

Package ‘n1qn1’

March 19, 2020

Title Port of the 'Scilab' 'n1qn1' and 'qnbdf' Modules for
(Un)constrained BFGS Optimization

Version 6.0.1-7

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Description Provides 'Scilab' 'n1qn1', or Quasi-Newton BFGS
``qn'' without constraints and 'qnbdf' or Quasi-Newton BFGS with constraints.
This takes more memory than traditional L-
BFGS. The n1qn1 routine is useful since it allows prespecification of a Hessian.
If the Hessian is near enough the truth in optimization it can speed up the optimization prob-
lem. Both algorithms are described in the
'Scilab' optimization documentation located at
<https://www.scilab.org/sites/default/files/optimization_in_scilab.pdf>.

URL <https://github.com/nlmixrdevelopment/n1qn1>

BugReports <https://github.com/nlmixrdevelopment/n1qn1/issues>

Depends R (>= 3.2)

Imports Rcpp (>= 0.12.3)

Suggests testthat, covr

License CeCILL-2

NeedsCompilation yes

LinkingTo RcppArmadillo (>= 0.5.600.2.0), Rcpp (>= 0.12.3)

LazyData true

RoxygenNote 7.0.2

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Date/Publication 2020-03-19 05:20:02 UTC

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n1qn1	<i>n1qn1 optimization</i>
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Description

This is an R port of the n1qn1 optimization procedure in scilab.

Usage

```
n1qn1(call_eval, call_grad, vars, environment = parent.frame(1), ...,
      epsilon = .Machine$double.eps, max_iterations = 100, nsim = 100,
      imp = 0, invisible = NULL, zm = NULL, restart = FALSE,
      assign = FALSE, print.functions = FALSE)
```

Arguments

call_eval	Objective function
call_grad	Gradient Function
vars	Initial starting point for line search
environment	Environment where call_eval/call_grad are evaluated.
...	Ignored additional parameters.
epsilon	Precision of estimate
max_iterations	Number of iterations
nsim	Number of function evaluations
imp	Verbosity of messages.
invisible	boolean to control if the output of the minimizer is suppressed.
zm	Prior Hessian (in compressed format; This format is output in c.hess).
restart	Is this an estimation restart?
assign	Assign hessian to c.hess in environment environment? (Default FALSE)
print.functions	Boolean to control if the function value and parameter estimates are echoed every time a function is called.

Value

The return value is a list with the following elements:

- value The value at the minimized function.
- par The parameter value that minimized the function.
- H The estimated Hessian at the final parameter estimate.
- c.hess Compressed Hessian for saving curvature.
- n.fn Number of function evaluations
- n.gr Number of gradient evaluations

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Examples

```
## Rosenbrock's banana function
n=3; p=100

fr = function(x)
{
  f=1.0
  for(i in 2:n) {
    f=f+p*(x[i]-x[i-1]**2)**2+(1.0-x[i])**2
  }
  f
}

grr = function(x)
{
  g = double(n)
  g[1]=-4.0*p*(x[2]-x[1]**2)*x[1]
  if(n>2) {
    for(i in 2:(n-1)) {
      g[i]=2.0*p*(x[i]-x[i-1]**2)-4.0*p*(x[i+1]-x[i]**2)*x[i]-2.0*(1.0-x[i])
    }
  }
  g[n]=2.0*p*(x[n]-x[n-1]**2)-2.0*(1.0-x[n])
  g
}

x = c(1.02,1.02,1.02)
eps=1e-3
n=length(x); niter=100L; nsim=100L; imp=3L;
nzm=as.integer(n*(n+13L)/2L)
zm=double(nzm)

(op1 <- n1qn1(fr, grr, x, imp=3))
```

```

## Note there are 40 function calls and 40 gradient calls in the above optimization

## Now assume we know something about the Hessian:
c.hess <- c(797.861115,
           -393.801473,
           -2.795134,
           991.271179,
           -395.382900,
           200.024349)
c.hess <- c(c.hess, rep(0, 24 - length(c.hess)))

(op2 <- n1qn1(fr, grr, x, imp=3, zm=c.hess))

## Note with this knowledge, there were only 29 function/gradient calls

(op3 <- n1qn1(fr, grr, x, imp=3, zm=op1$c.hess))

## The number of function evaluations is still reduced because the Hessian
## is closer to what it should be than the initial guess.

## With certain optimization procedures this can be helpful in reducing the
## Optimization time.

```

qnb

qnb optimization

Description

This is an R port of the qnb which is a BFGS-B optimization procedure in scilab. (R has L-BFGS-B in optim).

Usage

```

qnb(par, fn, gr, lower = -Inf, upper = Inf,
    environment = parent.frame(1),
    zero = sqrt(.Machine$double.eps/7e-07), maxFn = 10000L,
    maxIt = 10000L, epsf = sqrt(.Machine$double.eps),
    epsg = sqrt(.Machine$double.eps), epsx = sqrt(.Machine$double.eps),
    print.functions = FALSE)

```

Arguments

par	Initial parameter estimate
fn	Function
gr	Gradient
lower	Lower Bound for optimization

upper	Upper Bound for optimization
environment	Environment where call_eval/call_grad are evaluated.
zero	Tolerance for Zero
maxFn	Maximum function evaluations
maxIt	Maximum iterations
epsf	Function eps for exiting
epsg	Gradient eps for exiting
epsx	Parameter eps for exiting
print.functions	Boolean to control if the function value and parameter estimates are echoed every time a function is called.

Examples

```

## Rosenbrock's banana function
n=3; p=100

fr = function(x)
{
  f=1.0
  for(i in 2:n) {
    f=f+p*(x[i]-x[i-1]**2)**2+(1.0-x[i])**2
  }
  f
}

grr = function(x)
{
  g = double(n)
  g[1]=-4.0*p*(x[2]-x[1]**2)*x[1]
  if(n>2) {
    for(i in 2:(n-1)) {
      g[i]=2.0*p*(x[i]-x[i-1]**2)-4.0*p*(x[i+1]-x[i]**2)*x[i]-2.0*(1.0-x[i])
    }
  }
  g[n]=2.0*p*(x[n]-x[n-1]**2)-2.0*(1.0-x[n])
  g
}

x = c(1.02,1.02,1.02)

op1 <- qnb(x, fr, grr)

```

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