0), nrow = 2, byrow = TRUE)
colnames(x) <- c("Andy", "Bradley", "Ci")
alignStrings(x, pad = "c")
```

checkCores	<i>Number of core checks and changes</i>
------------	--

Description

Number of core checks and changes

Usage

```
checkCores(cores = NULL)
```

Arguments

cores Integer specifying the number of CPU cores used for parallel computation. Recommended to be set to the number of physical cores on the machine minus one.

Value

An integer with the number of cores.

Examples

```
checkCores()
checkCores(2)
suppressWarnings(checkCores(1000))
```

checkDimensions

*Determine function dimensionality and vectorisation***Description**

Determine function dimensionality and vectorisation

Usage

```

checkDimensions(
  FUN,
  x,
  f0 = NULL,
  func = NULL,
  elementwise = NA,
  vectorised = NA,
  multivalued = NA,
  deriv.order = 1,
  acc.order = 2,
  side = 0,
  h = NULL,
  zero.tol = sqrt(.Machine$double.eps),
  cores = 1,
  preschedule = TRUE,
  cl = NULL,
  ...
)

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

Arguments

FUN	A function returning a numeric scalar or a vector whose derivatives are to be computed. If the function returns a vector, the output will be a Jacobian.
x	Numeric vector or scalar: the point(s) at which the derivative is estimated. FUN(x) must be finite.
f0	Optional numeric: if provided, used to determine the vectorisation type to save time. If FUN(x) must be evaluated (e.g. second derivatives), saves one evaluation.
func	For compatibility with numDeriv::grad() only. If instead of FUN, func is used, it will be reassigned to FUN with a warning.
elementwise	Logical: is the domain effectively 1D, i.e. is this a mapping $\mathbb{R} \mapsto \mathbb{R}$ or $\mathbb{R}^n \mapsto \mathbb{R}^n$. If NA, compares the output length of the input length.

vectorised	Logical: if TRUE, the function is assumed to be vectorised: it will accept a vector of parameters and return a vector of values of the same length. Use FALSE or "no" for functions that take vector arguments and return outputs of arbitrary length (for $\mathbb{R}^n \mapsto \mathbb{R}^k$ functions). If NA, checks the output length and assumes vectorisation if it matches the input length; this check is necessary and potentially slow.
multivalued	Logical: if TRUE, the function is assumed to return vectors longer than 1. Use FALSE for element-wise functions. If NA, attempts inferring it from the function output.
deriv.order	Integer or vector of integers indicating the desired derivative order, d^m/dx^m , for each element of x.
acc.order	Integer or vector of integers specifying the desired accuracy order for each element of x. The final error will be of the order $O(h^{\text{acc.order}})$.
side	Integer scalar or vector indicating the type of finite difference: 0 for central, 1 for forward, and -1 for backward differences. Central differences are recommended unless computational cost is prohibitive.
h	Numeric or character specifying the step size(s) for the numerical difference or a method of automatic step determination ("CR", "CRm", "DV", or "SW" to be used in gradstep()). The default value is described in ?GenD.
zero.tol	Small positive integer: if $\text{abs}(x) \geq \text{zero.tol}$, then, the automatically guessed step size is relative (x multiplied by the step), unless an auto-selection procedure is requested; otherwise, it is absolute.
cores	Integer specifying the number of CPU cores used for parallel computation. Recommended to be set to the number of physical cores on the machine minus one.
preschedule	Logical: if TRUE, disables pre-scheduling for mclapply() or enables load balancing with parLapplyLB(). Recommended for functions that take less than 0.1 s per evaluation.
cl	An optional user-supplied cluster object (created by makeCluster or similar functions). If not NULL, the code uses parLapply() (if preschedule is TRUE) or parLapplyLB() on that cluster on Windows, and mclapply (fork cluster) on everything else.
...	Ignored.

Details

The following combinations of parameters are allowed, suggesting specific input and output handling by other functions:

	elementwise	!elementwise
!multivalued, vectorised	FUN(xgrid)	(undefined)
!multivalued, !vectorised	[mc]lapply(xgrid, FUN)	[mc]lapply gradient
multivalued, vectorised	(undefined)	FUN(xgrid) Jacobian
multivalued, !vectorised	(undefined)	[mc]lapply Jacobian

Some combinations are impossible: multi-valued functions cannot be element-wise, and single-valued vectorised functions must element-wise.

In brief, testing the input and output length and vectorisation capabilities may result in five cases, unlike 3 in `numDeriv::grad()` that does not provide checks for Jacobians.

Value

A named logical vector indicating if a function is element-wise or not, vectorised or not, and multivalued or not.

Examples

```
checkDimensions(sin, x = 1:4, h = 1e-5) # Rn -> Rn vectorised
checkDimensions(function(x) integrate(sin, 0, x)$value, x = 1:4, h = 1e-5) # non vec
checkDimensions(function(x) sum(sin(x)), x = 1:4, h = 1e-5) # Rn -> R, gradient
checkDimensions(function(x) c(sin(x), cos(x)), x = 1, h = 1e-5) # R -> Rn, Jacobian
checkDimensions(function(x) c(sin(x), cos(x)), x = 1:4, h = 1e-5) # vec Jac
checkDimensions(function(x) c(integrate(sin, 0, x)$value, integrate(sin, -x, 0)$value),
  x = 1:4, h = 1e-5) # non-vectorised Jacobian
```

dupRowInds

Repeated indices of the first unique value

Description

Repeated indices of the first unique value

Usage

```
dupRowInds(m)
```

Arguments

m A matrix or a data frame.

This function is an inverse function to such operations as `m[c(1:3, 1, 1, 2),]`: the matrix with potentially duplicated rows is taken as input, and repeated indices of the first occurrence of each row are returned.

This function is faster – at least in the examples tested so far – than `match(data.frame(t(m)), data.frame(t(unique(m))))`.

Value

A vector of row indices corresponding to the first occurrence of a given row.

Examples

```
dupRowInds(mtcars[rep(1:10, 10), rep(1:10, 10)])
dupRowInds(matrix(rnorm(1000), ncol = 10))
```

fdCoef

*Finite-difference coefficients for arbitrary grids***Description**

This function computes the coefficients for a numerical approximation to derivatives of any specified order. It provides the minimally sufficient stencil for the chosen derivative order and desired accuracy order. It can also use any user-supplied stencil (uniform or non-uniform). For that stencil $\{b_i\}_{i=1}^n$, it computes the optimal weights $\{w_i\}$ that yield the numerical approximation of the derivative:

$$\frac{d^m f}{dx^m} \approx h^{-m} \sum_{i=1}^n w_i f(x + b_i \cdot h)$$

Usage

```
fdCoef(
  deriv.order = 1L,
  side = c(0L, 1L, -1L),
  acc.order = 2L,
  stencil = NULL,
  zero.action = c("drop", "round", "none"),
  zero.tol = NULL
)
```

Arguments

deriv.order	Order of the derivative (m in $\frac{d^m f}{dx^m}$) for which a numerical approximation is needed.
side	Integer that determines the type of finite-difference scheme: 0 for central (AKA symmetrical or two-sided; the default), 1 for forward, and -1 for backward. Using 2 (for 'two-sided') triggers a warning and is treated as 0. with a warning. Unless the function is computationally prohibitively, central differences are strongly recommended for their accuracy.
acc.order	Order of accuracy: defines how the approximation error scales with the step size h , specifically $O(h^{a+1})$, where a is the accuracy order and depends on the higher-order derivatives of the function.
stencil	Optional custom vector of points for function evaluation. Must include at least $m+1$ points for the m -th order derivative.
zero.action	Character string specifying how to handle near-zero weights: "drop" to omit small (less in absolute value than zero.tol times the median weight) weights and corresponding stencil points, "round" to round small weights to zero, and "none" to leave all weights as calculated. E.g. the stencil for $f'(x)$ is $(-1, 0, 1)$ with weights $(-0.5, 0, 0.5)$; using "drop" eliminates the zero weight, and the redundant $f(x)$ is not computed.
zero.tol	Non-negative scalar defining the threshold: weights below zero.tol times the median weight are considered near-zero.

Details

This function relies on the approach of approximating numerical derivatives by weighted sums of function values described in (Fornberg 1988). It reproduces all tables from this paper exactly; see the example below to create Table 1.

The finite-difference coefficients for any given stencil are given as a solution of a linear system. The capabilities of this function are similar to those of (Taylor 2016), but instead of matrix inversion, the (Björck and Pereyra 1970) algorithm is used because the left-hand-side matrix is a Vandermonde matrix, and its inverse may be very inaccurate, especially for long one-sided stencils.

The weights computed for the stencil via this algorithm are very reliable; numerical simulations in (Higham 1987) show that the relative error is low even for ill-conditioned systems. (Kostyrka 2025) computes the exact relative error of the weights on the stencils returned by this function; the zero tolerance is based on these calculations.

Value

A list containing the stencil used and the corresponding weights for each point.

References

Björck Å, Pereyra V (1970). “Solution of Vandermonde systems of equations.” *Mathematics of computation*, **24**(112), 893–903.

Fornberg B (1988). “Generation of Finite Difference Formulas on Arbitrarily Spaced Grids.” *Mathematics of Computation*, **51**(184), 699–706. doi:10.1090/S00255718198809350770.

Higham NJ (1987). “Error analysis of the Björck-Pereyra algorithms for solving Vandermonde systems.” *Numerische Mathematik*, **50**(5), 613–632.

Kostyrka AV (2025). “What are you doing, step size: fast computation of accurate numerical derivatives with finite precision.” Working paper.

Taylor CR (2016). “Finite Difference Coefficients Calculator.” <https://web.media.mit.edu/~crtaylor/calculator.html>.

Examples

```
fdCoef() # Simple two-sided derivative
fdCoef(2) # Simple two-sided second derivative
fdCoef(acc.order = 4)$weights * 12 # Should be (1, -8, 8, -1)

# Using an custom stencil for the first derivative: x-2h and x+h
fdCoef(stencil = c(-2, 1), acc.order = 1)

# Reproducing Table 1 from Fornberg (1988) (cited above)
pad9 <- function(x) {l <- length(x); c(a <- rep(0, (9-l)/2), x, a)}
f <- function(d, a) pad9(fdCoef(deriv.order = d, acc.order = a,
                                zero.action = "round")$weights)
t11 <- t(sapply((1:4)*2, function(a) f(d = 1, a)))
t12 <- t(sapply((1:4)*2, function(a) f(d = 2, a)))
```



```

t13 <- t(sapply((1:3)*2, function(a) f(d = 3, a)))
t14 <- t(sapply((1:3)*2, function(a) f(d = 4, a)))
t11 <- cbind(t11[, 1:4], 0, t11[, 5:8])
t13 <- cbind(t13[, 1:4], 0, t13[, 5:8])
t1 <- data.frame(OrdDer = rep(1:4, times = c(4, 4, 3, 3)),
                 OrdAcc = c((1:4)*2, (1:4)*2, (1:3)*2, (1:3)*2),
                 rbind(t11, t12, t13, t14))
colnames(t1)[3:11] <- as.character(-4:4)
print(t1, digits = 4)

```

formatMat

*Round a matrix to N significant digits in mixed FP/exp notation***Description**

Round a matrix to N significant digits in mixed FP/exp notation

Usage

```
formatMat(x, digits = 3, shave.spaces = TRUE)
```

Arguments

x	Numeric matrix.
digits	Positive integer: the number of digits after the decimal comma to round to (i.e. one less than the number of significant digits).
shave.spaces	Logical: if true, removes spaces to ensure compact output; if false, results in nearly fixed-width output (almost).

Value

A numeric matrix with all entries of equal width with the same number of characters

Examples

```

x <- matrix(c(1234567, 12345.67, 123.4567,
              1.23456, -1.23456e-1, 0,
              -1.23456e-4, 1.23456e-2, -1.23456e-6), nrow = 3)
print(formatMat(x), quote = FALSE)
print(formatMat(x, digits = 1), quote = FALSE)

```

GenD

*Numerical derivative matrices with parallel capabilities***Description**

Computes numerical derivatives of a scalar or vector function using finite-difference methods. This function serves as a backbone for [Grad\(\)](#) and [Jacobian\(\)](#), allowing for detailed control over the derivative computation process, including order of derivatives, accuracy, and step size. GenD is fully vectorised over different coordinates of the function argument, allowing arbitrary accuracies, sides, and derivative orders for different coordinates.

Usage

```
GenD(
  FUN,
  x,
  elementwise = NA,
  vectorised = NA,
  multivalued = NA,
  deriv.order = 1L,
  side = 0,
  acc.order = 2L,
  stencil = NULL,
  h = NULL,
  zero.tol = sqrt(.Machine$double.eps),
  h0 = NULL,
  control = list(),
  f0 = NULL,
  cores = 1,
  preschedule = TRUE,
  cl = NULL,
  func = NULL,
  method = NULL,
  method.args = list(),
  ...
)

## S3 method for class 'GenD'
print(
  x,
  digits = 4,
  shave.spaces = TRUE,
  begin = "",
  sep = " ",
  end = "",
  ...
)
```

Arguments

<code>FUN</code>	A function returning a numeric scalar or a vector whose derivatives are to be computed. If the function returns a vector, the output will be a Jacobian.
<code>x</code>	Numeric vector or scalar: the point(s) at which the derivative is estimated. <code>FUN(x)</code> must be finite.
<code>elementwise</code>	Logical: is the domain effectively 1D, i.e. is this a mapping $\mathbb{R} \mapsto \mathbb{R}$ or $\mathbb{R}^n \mapsto \mathbb{R}^n$. If NA, compares the output length of the input length.
<code>vectorised</code>	Logical: if TRUE, the function is assumed to be vectorised: it will accept a vector of parameters and return a vector of values of the same length. Use FALSE or "no" for functions that take vector arguments and return outputs of arbitrary length (for $\mathbb{R}^n \mapsto \mathbb{R}^k$ functions). If NA, checks the output length and assumes vectorisation if it matches the input length; this check is necessary and potentially slow.
<code>multivalued</code>	Logical: if TRUE, the function is assumed to return vectors longer than 1. Use FALSE for element-wise functions. If NA, attempts inferring it from the function output.
<code>deriv.order</code>	Integer or vector of integers indicating the desired derivative order, d^m/dx^m , for each element of <code>x</code> .
<code>side</code>	Integer scalar or vector indicating the type of finite difference: 0 for central, 1 for forward, and -1 for backward differences. Central differences are recommended unless computational cost is prohibitive.
<code>acc.order</code>	Integer or vector of integers specifying the desired accuracy order for each element of <code>x</code> . The final error will be of the order $O(h^{\text{acc.order}})$.
<code>stencil</code>	Optional custom vector of points for function evaluation. Must include at least <code>m+1</code> points for the <code>m</code> -th order derivative.
<code>h</code>	Numeric or character specifying the step size(s) for the numerical difference or a method of automatic step determination ("CR", "CRm", "DV", or "SW" to be used in <code>gradstep()</code>). The default value is described in <code>?GenD</code> .
<code>zero.tol</code>	Small positive integer: if <code>abs(x) >= zero.tol</code> , then, the automatically guessed step size is relative (<code>x</code> multiplied by the step), unless an auto-selection procedure is requested; otherwise, it is absolute.
<code>h0</code>	Numeric scalar or vector: initial step size for automatic search with <code>gradstep()</code> .
<code>control</code>	A named list of tuning parameters passed to <code>gradstep()</code> .
<code>f0</code>	Optional numeric: if provided, used to determine the vectorisation type to save time. If <code>FUN(x)</code> must be evaluated (e.g. second derivatives), saves one evaluation.
<code>cores</code>	Integer specifying the number of CPU cores used for parallel computation. Recommended to be set to the number of physical cores on the machine minus one.
<code>preschedule</code>	Logical: if TRUE, disables pre-scheduling for <code>mclapply()</code> or enables load balancing with <code>parLapplyLB()</code> . Recommended for functions that take less than 0.1 s per evaluation.
<code>cl</code>	An optional user-supplied cluster object (created by <code>makeCluster</code> or similar functions). If not NULL, the code uses <code>parLapply()</code> (if <code>preschedule</code> is TRUE) or <code>parLapplyLB()</code> on that cluster on Windows, and <code>mclapply</code> (fork cluster) on everything else.

<code>func</code>	For compatibility with <code>numDeriv::grad()</code> only. If instead of <code>FUN</code> , <code>func</code> is used, it will be reassigned to <code>FUN</code> with a warning.
<code>method</code>	For compatibility with <code>numDeriv::grad()</code> only. Supported values: <code>"simple"</code> and <code>"Richardson"</code> . Non-null values result in a warning.
<code>method.args</code>	For compatibility with <code>numDeriv::grad()</code> only. Check <code>?numDeriv::grad</code> for a list of values. Non-empty lists result in a warning.
<code>...</code>	Ignored.
<code>digits</code>	Positive integer: the number of digits after the decimal comma to round to (i.e. one less than the number of significant digits).
<code>shave.spaces</code>	Logical: if true, removes spaces to ensure compact output; if false, results in nearly fixed-width output (almost).
<code>begin</code>	A character to put at the beginning of each line, usually <code>"", "("</code> , or <code>"c("</code> (the latter is useful if console output is used in calculations).
<code>sep</code>	The column delimiter, usually <code>" ", " ", "&"</code> (for LaTeX), or <code>", "</code> .
<code>end</code>	A character to put at the end of each line, usually <code>""</code> or <code>)"</code> .

Details

For computing gradients and Jacobians, use convenience wrappers `Jacobian` and `Grad`.

If the step size is too large, the slope of the secant poorly estimates the derivative; if it is too small, it leads to numerical instability due to the function value rounding.

The optimal step size for one-sided differences typically approaches $\text{Mach.eps}^{(1/2)}$ to balance the Taylor series truncation error with the rounding error due to storing function values with limited precision. For two-sided differences, it is proportional to $\text{Mach.eps}^{(1/3)}$. However, selecting the best step size typically requires knowledge of higher-order derivatives, which may not be readily available. Luckily, using `step = "SW"` invokes a reliable automatic data-driven step-size selection. Other options include `"DV"`, `"CR"`, and `"CRm"`. The step size defaults to $1.5\text{e-}8$ (the square root of machine epsilon) for x less than $1.5\text{e-}8$, $(2.2\text{e-}16)^{(1/(\text{deriv.order} + \text{acc.order}))} * x$ for $x > 1$, and interpolates exponentially between these values for $1.5\text{e-}8 < x < 1$.

The use of `f0` can reduce computation time similar to the use of `f.lower` and `f.upper` in `uniroot()`.

Unlike `numDeriv::grad()` and `numDeriv::jacobian()`, this function fully preserves the names of `x` and `FUN(x)`.

Value

A vector or matrix containing the computed derivatives, structured according to the dimensionality of `x` and `FUN`. If `FUN` is scalar-valued, returns a gradient vector. If `FUN` is vector-valued, returns a Jacobian matrix.

See Also

[gradstep\(\)](#) for automatic step-size selection.

Examples

```
# Case 1: Vector argument, vector output
f1 <- sin
g1 <- GenD(FUN = f1, x = 1:100)
g1.true <- cos(1:100)
plot(1:100, g1 - g1.true, main = "Approximation error of d/dx sin(x)")

# Case 2: Vector argument, scalar result
f2 <- function(x) sum(sin(x))
g2 <- GenD(FUN = f2, x = 1:4)
g2.h2 <- Grad(FUN = f2, x = 1:4, h = 7e-6)
g2 - g2.h2 # Tiny differences due to different step sizes
g2.auto <- Grad(FUN = f2, x = 1:4, h = "SW")
print(attr(g2.auto, "step.search")$exitcode) # Success

# Case 3: vector input, vector argument of different length
f3 <- function(x) c(sum(sin(x)), prod(cos(x)))
x3 <- 1:3
j3 <- GenD(f3, x3, multivalued = TRUE)
print(j3)

# Gradients for vectorised functions -- e.g. leaky ReLU
LReLU <- function(x) ifelse(x > 0, x, 0.01*x)
system.time(replicate(10, suppressMessages(GenD(LReLU, runif(30, -1, 1)))))
system.time(replicate(10, suppressMessages(GenD(LReLU, runif(30, -1, 1)))))

# Saving time for slow functions by using pre-computed values
x <- 1:4
finner <- function(x) sin(x*log(abs(x)+1))
fouter <- function(x) integrate(finner, 0, x, rel.tol = 1e-12, abs.tol = 0)$value
# The outer function is non-vectorised
fslow <- function(x) {Sys.sleep(0.01); fouter(x)}
f0 <- sapply(x, fouter)
system.time(GenD(fslow, x, side = 1, acc.order = 2, f0 = f0))
# Now, with extra checks, it will be slower
system.time(GenD(fslow, x, side = 1, acc.order = 2))
# Printing whilst preserving names
x <- structure(1:3, names = c("Andy", "Bradley", "Ca"))
print(Grad(function(x) prod(sin(x)), 1)) # 1D derivative
print(Grad(function(x) prod(sin(x)), x))
print(Jacobian(function(x) c(prod(sin(x)), sum(exp(x))), x))
```

generateGrid

Create a grid of points for a gradient / Jacobian

Description

Create a grid of points for a gradient / Jacobian

Usage

```
generateGrid(x, h, stencils, elementwise, vectorised)
```

Arguments

x	Numeric vector or scalar: the point(s) at which the derivative is estimated. FUN(x) must be finite.
h	Numeric or character specifying the step size(s) for the numerical difference or a method of automatic step determination ("CR", "CRm", "DV", or "SW" to be used in <code>gradstep()</code>). The default value is described in ?GenD.
stencils	A list of outputs from <code>fdCoef()</code> for each coordinate of x.
elementwise	Logical: is the domain effectively 1D, i.e. is this a mapping $\mathbb{R} \mapsto \mathbb{R}$ or $\mathbb{R}^n \mapsto \mathbb{R}^n$. If NA, compares the output length of the input length.
vectorised	Logical: if TRUE, the function is assumed to be vectorised: it will accept a vector of parameters and return a vector of values of the same length. Use FALSE or "no" for functions that take vector arguments and return outputs of arbitrary length (for $\mathbb{R}^n \mapsto \mathbb{R}^k$ functions). If NA, checks the output length and assumes vectorisation if it matches the input length; this check is necessary and potentially slow.

Value

A list with points for evaluation, summation weights for derivative computation, and indices for combining values.

Examples

```
generateGrid(1:4, h = 1e-5, elementwise = TRUE, vectorised = TRUE,
             stencils = lapply(1:4, function(a) fdCoef(acc.order = a)))
```

generateGrid2	<i>Generate grid points for Hessians</i>
---------------	--

Description

Creates a list of unique evaluation points for second derivatives: both diagonal ($\partial^2/\partial x_i^2$) and cross ($\partial^2/\partial x_i \partial x_j$).

Usage

```
generateGrid2(x, side, acc.order, h)
```

Arguments

<code>x</code>	Numeric vector or scalar: the point(s) at which the derivative is estimated. <code>FUN(x)</code> must be finite.
<code>side</code>	Integer scalar or vector indicating the type of finite difference: 0 for central, 1 for forward, and -1 for backward differences. Central differences are recommended unless computational cost is prohibitive.
<code>acc.order</code>	Integer or vector of integers specifying the desired accuracy order for each element of <code>x</code> . The final error will be of the order $O(h^{\text{acc.order}})$.
<code>h</code>	Numeric or character specifying the step size(s) for the numerical difference or a method of automatic step determination ("CR", "CRm", "DV", or "SW" to be used in <code>gradstep()</code>). The default value is described in <code>?GenD</code> .

Value

A list with elements:

- `xlist`: a list of unique coordinate shifts,
- `w`: the finite-difference weights (one per point),
- `i1`, `i2`: integer vectors giving partial-derivative indices.

The length of each vector matches `xlist`.

See Also

`GenD()`, `Hessian()`.

Examples

```
generateGrid2(1:4, side = rep(0, 4), acc.order = c(2, 6, 4, 2),
              h = c(1e-5, 1e-4, 2e-5, 1e-6))
```

Description

Computes numerical derivatives and gradients of scalar-valued functions using finite differences. This function supports both two-sided (central, symmetric) and one-sided (forward or backward) derivatives. It can utilise parallel processing to accelerate computation of gradients for slow functions or to attain higher accuracy faster.

Usage

```

Grad(
  FUN,
  x,
  elementwise = NA,
  vectorised = NA,
  multivalued = NA,
  deriv.order = 1L,
  side = 0,
  acc.order = 2,
  stencil = NULL,
  h = NULL,
  zero.tol = sqrt(.Machine$double.eps),
  h0 = NULL,
  control = list(),
  f0 = NULL,
  cores = 1,
  preschedule = TRUE,
  cl = NULL,
  func = NULL,
  method = NULL,
  method.args = list(),
  ...
)

```

Arguments

<code>FUN</code>	A function returning a numeric scalar or a vector whose derivatives are to be computed. If the function returns a vector, the output will be a Jacobian.
<code>x</code>	Numeric vector or scalar: the point(s) at which the derivative is estimated. <code>FUN(x)</code> must be finite.
<code>elementwise</code>	Logical: is the domain effectively 1D, i.e. is this a mapping $\mathbb{R} \mapsto \mathbb{R}$ or $\mathbb{R}^n \mapsto \mathbb{R}^n$. If NA, compares the output length of the input length.
<code>vectorised</code>	Logical: if TRUE, the function is assumed to be vectorised: it will accept a vector of parameters and return a vector of values of the same length. Use FALSE or "no" for functions that take vector arguments and return outputs of arbitrary length (for $\mathbb{R}^n \mapsto \mathbb{R}^k$ functions). If NA, checks the output length and assumes vectorisation if it matches the input length; this check is necessary and potentially slow.
<code>multivalued</code>	Logical: if TRUE, the function is assumed to return vectors longer than 1. Use FALSE for element-wise functions. If NA, attempts inferring it from the function output.
<code>deriv.order</code>	Integer or vector of integers indicating the desired derivative order, d^m/dx^m , for each element of <code>x</code> .
<code>side</code>	Integer scalar or vector indicating the type of finite difference: 0 for central, 1 for forward, and -1 for backward differences. Central differences are recommended unless computational cost is prohibitive.

<code>acc.order</code>	Integer or vector of integers specifying the desired accuracy order for each element of <code>x</code> . The final error will be of the order $O(h^{\text{acc.order}})$.
<code>stencil</code>	Optional custom vector of points for function evaluation. Must include at least $m+1$ points for the m -th order derivative.
<code>h</code>	Numeric or character specifying the step size(s) for the numerical difference or a method of automatic step determination ("CR", "CRm", "DV", or "SW" to be used in <code>gradstep()</code>). The default value is described in <code>?GenD</code> .
<code>zero.tol</code>	Small positive integer: if $\text{abs}(x) \geq \text{zero.tol}$, then, the automatically guessed step size is relative (x multiplied by the step), unless an auto-selection procedure is requested; otherwise, it is absolute.
<code>h0</code>	Numeric scalar or vector: initial step size for automatic search with <code>gradstep()</code> .
<code>control</code>	A named list of tuning parameters passed to <code>gradstep()</code> .
<code>f0</code>	Optional numeric: if provided, used to determine the vectorisation type to save time. If <code>FUN(x)</code> must be evaluated (e.g. second derivatives), saves one evaluation.
<code>cores</code>	Integer specifying the number of CPU cores used for parallel computation. Recommended to be set to the number of physical cores on the machine minus one.
<code>preschedule</code>	Logical: if TRUE, disables pre-scheduling for <code>mclapply()</code> or enables load balancing with <code>parLapplyLB()</code> . Recommended for functions that take less than 0.1 s per evaluation.
<code>cl</code>	An optional user-supplied cluster object (created by <code>makeCluster</code> or similar functions). If not NULL, the code uses <code>parLapply()</code> (if <code>preschedule</code> is TRUE) or <code>parLapplyLB()</code> on that cluster on Windows, and <code>mclapply</code> (fork cluster) on everything else.
<code>func</code>	For compatibility with <code>numDeriv::grad()</code> only. If instead of <code>FUN</code> , <code>func</code> is used, it will be reassigned to <code>FUN</code> with a warning.
<code>method</code>	For compatibility with <code>numDeriv::grad()</code> only. Supported values: "simple" and "Richardson". Non-null values result in a warning.
<code>method.args</code>	For compatibility with <code>numDeriv::grad()</code> only. Check <code>?numDeriv::grad</code> for a list of values. Non-empty lists result in a warning.
<code>...</code>	Ignored.

Details

This function aims to be 100% compatible with the syntax of `numDeriv::Grad()`, but there might be differences in the step size because some choices made in `numDeriv` are not consistent with theory.

There is one feature of the default step size in `numDeriv` that deserves an explanation. In that package (but not in `pnd`),

- If `method = "simple"`, then, simple forward differences are used with a fixed step size eps , which we denote by ε .
- If `method = "Richardson"`, then, central differences are used with a fixed step $h := |d \cdot x| + \varepsilon(|x| < \text{zero.tol})$, where $d = 1\text{e-}4$ is the relative step size and eps becomes an extra addition to the step size for the argument that are closer to zero than `zero.tol`.

We believe that the latter may lead to mistakes when the user believes that they can set the step size for near-zero arguments, whereas in reality, a combination of `d` and `eps` is used.

Here is the synopsis of the old arguments:

side `numDeriv` uses NA for handling two-sided differences. The `pnd` equivalent is 0, and NA is replaced with 0.

eps If `numDeriv method = "simple"`, then, `eps = 1e-4` is the absolute step size and forward differences are used. If `method = "Richardson"`, then, `eps = 1e-4` is the absolute increment of the step size for small arguments below the zero tolerance.

d If `numDeriv method = "Richardson"`, then, `d*abs(x)` is the step size for arguments above the zero tolerance and the baseline step size for small arguments that gets incremented by `eps`.

r The number of Richardson extrapolations that successively reduce the initial step size. For two-sided differences, each extrapolation increases the accuracy order by 2.

v The reduction factor in Richardson extrapolations.

Here are the differences in the new compatible implementation.

eps If `numDeriv method = "simple"`, then, `ifelse(x!=0, abs(x), 1) * sqrt(.Machine$double.eps) * 2` is used because one-sided differences require a smaller step size to reduce the truncation error. If `method = "Richardson"`, then, `eps = 1e-5`.

d If `numDeriv method = "Richardson"`, then, `d*abs(x)` is the step size for arguments above the zero tolerance and the baseline step size for small arguments that gets incremented by `eps`.

r The number of Richardson extrapolations that successively reduce the initial step size. For two-sided differences, each extrapolation increases the accuracy order by 2.

v The reduction factor in Richardson extrapolations.

`Grad` does an initial check (if `f0 = FUN(x)` is not provided) and calls `GenD()` with a set of appropriate parameters (`multivalued = FALSE` if the check succeeds). In case of parameter mismatch, throws an error.

Value

Numeric vector of the gradient. If `FUN` returns a vector, a warning is issued suggesting the use of `Jacobian()`.

See Also

`GenD()`, `Jacobian()`

Examples

```
f <- function(x) sum(sin(x))
g1 <- Grad(FUN = f, x = 1:4)
g2 <- Grad(FUN = f, x = 1:4, h = 7e-6)
g2 - g1 # Tiny differences due to different step sizes
g.auto <- Grad(FUN = f, x = 1:4, h = "SW")
print(g.auto)
attr(g.auto, "step.search")$exitcode # Success
```

```
# Gradients for vectorised functions -- e.g. leaky ReLU
LReLU <- function(x) ifelse(x > 0, x, 0.01*x)
Grad(LReLU, seq(-1, 1, 0.1))
```

gradstep

*Automatic step selection for numerical derivatives***Description**

Automatic step selection for numerical derivatives

Usage

```
gradstep(
  FUN,
  x,
  h0 = NULL,
  method = c("plugin", "SW", "CR", "CRm", "DV", "M", "K"),
  control = NULL,
  cores = 1,
  preschedule = getOption("pnd.preschedule", TRUE),
  cl = NULL,
  ...
)

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

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

Arguments

<code>FUN</code>	Function for which the optimal numerical derivative step size is needed.
<code>x</code>	Numeric vector or scalar: the point at which the derivative is computed and the optimal step size is estimated.
<code>h0</code>	Numeric vector or scalar: initial step size, defaulting to a relative step of slightly greater than $.Machine$double.eps^{(1/3)}$ (or absolute step if $x == 0$).
<code>method</code>	Character indicating the method: "CR" for (Curtis and Reid 1974), "CRm" for modified Curtis–Reid, "DV" for (Dumontet and Vignes 1977), "SW" (Stepleman and Winarsky 1979), "M" for (Mathur 2012), "K" for Kostyrka (2026, experimental), and "plugin" for the single-step plug-in estimator.
<code>control</code>	A named list of tuning parameters for the method. If NULL, default values are used. See the documentation for the respective methods. Note that full iteration history including all function evaluations is returned, but different methods have slightly different diagnostic outputs.

cores	Integer specifying the number of CPU cores used for parallel computation. Recommended to be set to the number of physical cores on the machine minus one.
preschedule	Logical: if TRUE, disables pre-scheduling for <code>mclapply()</code> or enables load balancing with <code>parLapplyLB()</code> . Recommended for functions that take less than 0.1 s per evaluation.
cl	An optional user-supplied cluster object (created by <code>makeCluster</code> or similar functions). If not NULL, the code uses <code>parLapply()</code> (if <code>preschedule</code> is TRUE) or <code>parLapplyLB()</code> on that cluster on Windows, and <code>mclapply</code> (fork cluster) on everything else.
...	Passed to FUN.

Details

We recommend using the Stepleman–Winarsky algorithm because it does not suffer from over-estimation of the truncation error in the Curtis–Reid approach and from sensitivity to near-zero third derivatives in the Dumontet–Vignes approach. It really tries multiple step sizes and handles missing values due to bad evaluations for inadequate step sizes really in a robust manner.

Value

A list similar to the one returned by `optim()` and made of concatenated individual elements coordinate-wise lists: `par` – the optimal step sizes found, `value` – the estimated numerical gradient, `counts` – the number of iterations for each coordinate, `abs.error` – an estimate of the total approximation error (sum of truncation and rounding errors), `exitcode` – an integer code indicating the termination status: 0 indicates optimal termination within tolerance, 1 means that the truncation error (CR method) or the third derivative (DV method) is zero and large step size is preferred, 2 is returned if there is no change in step size within tolerance, 3 indicates a solution at the boundary of the allowed value range, 4 signals that the maximum number of iterations was reached. `message` – summary messages of the exit status. `iterations` is a list of lists including the full step size search path, argument grids, function values on those grids, estimated error ratios, and estimated derivative values for each coordinate.

References

- Curtis AR, Reid JK (1974). “The Choice of Step Lengths When Using Differences to Approximate Jacobian Matrices.” *IMA Journal of Applied Mathematics*, **13**(1), 121–126. doi:10.1093/imamat/13.1.121.
- Dumontet J, Vignes J (1977). “Détermination du pas optimal dans le calcul des dérivées sur ordinateur.” *RAIRO. Analyse numérique*, **11**(1), 13–25. doi:10.1051/m2an/1977110100131.
- Mathur R (2012). *An Analytical Approach to Computing Step Sizes for Finite-Difference Derivatives*. Ph.D. thesis, University of Texas at Austin. <http://hdl.handle.net/2152/ETD-UT-2012-05-5275>.
- Stepleman RS, Winarsky ND (1979). “Adaptive numerical differentiation.” *Mathematics of Computation*, **33**(148), 1257–1264. doi:10.1090/s00255718197905379698.

See Also

[step.CR\(\)](#) for Curtis–Reid (1974) and its modification, [step.plugin\(\)](#) for the one-step plug-in solution, [step.DV\(\)](#) for Dumontet–Vignes (1977), [step.SW\(\)](#) for Stepleman–Winarsky (1979), [step.M\(\)](#) for Mathur (2012), and [step.K\(\)](#) for Kostyrka (2026).

Examples

```
gradstep(x = 1, FUN = sin, method = "CR")
gradstep(x = 1, FUN = sin, method = "CRm")
gradstep(x = 1, FUN = sin, method = "DV")
gradstep(x = 1, FUN = sin, method = "SW")
gradstep(x = 1, FUN = sin, method = "M")
# Works for gradients
gradstep(x = 1:4, FUN = function(x) sum(sin(x)))
print(step.CR(x = 1, sin))
print(step.DV(x = 1, sin))
print(step.plugin(x = 1, sin))
print(step.SW(x = 1, sin))
print(step.M(x = 1, sin))
print(step.K(x = 1, sin))
f <- function(x) x[1]^3 + sin(x[2])*exp(x[3])
print(gradstep(x = c(2, pi/4, 0.5), f))
```

Jacobian

Jacobian matrix computation with parallel capabilities s Computes the numerical Jacobian for vector-valued functions. Its columns are partial derivatives of the function with respect to the input elements. This function supports both two-sided (central, symmetric) and one-sided (forward or backward) derivatives. It can utilise parallel processing to accelerate computation of gradients for slow functions or to attain higher accuracy faster. Currently, only Mac and Linux are supported `parallel::mclapply()`. Windows support with `parallel::parLapply()` is under development.

Description

Jacobian matrix computation with parallel capabilities s Computes the numerical Jacobian for vector-valued functions. Its columns are partial derivatives of the function with respect to the input elements. This function supports both two-sided (central, symmetric) and one-sided (forward or backward) derivatives. It can utilise parallel processing to accelerate computation of gradients for slow functions or to attain higher accuracy faster. Currently, only Mac and Linux are supported `parallel::mclapply()`. Windows support with `parallel::parLapply()` is under development.

Usage

```
Jacobian(
  FUN,
  x,
```

```

    elementwise = NA,
    vectorised = NA,
    multivalued = NA,
    deriv.order = 1L,
    side = 0,
    acc.order = 2,
    stencil = NULL,
    h = NULL,
    zero.tol = sqrt(.Machine$double.eps),
    h0 = NULL,
    control = list(),
    f0 = NULL,
    cores = 1,
    preschedule = TRUE,
    cl = NULL,
    func = NULL,
    method = NULL,
    method.args = list(),
    ...
)

```

Arguments

<code>FUN</code>	A function returning a numeric scalar or a vector whose derivatives are to be computed. If the function returns a vector, the output will be a Jacobian.
<code>x</code>	Numeric vector or scalar: the point(s) at which the derivative is estimated. <code>FUN(x)</code> must be finite.
<code>elementwise</code>	Logical: is the domain effectively 1D, i.e. is this a mapping $\mathbb{R} \mapsto \mathbb{R}$ or $\mathbb{R}^n \mapsto \mathbb{R}^n$. If NA, compares the output length of the input length.
<code>vectorised</code>	Logical: if TRUE, the function is assumed to be vectorised: it will accept a vector of parameters and return a vector of values of the same length. Use FALSE or "no" for functions that take vector arguments and return outputs of arbitrary length (for $\mathbb{R}^n \mapsto \mathbb{R}^k$ functions). If NA, checks the output length and assumes vectorisation if it matches the input length; this check is necessary and potentially slow.
<code>multivalued</code>	Logical: if TRUE, the function is assumed to return vectors longer than 1. Use FALSE for element-wise functions. If NA, attempts inferring it from the function output.
<code>deriv.order</code>	Integer or vector of integers indicating the desired derivative order, d^m/dx^m , for each element of <code>x</code> .
<code>side</code>	Integer scalar or vector indicating the type of finite difference: 0 for central, 1 for forward, and -1 for backward differences. Central differences are recommended unless computational cost is prohibitive.
<code>acc.order</code>	Integer or vector of integers specifying the desired accuracy order for each element of <code>x</code> . The final error will be of the order $O(h^{\text{acc.order}})$.
<code>stencil</code>	Optional custom vector of points for function evaluation. Must include at least <code>m+1</code> points for the <code>m</code> -th order derivative.

<code>h</code>	Numeric or character specifying the step size(s) for the numerical difference or a method of automatic step determination ("CR", "CRm", "DV", or "SW" to be used in <code>gradstep()</code>). The default value is described in <code>?GenD</code> .
<code>zero.tol</code>	Small positive integer: if <code>abs(x) >= zero.tol</code> , then, the automatically guessed step size is relative (x multiplied by the step), unless an auto-selection procedure is requested; otherwise, it is absolute.
<code>h0</code>	Numeric scalar or vector: initial step size for automatic search with <code>gradstep()</code> .
<code>control</code>	A named list of tuning parameters passed to <code>gradstep()</code> .
<code>f0</code>	Optional numeric: if provided, used to determine the vectorisation type to save time. If <code>FUN(x)</code> must be evaluated (e.g. second derivatives), saves one evaluation.
<code>cores</code>	Integer specifying the number of CPU cores used for parallel computation. Recommended to be set to the number of physical cores on the machine minus one.
<code>preschedule</code>	Logical: if TRUE, disables pre-scheduling for <code>mclapply()</code> or enables load balancing with <code>parLapplyLB()</code> . Recommended for functions that take less than 0.1 s per evaluation.
<code>cl</code>	An optional user-supplied cluster object (created by <code>makeCluster</code> or similar functions). If not NULL, the code uses <code>parLapply()</code> (if <code>preschedule</code> is TRUE) or <code>parLapplyLB()</code> on that cluster on Windows, and <code>mclapply</code> (fork cluster) on everything else.
<code>func</code>	For compatibility with <code>numDeriv::grad()</code> only. If instead of <code>FUN</code> , <code>func</code> is used, it will be reassigned to <code>FUN</code> with a warning.
<code>method</code>	For compatibility with <code>numDeriv::grad()</code> only. Supported values: "simple" and "Richardson". Non-null values result in a warning.
<code>method.args</code>	For compatibility with <code>numDeriv::grad()</code> only. Check <code>?numDeriv::grad</code> for a list of values. Non-empty lists result in a warning.
<code>...</code>	Ignored.

Value

Matrix where each row corresponds to a function output and each column to an input coordinate. For scalar-valued functions, a warning is issued and the output is returned as a row matrix.

See Also

[GenD\(\)](#), [Grad\(\)](#)

Examples

```
slowFun <- function(x) {Sys.sleep(0.01); sum(sin(x))}
slowFunVec <- function(x) {Sys.sleep(0.01);
  c(sin = sum(sin(x)), exp = sum(exp(x)))}
true.g <- cos(1:4) # Analytical gradient
true.j <- rbind(cos(1:4), exp(1:4)) # Analytical Jacobian
x0 <- c(each = 1, par = 2, is = 3, named = 4)
```

```
# Compare computation times
system.time(g.slow <- numDeriv::grad(slowFun, x = x0) - true.g)
system.time(j.slow <- numDeriv::jacobian(slowFunVec, x = x0) - true.j)
system.time(g.fast <- Grad(slowFun, x = x0, cores = 2) - true.g)
system.time(j.fast <- Jacobian(slowFunVec, x = x0, cores = 2) - true.j)
system.time(j.fast4 <- Jacobian(slowFunVec, x = x0, acc.order = 4, cores = 2) - true.j)

# Compare accuracy
rownames(j.slow) <- paste0("numDeriv.jac.", c("sin", "exp"))
rownames(j.fast) <- paste0("pnd.jac.order2.", rownames(j.fast))
rownames(j.fast4) <- paste0("pnd.jac.order4.", rownames(j.fast4))
# Discrepancy
print(rbind(numDeriv.grad = g.slow, pnd.Grad = g.fast, j.slow, j.fast, j.fast4), 2)
# The order-4 derivative is more accurate for functions
# with non-zero third and higher derivatives -- look at pnd.jac.order.4
```

plotTE

Estimated total error plot

Description

Visualises the estimated truncation error, rounding error, and total error used in automatic step-size selection for numerical differentiation. The plot follows the approach used in Mathur (2012) and other step-selection methods.

Usage

```
plotTE(
  hgrid,
  etotal,
  eround,
  hopt = NULL,
  elabels = NULL,
  echeck = NULL,
  epsilon = .Machine$double.eps^(7/8),
  ...
)
```

Arguments

hgrid	Numeric vector: a sequence of step sizes used as the horizontal positions (usually exponentially spaced).
etotal	Numeric vector: estimated combined error at each step size. This is typically computed by subtracting a more accurate finite-difference approximation from a less accurate one.

eround	Numeric vector: estimated rounding error at each step size; usually the best guess or the upper bound is used.
hopt	Numeric scalar (optional): selected optimal step size. If provided, a vertical line is drawn at this value.
elabels	Character vector of the same length as hgrid containing the following values: "r", "g", "o", "i", "b" for r ounding, g ood truncation, o kay truncation, i ncreasing truncation and b ad truncation.
echeck	Numeric vector: estimated V-shaped check, usually from a fit.
epsilon	Numeric scalar: condition error, i.e. the error bound for the accuracy of the evaluated function; used for labelling rounding error assumptions.
...	Additional graphical parameters passed to plot().

Value

Nothing (invisible null).

Examples

```
a <- step.K(sin, 1)
hgrid <- a$iterations$h
etotal <- a$iterations$est.error[, 1]
eround <- a$iterations$est.error[, 2]
elabels <- c(rep("r", 32), rep("i", 2), rep("g", 12), rep("b", 15))
hopt <- a$par
plotTE(hgrid, etotal = 2e-12 * hgrid^2 + 1e-14 / hgrid,
       eround = 1e-14 / hgrid, hopt = 0.4, i.increasing = 30:45, i.good = 32:45,
       abline.round = c(-46.5, -1))
```

print.Hessian

Numerical Hessians

Description

Computes the second derivatives of a function with respect to all combinations of its input coordinates. Arbitrary accuracies and sides for different coordinates of the argument vector are supported.

Usage

```
## S3 method for class 'Hessian'
print(
  x,
  digits = 4,
  shave.spaces = TRUE,
  begin = "",
  sep = " ",
  end = "",
  ...
)
```

```

)

Hessian(
  FUN,
  x,
  side = 0,
  acc.order = 2,
  h = NULL,
  h0 = NULL,
  control = list(),
  f0 = NULL,
  cores = 1,
  preschedule = TRUE,
  cl = NULL,
  func = NULL,
  ...
)

```

Arguments

x	Numeric vector or scalar: point at which the derivative is estimated. FUN(x) must return a finite value.
digits	Positive integer: the number of digits after the decimal comma to round to (i.e. one less than the number of significant digits).
shave.spaces	Logical: if true, removes spaces to ensure compact output; if false, results in nearly fixed-width output (almost).
begin	A character to put at the beginning of each line, usually "", "(", or "c(" (the latter is useful if console output is used in calculations).
sep	The column delimiter, usually " ", " ", "&" (for LaTeX), or ", ".
end	A character to put at the end of each line, usually "" or ")".
...	Additional arguments passed to FUN.
FUN	A function returning a numeric scalar. If the function returns a vector, the output will be is a Jacobian. If instead of FUN, func is passed, as in numDeriv::grad, it will be reassigned to FUN with a warning.
side	Integer scalar or vector indicating difference type: 0 for central, 1 for forward, and -1 for backward differences. Central differences are recommended unless computational cost is prohibitive.
acc.order	Integer specifying the desired accuracy order. The error typically scales as $O(h^{\text{acc.order}})$.
h	Numeric scalar, vector, or character specifying the step size for the numerical difference. If character ("CR", "CRm", "DV", or "SW"), calls gradstep() with the appropriate step-selection method. Must be length 1 or match length(x). Matrices of step sizes are not supported. Suggestions how to handle all pairs of coordinates are welcome.
h0	Numeric scalar or vector: initial step size for automatic search with gradstep().Hessian(f, 1:100)

control	A named list of tuning parameters passed to gradstep().
f0	Optional numeric scalar or vector: if provided and applicable, used where the stencil contains zero (i.e. FUN(x) is part of the sum) to save time. TODO: Currently ignored.
cores	Integer specifying the number of CPU cores used for parallel computation. Recommended to be set to the number of physical cores on the machine minus one.
preschedule	Logical: if TRUE, disables pre-scheduling for mclapply() or enables load balancing with parLapplyLB(). Recommended for functions that take less than 0.1 s per evaluation.
cl	An optional user-supplied cluster object (created by makeCluster or similar functions). If not NULL, the code uses parLapply() (if preschedule is TRUE) or parLapplyLB() on that cluster on Windows, and mclapply (fork cluster) on everything else.
func	Deprecated; for numDeriv::grad() compatibility only.

Details

The optimal step size for 2nd-order-accurate central-differences-based Hessians is of the order $\text{Mach.eps}^{1/4}$ to balance the Taylor series truncation error with the rounding error. However, selecting the best step size typically requires knowledge of higher-order cross derivatives and is highly technically involved. Future releases will allow character arguments to invoke automatic data-driven step-size selection.

The use of `f0` can reduce computation time similar to the use of `f.lower` and `f.upper` in `uniroot()`.

Some numerical packages use the option (or even the default behaviour) of computing not only the $i < j$ cross-partials for the Hessian, but all pairs of i and j . The upper and lower triangular matrices are filled, and the matrix is averaged with its transpose to obtain a Hessian – this is the behaviour of `optimHess()`. However, it can be shown that $H[i, j]$ and $H[j, i]$ use the same evaluation grid, and with a single parallelisable evaluation of the function on that grid, no symmetrisation is necessary because the result is mathematically and computationally identical. In `pnd`, only the upper triangular matrix is computed, saving time and ensuring unambiguous results owing to the interchangeability of summation terms (ignoring the numerical error in summation as there is nothing that can be done apart from compensation summation, e.g. via Kahan's algorithm).

Value

A matrix with as many rows and columns as `length(x)`. Unlike the output of `numDeriv::hessian()`, this output preserves the names of `x`.

See Also

[Grad\(\)](#) for gradients, [GenD\(\)](#) for generalised numerical differences.

Examples

```
f <- function(x) prod(sin(x))
Hessian(f, 1:4)
# Large matrices
```

```
system.time(Hessian(f, 1:100))
```

printMat	<i>Print a matrix with separators</i>
----------	---------------------------------------

Description

Print a matrix with separators

Usage

```
printMat(
  x,
  digits = 3,
  shave.spaces = TRUE,
  begin = "",
  sep = " ",
  end = "",
  print = TRUE,
  format = TRUE
)
```

Arguments

x	A numeric matrix to print line by line.
digits	Positive integer: the number of digits after the decimal comma to round to (i.e. one less than the number of significant digits).
shave.spaces	Logical: if true, removes spaces to ensure compact output; if false, results in nearly fixed-width output (almost).
begin	A character to put at the beginning of each line, usually "", "(", or "c(" (the latter is useful if console output is used in calculations).
sep	The column delimiter, usually " ", " ", "&" (for LaTeX), or ", ".
end	A character to put at the end of each line, usually "" or ")".
print	If TRUE, outputs the lines of the matrix rows into the console.
format	If FALSE, skips the formatting part.

Value

The same x that was passed as the first input.

Examples

```
x <- matrix(c(-1234567, 12345.67, 123.4567,
              1.23456, -1.23456e-1, 0,
              1.23456e-4, 1.23456e-2, -1.23456e-6), nrow = 3)
printMat(x)
printMat(x, 2, TRUE, "c(", " ", " ", ")") # Ready row vectors
```

runParallel	<i>Run a function in parallel over a list (internal use only)</i>
-------------	---

Description

Run a function in parallel over a list (internal use only)

Usage

```
runParallel(FUN, x, cores = 1L, cl = NULL, preschedule = FALSE)
```

Arguments

<code>FUN</code>	A function of only one argument. If there are more arguments, use the <code>FUN2 <- do.call(FUN, c(list(x), ...))</code> and call it.
<code>x</code>	A list to parallelise the evaluation of <code>FUN</code> over: either numbers or expressions.
<code>cores</code>	Integer specifying the number of CPU cores used for parallel computation. Recommended to be set to the number of physical cores on the machine minus one.
<code>cl</code>	An optional user-supplied cluster object (created by <code>makeCluster</code> or similar functions). If not <code>NULL</code> , the code uses <code>parLapply()</code> (if <code>preschedule</code> is <code>TRUE</code>) or <code>parLapplyLB()</code> on that cluster on Windows, and <code>mclapply</code> (fork cluster) on everything else.
<code>preschedule</code>	Logical: if <code>TRUE</code> , disables pre-scheduling for <code>mclapply()</code> or enables load balancing with <code>parLapplyLB()</code> . Recommended for functions that take less than 0.1 s per evaluation.

Value

The value that `lapply(x, FUN)` would have returned.

Examples

```
fslow <- function(x) Sys.sleep(x)
x <- rep(0.05, 6)
cl <- parallel::makeCluster(2)
print(t1 <- system.time(runParallel(fslow, x)))
print(t2 <- system.time(runParallel(fslow, x, cl = cl)))
print(t3 <- system.time(runParallel(fslow, x, cores = 2)))
parallel::stopCluster(cl)
cat("Parallel overhead at 2 cores: ", round(t2[3]*200/t1[3]-100), "%\n", sep = "")
# Ignore on Windows
cat("makeCluster() overhead at 2 cores: ", round(100*t2[3]/t3[3]-100), "%\n", sep = "")
```

solveVandermonde	<i>Numerically stable non-confluent Vandermonde system solver</i>
------------------	---

Description

Numerically stable non-confluent Vandermonde system solver

Usage

```
solveVandermonde(s, b)
```

Arguments

- | | |
|---|--|
| s | Numeric vector of stencil points defining the Vandermonde matrix on the left-hand side, where each element $S_{i,j}$ is calculated as $s[j]^{(i-1)}$. |
| b | Numeric vector of the right-hand side of the equation. This vector must be the same length as s. |

Details

This function utilises the (Björck and Pereyra 1970) algorithm for an accurate solution to non-confluent Vandermonde systems, which are known for their numerical instability. Unlike Gaussian elimination, which suffers from ill conditioning, this algorithm achieves numerical stability through exploiting the ordering of the stencil. An unsorted stencils will trigger a warning. Additionally, the stencil must contain unique points, as repeated values make the Vandermonde matrix confluent and therefore non-invertible.

This implementation is a verbatim translation of Algorithm 4.6.2 from (Golub and Van Loan 2013), which is robust against the issues typically associated with Vandermonde systems.

See (Higham 1987) for an in-depth error analysis of this algorithm.

Value

A numeric vector of coefficients solving the Vandermonde system, matching the length of s.

References

- Björck Å, Pereyra V (1970). “Solution of Vandermonde systems of equations.” *Mathematics of computation*, **24**(112), 893–903.
- Golub GH, Van Loan CF (2013). *Matrix computations*, 4 edition. Johns Hopkins University Press.
- Higham NJ (1987). “Error analysis of the Björck-Pereyra algorithms for solving Vandermonde systems.” *Numerische Mathematik*, **50**(5), 613–632.

Examples

```
# Approximate the 4th derivatives on a non-negative stencil
solveVandermonde(s = 0:5, b = c(0, 0, 0, 0, 24, 0))

# Small numerical inaccuracies: note the 6.66e-15 in the 4th position --
# it should be rounded towards zero:
solveVandermonde(s = -3:3, b = c(0, 1, rep(0, 5))) * 60
```

step.CR

*Curtis–Reid automatic step selection***Description**

Curtis–Reid automatic step selection

Usage

```
step.CR(
  FUN,
  x,
  h0 = 1e-05 * max(abs(x), sqrt(.Machine$double.eps)),
  max.rel.error = .Machine$double.eps^(7/8),
  version = c("original", "modified"),
  aim = if (version[1] == "original") 100 else 1,
  acc.order = c(2L, 4L),
  tol = if (version[1] == "original") 10 else 4,
  range = h0/c(1e+05, 1e-05),
  maxit = 20L,
  seq.tol = 1e-04,
  cores = 1,
  preschedule = getOption("pnd.preschedule", TRUE),
  cl = NULL,
  ...
)
```

Arguments

<code>FUN</code>	Function for which the optimal numerical derivative step size is needed.
<code>x</code>	Numeric scalar: the point at which the derivative is computed and the optimal step size is estimated.
<code>h0</code>	Numeric scalar: initial step size, defaulting to a relative step of slightly greater than $\text{.Machine\$double.eps}^{1/3}$ (or absolute step if $x == 0$).
<code>max.rel.error</code>	Error bound for the relative function-evaluation error $(\frac{\hat{f}(x) - f(x)}{f(x)})$. Measures how noisy a function is. If the function is relying on numerical optimisation routines, consider setting to $\text{sqrt}(\text{.Machine\$double.eps})$. If the function has full precision to the last bit, set to $\text{.Machine\$double.eps}/2$.

version	Character scalar: "original" for the original 1974 version by Curtis and Reid; "modified" for Kostyrka's 2025 modification, which adds an extra evaluation for a more accurate estimate of the truncation error.
aim	Positive real scalar: desired ratio of truncation-to-rounding error. The "original" version over-estimates the truncation error, hence a higher aim is recommended. For the "modified" version, aim should be close to 1.
acc.order	Numeric scalar: in the modified version, allows searching for a step size that would be optimal for a 4th-order-accurate central difference See the Details section below.
tol	Numeric scalar greater than 1: tolerance multiplier for determining when to stop the algorithm based on the current estimate being between aim/tol and $\text{aim}*\text{tol}$.
range	Numeric vector of length 2 defining the valid search range for the step size.
maxit	Integer: maximum number of algorithm iterations to prevent infinite loops in degenerate cases.
seq.tol	Numeric scalar: maximum relative difference between old and new step sizes for declaring convergence.
cores	Integer specifying the number of CPU cores used for parallel computation. Recommended to be set to the number of physical cores on the machine minus one.
preschedule	Logical: if TRUE, disables pre-scheduling for <code>mclapply()</code> or enables load balancing with <code>parLapplyLB()</code> . Recommended for functions that take less than 0.1 s per evaluation.
cl	An optional user-supplied cluster object (created by <code>makeCluster</code> or similar functions). If not NULL, the code uses <code>parLapply()</code> (if <code>preschedule</code> is TRUE) or <code>parLapplyLB()</code> on that cluster on Windows, and <code>mclapply</code> (fork cluster) on everything else.
...	Passed to FUN.

Details

This function computes the optimal step size for central differences using the (Curtis and Reid 1974) algorithm. If the estimated third derivative is exactly zero, then, the initial step size is multiplied by 4 and returned.

If 4th-order accuracy (4OA) is requested, then, two things happen. Firstly, since 4OA differences requires a larger step size and the truncation error for the 2OA differences grows if the step size is larger than the optimal one, a higher ratio of truncation-to-rounding errors should be targeted. Secondly, a 4OA numerical derivative is returned, but the truncation and rounding errors are still estimated for the 2OA differences. Therefore, the estimating truncation error is higher and the real truncation error of 4OA differences is lower.

TODO: mention that `f` must be one-dimensional

The arguments passed to ... must not partially match those of `step.CR()`. For example, if `cl` exists, then, attempting to avoid cluster export by using `step.CR(f, x, h = 1e-4, cl = cl, a = a)` will result in an error: `a` matches `aim` and `acc.order`. Redefine the function for this argument to have a name that is not equal to the beginning of one of the arguments of `step.CR()`.

Value

A list similar to the one returned by `optim()`:

- `par` – the optimal step size found.
- `value` – the estimated numerical first derivative (using central differences; especially useful for computationally expensive functions).
- `counts` – the number of iterations (each iteration includes three function evaluations).
- `abs.error` – an estimate of the truncation and rounding errors.
- `exitcode` – an integer code indicating the termination status:
 - 0 – Optimal termination within tolerance.
 - 1 – Third derivative is zero; large step size preferred.
 - 2 – No change in step size within tolerance.
 - 3 – Solution lies at the boundary of the allowed value range.
 - 4 – Maximum number of iterations reached.
- `message` – A summary message of the exit status.
- `iterations` – A list including the full step size search path, argument grids, function values on those grids, estimated error ratios, and estimated derivative values.

References

Curtis AR, Reid JK (1974). “The Choice of Step Lengths When Using Differences to Approximate Jacobian Matrices.” *IMA Journal of Applied Mathematics*, **13**(1), 121–126. doi:10.1093/imamat/13.1.121.

Examples

```
f <- function(x) x^4
step.CR(x = 2, f)
step.CR(x = 2, f, h0 = 1e-3)
step.CR(x = 2, f, version = "modified")
step.CR(x = 2, f, version = "modified", acc.order = 4)

# A bad start: too far away
step.CR(x = 2, f, h0 = 1000) # Bad exit code + a suggestion to extend the range
step.CR(x = 2, f, h0 = 1000, range = c(1e-10, 1e5)) # Problem solved

library(parallel)
cl <- makePSOCKcluster(names = 2, outfile = "")
abc <- 2
f <- function(x, abc) {Sys.sleep(0.02); abc*sin(x)}
x <- pi/4
system.time(step.CR(f, x, h = 1e-4, cores = 1, abc = abc)) # To remove speed-ups
system.time(step.CR(f, x, h = 1e-4, cores = 2, abc = abc)) # Faster
f2 <- function(x) f(x, abc)
clusterExport(cl, c("f2", "f", "abc"))
system.time(step.CR(f2, x, h = 1e-4, cl = cl)) # Also fast
stopCluster(cl)
```

step.DV

*Dumontet–Vignes automatic step selection***Description**

Dumontet–Vignes automatic step selection

Usage

```

step.DV(
  FUN,
  x,
  h0 = 1e-05 * max(abs(x), sqrt(.Machine$double.eps)),
  range = h0/c(1e+06, 1e-06),
  max.rel.error = .Machine$double.eps^(7/8),
  ratio.limits = c(2, 15),
  maxit = 40L,
  cores = 1,
  preschedule = getOption("pnd.preschedule", TRUE),
  cl = NULL,
  ...
)

```

Arguments

FUN	Function for which the optimal numerical derivative step size is needed.
x	Numeric scalar: the point at which the derivative is computed and the optimal step size is estimated.
h0	Numeric scalar: initial step size, defaulting to a relative step of slightly greater than $\text{.Machine\$double.eps}^{1/3}$ (or absolute step if $x == 0$). This step size for first derivatives is internally translated into the initial step size for third derivatives by multiplying it by the machine epsilon raised to the power $-2/15$.
range	Numeric vector of length 2 defining the valid search range for the step size.
max.rel.error	Positive numeric scalar > 0 indicating the maximum relative error of function evaluation. For highly accurate functions with all accurate bits is equal to $\text{.Machine\$double.eps}/2$. For noisy functions (derivatives, integrals, output of optimisation routines etc.), it is higher, typically $\text{sqrt}(\text{.Machine\$double.eps})$. Dumontet and Vignes recommend $\text{.Machine\$double.eps}^{3/4} = 2\text{e-}12$ for common functions.
ratio.limits	Numeric vector of length 2 defining the acceptable ranges for step size: the algorithm stops if the relative perturbation of the third derivative by amplified rounding errors falls within this range.
maxit	Maximum number of algorithm iterations to avoid infinite loops in cases the desired relative perturbation factor cannot be achieved within the given range. Consider extending the range if this limit is reached.

cores	Integer specifying the number of CPU cores used for parallel computation. Recommended to be set to the number of physical cores on the machine minus one.
preschedule	Logical: if TRUE, disables pre-scheduling for <code>mclapply()</code> or enables load balancing with <code>parLapplyLB()</code> . Recommended for functions that take less than 0.1 s per evaluation.
cl	An optional user-supplied cluster object (created by <code>makeCluster</code> or similar functions). If not NULL, the code uses <code>parLapply()</code> (if <code>preschedule</code> is TRUE) or <code>parLapplyLB()</code> on that cluster on Windows, and <code>mclapply</code> (fork cluster) on everything else.
...	Passed to FUN.

Details

This function computes the optimal step size for central differences using the (Dumontet and Vignes 1977) algorithm. If the estimated third derivative is exactly zero, the function assumes a third derivative of 1 to prevent division-by-zero errors.

Note: the iteration history tracks the third derivative, not the first.

Value

A list similar to the one returned by `optim()`:

- `par` – the optimal step size found.
- `value` – the estimated numerical first derivative (using central differences).
- `counts` – the number of iterations (each iteration includes four function evaluations).
- `abs.error` – an estimate of the truncation and rounding errors.
- `exitcode` – an integer code indicating the termination status:
 - 0 – Optimal termination within tolerance.
 - 1 – Third derivative is zero; large step size preferred.
 - 3 – Solution lies at the boundary of the allowed value range.
 - 4 – Maximum number of iterations reached; optimal step size is within the allowed range.
 - 5 – Maximum number of iterations reached; optimal step size was outside allowed range and had to be snapped to a boundary.
 - 6 – No search was performed (used when `maxit` = 1).
- `message` – A summary message of the exit status.
- `iterations` – A list including the full step size search path (note: for the third derivative), argument grids, function values on those grids, and estimated third derivative values.

References

Dumontet J, Vignes J (1977). “Détermination du pas optimal dans le calcul des dérivées sur ordinateur.” *RAIRO. Analyse numérique*, **11**(1), 13–25. doi:[10.1051/m2an/1977110100131](https://doi.org/10.1051/m2an/1977110100131).

Examples

```
f <- function(x) x^4
step.DV(x = 2, f)
step.DV(x = 2, f, h0 = 1e-3)

# Plug-in estimator with only one evaluation of f'''
step.DV(x = 2, f, maxit = 1)
step.plugin(x = 2, f)
```

step.K

Kink-based step selection (experimental!)

Description

Optimal step-size search using the full range of practical error estimates and numerical optimisation to find the spot where the theoretical total-error shape is best described by the data, and finds the step size where the ratio of rounding-to-truncation error is optimal.

Usage

```
step.K(
  FUN,
  x,
  h0 = NULL,
  deriv.order = 1,
  acc.order = 2,
  range = NULL,
  shrink.factor = 0.5,
  max.rel.error = .Machine$double.eps^(7/8),
  plot = FALSE,
  cores = 1,
  preschedule = getOption("pnd.preschedule", TRUE),
  cl = NULL,
  ...
)
```

Arguments

FUN	Function for which the optimal numerical derivative step size is needed.
x	Numeric scalar: the point at which the derivative is computed and the optimal step size is estimated.
h0	Numeric scalar: initial step size, defaulting to a relative step of slightly greater than $.Machine$double.eps^{1/3}$ (or absolute step if $x == 0$).
deriv.order	Integer or vector of integers indicating the desired derivative order, d^m/dx^m , for each element of x.

acc.order	Integer or vector of integers specifying the desired accuracy order for each element of x . The final error will be of the order $O(h^{\text{acc.order}})$.
range	Numeric vector of length 2 defining the valid search range for the step size.
shrink.factor	A scalar less than 1 that is used to create a sequence of step sizes. The recommended value is 0.5. Change to 0.25 for a faster search. This number should be a negative power of 2 for the most accurate representation.
max.rel.error	Error bound for the relative function-evaluation error ($\frac{\hat{f}(\hat{x}) - f(x)}{f(x)}$). Measures how noisy a function is. If the function is relying on numerical optimisation routines, consider setting to <code>sqrt(.Machine\$double.eps)</code> . If the function has full precision to the last bit, set to <code>.Machine\$double.eps/2</code> .
plot	Logical: if TRUE, plots the estimated truncation and round-off errors.
cores	Integer specifying the number of CPU cores used for parallel computation. Recommended to be set to the number of physical cores on the machine minus one.
preschedule	Logical: if TRUE, disables pre-scheduling for <code>mclapply()</code> or enables load balancing with <code>parLapplyLB()</code> . Recommended for functions that take less than 0.1 s per evaluation.
cl	An optional user-supplied cluster object (created by <code>makeCluster</code> or similar functions). If not NULL, the code uses <code>parLapply()</code> (if <code>preschedule</code> is TRUE) or <code>parLapplyLB()</code> on that cluster on Windows, and <code>mclapply</code> (fork cluster) on everything else.
...	Passed to FUN.

Details

This function computes the optimal step size for central differences using the statistical kink-search approach. The optimal step size is determined as the minimiser of the total error, which for central differences is V-shaped with the left-branch slope equal to the negative derivation order and the right-branch slope equal to the accuracy order. For standard simple central differences, the slopes are -1 and 2, respectively. The algorithm uses the least-median-of-squares (LMS) penalty and searches for the optimal position of the check that fits the data the best on a bounded 2D rectangle using derivative-free (Nelder–Mead) optimisation. Unlike other algorithms, if the estimated third derivative is too small, the function shape will be different, and two checks are made for the existence of two branches.

Value

A list similar to the one returned by `optim()`:

- `par` – the optimal step size found.
- `value` – the estimated numerical first derivative (using central differences).
- `counts` – the number of iterations (each iteration includes two function evaluations).
- `abs.error` – an estimate of the truncation and rounding errors.
- `exitcode` – an integer code indicating the termination status:
 - 0 – Optimal termination; a minimum of the V-shaped function was found.
 - 1 – Third derivative is too small or noisy; a fail-safe value is returned.

- 2 – Third derivative is nearly zero; a fail-safe value is returned.
- 3 – There is no left branch of the V shape; a fail-safe value is returned.
- message – A summary message of the exit status.
- iterations – A list including the step and argument grids, function values on those grids, estimated derivative values, estimated error values, and predicted model-based errors.

References

There are no references for Rd macro `\insertAllCites` on this help page.

Examples

```
step.K(sin, 1, plot = TRUE)
step.K(exp, 1, range = c(1e-12, 1e-0), plot = TRUE)
step.K(atan, 1, plot = TRUE)

# Edge case 1: function symmetric around x0, zero truncation error
step.K(sin, pi/2, plot = TRUE)
step.K(sin, pi/2, shrink.factor = 0.8, plot = TRUE)

# Edge case 1: the truncation error is always zero and f(x0) = 0
suppressWarnings(step.K(function(x) x^2, 0, plot = TRUE))
# Edge case 2: the truncation error is always zero
step.K(function(x) x^2, 1, plot = TRUE)
step.K(function(x) x^4, 0, plot = TRUE)
step.K(function(x) x^4, 0.1, plot = TRUE)
step.K(function(x) x^6 - x^4, 0.1, plot = TRUE)
step.K(atan, 3/4, plot = TRUE)
step.K(exp, 2, plot = TRUE, range = c(1e-16, 1e+1))
```

step.M

Mathur's AutoDX-like automatic step selection

Description

Mathur's AutoDX-like automatic step selection

Usage

```
step.M(
  FUN,
  x,
  h0 = NULL,
  max.rel.error = .Machine$double.eps^(7/8),
  range = NULL,
  shrink.factor = 0.5,
  min.valid.slopes = 5L,
  seq.tol = 0.01,
```

```

correction = TRUE,
plot = FALSE,
cores = 1,
preschedule = getOption("pnd.preschedule", TRUE),
cl = NULL,
...
)

```

Arguments

<code>FUN</code>	Function for which the optimal numerical derivative step size is needed.
<code>x</code>	Numeric scalar: the point at which the derivative is computed and the optimal step size is estimated.
<code>h0</code>	Numeric scalar: initial step size, defaulting to a relative step of slightly greater than $.Machine$double.eps^{1/3}$ (or absolute step if <code>x == 0</code>).
<code>max.rel.error</code>	Error bound for the relative function-evaluation error ($\frac{\hat{f}(x) - f(x)}{f(x)}$). Measures how noisy a function is. If the function is relying on numerical optimisation routines, consider setting to <code>sqrt(.Machine\$double.eps)</code> . If the function has full precision to the last bit, set to <code>.Machine\$double.eps/2</code> .
<code>range</code>	Numeric vector of length 2 defining the valid search range for the step size.
<code>shrink.factor</code>	A scalar less than 1 that is used to create a sequence of step sizes. The recommended value is 0.5. Change to 0.25 for a faster search. This number should be a negative power of 2 for the most accurate representation.
<code>min.valid.slopes</code>	Positive integer: how many points must form a sequence with the correct slope with relative difference from 2 less than <code>seq.tol</code> . If <code>shrink.factor</code> is small (< 0.33), consider reducing this to 4.
<code>seq.tol</code>	Numeric scalar: maximum relative difference between old and new step sizes for declaring convergence.
<code>correction</code>	Logical: if TRUE, returns the corrected step size (last point in the sequence times a less-than-1 number to account for the possible continuation of the downwards slope of the total error); otherwise, returns the grid point that is lowest in the increasing sequence of valid error estimates.
<code>plot</code>	Logical: if TRUE, plots the estimated truncation and round-off errors.
<code>cores</code>	Integer specifying the number of CPU cores used for parallel computation. Recommended to be set to the number of physical cores on the machine minus one.
<code>preschedule</code>	Logical: if TRUE, disables pre-scheduling for <code>mclapply()</code> or enables load balancing with <code>parLapplyLB()</code> . Recommended for functions that take less than 0.1 s per evaluation.
<code>cl</code>	An optional user-supplied cluster object (created by <code>makeCluster</code> or similar functions). If not NULL, the code uses <code>parLapply()</code> (if <code>preschedule</code> is TRUE) or <code>parLapplyLB()</code> on that cluster on Windows, and <code>mclapply</code> (fork cluster) on everything else.
<code>...</code>	Passed to <code>FUN</code> .

Details

This function computes the optimal step size for central differences using the (Mathur 2012) algorithm.

Value

A list similar to the one returned by `optim()`:

- `par` – the optimal step size found.
- `value` – the estimated numerical first derivative (using central differences).
- `counts` – the number of iterations (each iteration includes two function evaluations).
- `abs.error` – an estimate of the truncation and rounding errors.
- `exitcode` – an integer code indicating the termination status:
 - 0 – Optimal termination due to a sequence of correct reductions.
 - 1 – Reductions are slightly outside the tolerance.
 - 2 – Tolerances are significantly violated; an approximate minimum is returned.
 - 3 – Not enough finite function values; a rule-of-thumb value is returned.
- `message` – A summary message of the exit status.
- `iterations` – A list including the step and argument grids, function values on those grids, estimated derivative values, and estimated error values.

References

Mathur R (2012). *An Analytical Approach to Computing Step Sizes for Finite-Difference Derivatives*. Ph.D. thesis, University of Texas at Austin. <http://hdl.handle.net/2152/ETD-UT-2012-05-5275>.

Examples

```
f <- function(x) x^4 # The derivative at 1 is 4
step.M(x = 1, f, plot = TRUE)
step.M(x = 1, f, h0 = 1e-9) # Starting low
step.M(x = 1, f, h0 = 1000) # Starting high

f <- sin # The derivative at pi/4 is sqrt(2)/2
step.M(x = pi/2, f, plot = TRUE) # Bad case -- TODO a fix
step.M(x = pi/4, f, plot = TRUE)
step.M(x = pi/4, f, h0 = 1e-9) # Starting low
step.M(x = pi/4, f, h0 = 1000) # Starting high
# where the truncation error estimate is invalid
```

step.plugin	<i>Plug-in step selection</i>
-------------	-------------------------------

Description

Plug-in step selection

Usage

```
step.plugin(
  FUN,
  x,
  h0 = 1e-05 * max(abs(x), sqrt(.Machine$double.eps)),
  max.rel.error = .Machine$double.eps^(7/8),
  range = h0/c(10000, 1e-04),
  cores = 1,
  preschedule = getOption("pnd.preschedule", TRUE),
  cl = NULL,
  ...
)
```

Arguments

<code>FUN</code>	Function for which the optimal numerical derivative step size is needed.
<code>x</code>	Numeric scalar: the point at which the derivative is computed and the optimal step size is estimated.
<code>h0</code>	Numeric scalar: initial step size, defaulting to a relative step of slightly greater than $.Machine$double.eps^{1/3}$ (or absolute step if $x == 0$). This step size for first derivatives is internally translated into the initial step size for third derivatives by multiplying it by the machine epsilon raised to the power $-2/15$.
<code>max.rel.error</code>	Positive numeric scalar > 0 indicating the maximum relative error of function evaluation. For highly accurate functions with all accurate bits is equal to $.Machine$double.eps/2$. For noisy functions (derivatives, integrals, output of optimisation routines etc.), it is higher, typically $\sqrt{.Machine$double.eps}$. Dumontet and Vignes recommend $.Machine$double.eps^{3/4} = 2e-12$ for common functions.
<code>range</code>	Numeric vector of length 2 defining the valid search range for the step size.
<code>cores</code>	Integer specifying the number of CPU cores used for parallel computation. Recommended to be set to the number of physical cores on the machine minus one.
<code>preschedule</code>	Logical: if TRUE, disables pre-scheduling for <code>mclapply()</code> or enables load balancing with <code>parLapplyLB()</code> . Recommended for functions that take less than 0.1 s per evaluation.
<code>cl</code>	An optional user-supplied cluster object (created by <code>makeCluster</code> or similar functions). If not NULL, the code uses <code>parLapply()</code> (if <code>preschedule</code> is TRUE) or <code>parLapplyLB()</code> on that cluster on Windows, and <code>mclapply</code> (fork cluster) on everything else.

... Passed to FUN.

Details

This function computes the optimal step size for central differences using the plug-in approach. The optimal step size is determined as the minimiser of the total error, which for central finite differences is (assuming minimal bounds for relative rounding errors)

$$\sqrt[3]{1.5 \frac{f'(x)}{f'''(x)} \epsilon_{\text{mach}}}.$$

If the estimated third derivative is too small, the function assumes a third derivative of 1 to prevent division-by-zero errors.

Value

A list similar to the one returned by `optim()`:

- `par` – the optimal step size found.
- `value` – the estimated numerical first derivative (using central differences).
- `counts` – the number of iterations (in this case, it is 2).
- `abs.error` – an estimate of the truncation and rounding errors.
- `exitcode` – an integer code indicating the termination status:
 - 0 – Termination with checks passed tolerance.
 - 1 – Third derivative is exactly zero; large step size preferred.
 - 2 – Third derivative is too close to zero; large step size preferred.
 - 3 – Solution lies at the boundary of the allowed value range.
- `message` – A summary message of the exit status.
- `iterations` – A list including the two-step size search path, argument grids, function values on those grids, and estimated third derivative values.

References

There are no references for Rd macro `\insertAllCites` on this help page.

Examples

```
f <- function(x) x^4
step.plugin(x = 2, f)
step.plugin(x = 0, f) # f''' = 0, setting a large one
```

step.SW

*Stepleman–Winarsky automatic step selection***Description**

Stepleman–Winarsky automatic step selection

Usage

```

step.SW(
  FUN,
  x,
  h0 = 1e-05 * (abs(x) + (x == 0)),
  shrink.factor = 0.5,
  range = h0/c(1e+12, 1e-08),
  seq.tol = 1e-04,
  max.rel.error = .Machine$double.eps/2,
  maxit = 40L,
  cores = 1,
  preschedule = getOption("pnd.preschedule", TRUE),
  cl = NULL,
  ...
)

```

Arguments

<code>FUN</code>	Function for which the optimal numerical derivative step size is needed.
<code>x</code>	Numeric scalar: the point at which the derivative is computed and the optimal step size is estimated.
<code>h0</code>	Numeric scalar: initial step size, defaulting to a relative step of slightly greater than $.Machine$double.eps^{1/3}$ (or absolute step if $x == 0$).
<code>shrink.factor</code>	A scalar less than 1 that is used to multiply the step size during the search. The authors recommend 0.25, but this may result in earlier termination at slightly sub-optimal steps. Change to 0.5 for a more thorough search.
<code>range</code>	Numeric vector of length 2 defining the valid search range for the step size.
<code>seq.tol</code>	Numeric scalar: maximum relative difference between old and new step sizes for declaring convergence.
<code>max.rel.error</code>	Positive numeric scalar > 0 indicating the maximum relative error of function evaluation. For highly accurate functions with all accurate bits is equal to half of machine epsilon. For noisy functions (derivatives, integrals, output of optimisation routines etc.), it is higher.
<code>maxit</code>	Maximum number of algorithm iterations to avoid infinite loops. Consider trying some smaller or larger initial step size <code>h0</code> if this limit is reached.
<code>cores</code>	Integer specifying the number of CPU cores used for parallel computation. Recommended to be set to the number of physical cores on the machine minus one.

preschedule	Logical: if TRUE, disables pre-scheduling for mclapply() or enables load balancing with parLapplyLB(). Recommended for functions that take less than 0.1 s per evaluation.
cl	An optional user-supplied cluster object (created by makeCluster or similar functions). If not NULL, the code uses parLapply() (if preschedule is TRUE) or parLapplyLB() on that cluster on Windows, and mclapply (fork cluster) on everything else.
...	Passed to FUN.

Details

This function computes the optimal step size for central differences using the (Stepleman and Winarsky 1979) algorithm.

Value

A list similar to the one returned by optim():

- par – the optimal step size found.
- value – the estimated numerical first derivative (using central differences).
- counts – the number of iterations (each iteration includes four function evaluations).
- abs.error – an estimate of the truncation and rounding errors.
- exitcode – an integer code indicating the termination status:
 - 0 – Optimal termination within tolerance.
 - 2 – No change in step size within tolerance.
 - 3 – Solution lies at the boundary of the allowed value range.
 - 4 – Maximum number of iterations reached.
- message – A summary message of the exit status.
- iterations – A list including the full step size search path, argument grids, function values on those grids, estimated derivative values, estimated error values, and monotonicity check results.

References

Stepleman RS, Winarsky ND (1979). “Adaptive numerical differentiation.” *Mathematics of Computation*, **33**(148), 1257–1264. doi:[10.1090/s00255718197905379698](https://doi.org/10.1090/s00255718197905379698).

Examples

```
f <- function(x) x^4 # The derivative at 1 is 4
step.SW(x = 1, f)
step.SW(x = 1, f, h0 = 1e-9) # Starting too low
# Starting somewhat high leads to too many preliminary iterations
step.SW(x = 1, f, h0 = 10)
step.SW(x = 1, f, h0 = 1000) # Starting absurdly high

f <- sin # The derivative at pi/4 is sqrt(2)/2
```

```

step.SW(x = pi/4, f)
step.SW(x = pi/4, f, h0 = 1e-9) # Starting too low
step.SW(x = pi/4, f, h0 = 0.1) # Starting slightly high
# The following two example fail because the truncation error estimate is invalid
step.SW(x = pi/4, f, h0 = 10)   # Warning
step.SW(x = pi/4, f, h0 = 1000) # Warning

```

stepx

Default step size at given points

Description

Compute an appropriate finite-difference step size based on the magnitude of x , derivation order, and accuracy order. If the function and its higher derivatives belong to the same order of magnitude, this step is near-optimal. For small x , returns a hard bound to prevent large machine-rounding errors.

Usage

```
stepx(x, deriv.order = 1, acc.order = 2, zero.tol = sqrt(.Machine$double.eps))
```

Arguments

<code>x</code>	Numeric vector or scalar: the point(s) at which the derivative is estimated. $\text{FUN}(x)$ must be finite.
<code>deriv.order</code>	Integer or vector of integers indicating the desired derivative order, d^m/dx^m , for each element of x .
<code>acc.order</code>	Integer or vector of integers specifying the desired accuracy order for each element of x . The final error will be of the order $O(h^{\text{acc.order}})$.
<code>zero.tol</code>	Small positive integer: if $\text{abs}(x) \geq \text{zero.tol}$, then, the automatically guessed step size is relative (x multiplied by the step), unless an auto-selection procedure is requested; otherwise, it is absolute.

Value

A numeric vector of the same length as x with positive step sizes.

Examples

```

stepx(10^(-10:2))
stepx(10^(-10:2), deriv.order = 2, acc.order = 4)

```

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