Optimization and related uses of autodiffr: Illustrations

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Introduction

autodiffr is an **R** package to perform automatic differentiation of **R** functions by calling the automatic differentiation tools in the **Julia** programming language. The document **FandR** (??ref) describes how to install autodiffr.

Here we will illustrate how autodiffr can be used to provide gradients or other derivative information for use in optimization problems for which **R** is being used to attempt solutions.

Problem setup

Most methods for optimization require

- an objective function that is to be minimized or maximized. Because the minimum of f(x) is the maximum of -f(x), we will only talk of minimizing functions. Though the mathematics of this are trivial, the care and attention to avoid errors when translating a maximization problem to one involving minimization require serious effort and continuous checking.
- a starting set of values for the parameters to be optimized (and perhaps also for any exogenous data and/or fixed parameters)

We need to load the autodiffr package and then initiate it. NOTE: This is quite slow the first time it is run. WORSE: It is always treated as a "first time" when called in knitr during the processing of a vignette from an Rmd-type file.

```
library(autodiffr)
## Attaching package: 'autodiffr'
## The following object is masked from 'package:stats':
##
##
       deriv
ad_setup()
## Julia version 0.6.3 at location /usr/local/bin will be used.
## Loading setup script for JuliaCall...
## Finish loading setup script for JuliaCall.
And we can use package numDeriv to compare with autodiffr.
require(numDeriv)
## Loading required package: numDeriv
## Attaching package: 'numDeriv'
## The following objects are masked from 'package:autodiffr':
##
       grad, hessian, jacobian
##
```

Test problem - ViaRes

This is simply to test how we can get the gradient of a function that is defined as the sum of squares of residuals, BUT the residuals are computed in a subsidiary function that must be called.

At July 2, 2018, this gives an error that stops knitr, so evaluation is turned off in the following example.

```
require(autodiffr)
ad_setup() # to ensure it is established

ores <- function(x){
    x # Function will be the parameters. ofn is sum of squares
}

ofn0 <- function(x){ # original ofn
    res <- ores(x) # returns a vector of residual values
    val <- as.numeric(crossprod(res)) # as.numeric because crossprod is a matrix
    val
}

ofn <- function(x){ # But autodiffr does not understand crossprod()
    res <- ores(x) # returns a vector of residual values
    val <- sum(res*res) # NOT crossprod()
    val
}</pre>
```

Note that this works with eval=TRUE, but Chebyquad still failing.

```
## Now try to generate the gradient function
ogr <- autodiffr::grad(ofn)

# print(ogr) # this will be more or less meaningless link to Julia function
x0 <- c(1,2,3)
print(ofn(x0)) # should be 14

## [1] 14

print(ofn0(x0)) # should be 14

## [1] 14

ogr0<-ogr(x0) # should be 2, 4, 6
ogr0

## [1] 2 4 6</pre>
```

Test problem – Chebyquad

This problem was given prominence in the optimization literature by Fletcher (1965).

First let us define our Chebyquad function. Note that this is for the **vector** x. This version is expressed as the sum of squares of a vector of function values, which provide a nonlinear least squares problem. Note that **crossprod()** may cause difficulties as it is not written in **R**.

```
require(autodiffr)
ad_setup()

cyq.f <- function (x) {</pre>
```

```
rv<-cyq.res(x)
 f <- sum(rv*rv)
}
cyq.res <- function (x) {</pre>
# Fletcher's chebyquad function m = n -- residuals
  n<-length(x)
  res<-zeros(x) # need to use zeros() to do initialization instead of rep() otherwise autodiffr will b
  ## This is because later on res[i] <- rr,</pre>
  ## if res is a normal R vector and rr is some Julia Number used by autodiffr,
  ## then R doesn't know what to do.
  for (i in 1:n) { #loop over resids
     rr<-0.0
    for (k in 1:n) {
    z7 < -1.0
    z2<-2.0*x[k]-1.0
        z8<-z2
        j<-1
        while (j<i) {
            z6<-z7
            z7<-z8
            z8<-2*z2*z7-z6 # recurrence to compute Chebyshev polynomial
        } # end recurrence loop
        rr<-rr+z8
      } # end loop on k
      rr<-rr/n
      if (2*trunc(i/2) == i) \{ rr <- rr + 1.0/(i*i - 1) \}
      res[i]<-rr
    } # end loop on i
    res
}
```

Let us choose a single value for the number of parameters, and for illustration use n = 4.

```
## cyq.setup
n <- 4
  lower<-rep(-10.0, n)
  upper<-rep(10.0, n)
  x<-1:n
  x<-x/(n+1.0) # Initial value suggested by Fletcher</pre>
```

For safety, let us check the function and a numerical approximation to the gradient.

```
require(numDeriv)
cat("Initial parameters:")

## Initial parameters:

print(x)

## [1] 0.2 0.4 0.6 0.8

cat("Initial value of the function is ",cyq.f(x),"\n")
```

Initial value of the function is 0.07118393

```
gn <- numDeriv::grad(cyq.f, x) # using numDeriv
cat("Approximation to gradient at initial point:")
## Approximation to gradient at initial point:
print(gn)</pre>
```

```
## [1] 0.6170624 0.1882112 -0.1882112 -0.6170624
```

Using a modular approach to the problem, first specifying it via **residuals** and computing the function as a sum of squares, we can also generate the gradient.

```
# Ref: Fletcher, R. (1965) Function minimization without calculating derivatives -- a review,
           Computer J., 8, 33-41.
# Note we do not have all components here e.g., .jsd, .h
cyq.jac<- function (x) {</pre>
# Chebyquad Jacobian matrix
   n<-length(x)
   cj<-matrix(0.0, n, n)</pre>
   for (i in 1:n) { # loop over rows
     for (k in 1:n) { # loop over columns (parameters)
       z5 < -0.0
       cj[i,k] < -2.0
       z8<-2.0*x[k]-1.0
       z2<-z8
       z7 < -1.0
       j<- 1
       while (j<i) { # recurrence loop</pre>
         z4<-z5
         z5 < -cj[i,k]
         cj[i,k] < -4.0*z8+2.0*z2*z5-z4
         z6<-z7
         z7<-z8
         z8<-2.0*z2*z7-z6
         j<- j+1
       } # end recurrence loop
       cj[i,k] < -cj[i,k]/n
     } # end loop on k
   } # end loop on i
   сj
}
cyq.g <- function (x) {</pre>
   cj<-cyq.jac(x)</pre>
   rv<-cyq.res(x)
   gg<- 2.0 * as.vector(rv %*% cj)
}
# check gradient function cyq.g
gajn <- cyq.g(x)</pre>
```

```
print(gajn)
## [1] 0.6170624 0.1882112 -0.1882112 -0.6170624
We can now try to see if autodiffr matches this gradient. However, the following code gives an error in
Julia.
# Do not evaluate, as this fails # Now it should work
cyq.ag <- autodiffr::grad(cyq.f)</pre>
gaag <- cyq.ag(x)</pre>
print(gaag)
## [1] 0.6170624 0.1882112 -0.1882112 -0.6170624
As a workaround, we can get the Chebyquad function from the package funconstrain. The funconstrain
offering does NOT require a call to the residuals, but has a single level R function.
require(funconstrain)
## Loading required package: funconstrain
cat("funconstrain loaded\n")
## funconstrain loaded
cheb <- chebyquad() # Seem to need the brackets or doesn't return pieces
print(str(cheb))
## List of 4
## $ fn:function (par)
## $ gr:function (par)
## $ fg:function (par)
## $x0:function (n = 50)
## NULL
cyq2.f <- cheb$fn
## Note that funconstrain offers the starting value
## x0b <- cheb$x0(n=4) # Need the size of the vector
## x0b
## cyq2.f(x0b)
## same as
print(cyq2.f(x))
## [1] 0.07118393
## Try the gradient
cyq2.ag <- autodiffr::grad(cyq2.f) # Need autodiffr:: specified for knitr
## print(cyq2.g)
cat("Gradient at x")
## Gradient at x
g2ag <- cyq2.ag(x)
print(g2ag)
## [1] 0.6170624 0.1882112 -0.1882112 -0.6170624
require(microbenchmark)
```

Loading required package: microbenchmark

```
cat("cyq.f timing:\n")
## cyq.f timing:
tcyq.f <- microbenchmark(cyq.f(x))</pre>
tcyq.f
## Unit: microseconds
       expr min
                       lq
                             mean median
                                                uq
                                                        max neval
## cyq.f(x) 380.923 383.4005 399.5041 386.2025 401.7775 742.449
cat("cyq2.f timing:\n")
## cyq2.f timing:
tcyq2.f <- microbenchmark(cyq2.f(x))</pre>
tcyq2.f
## Unit: microseconds
## expr min lq mean median uq max neval
## cyq2.f(x) 6.282 6.47 6.82894 6.5945 6.7255 24.854
tcyq.g <- microbenchmark(cyq.g(x))</pre>
tcyq.g
## Unit: microseconds
       expr min
                             mean median
                        lq
                                                        max neval
                                               uq
## cyq.g(x) 419.802 424.5455 490.2539 428.6315 460.204 4627.867 100
tcyq.g <- microbenchmark(cyq.g(x))</pre>
tcyq.g
## Unit: microseconds
## expr min
                     lq mean median uq
                                                     max neval
## cyq.g(x) 419.766 424.2015 442.3977 427.682 453.912 721.357 100
cyq2.g <- cheb$gr
tcyq2.g <- microbenchmark(cyq2.g(x), unit="us")
# microseconds
tcyq2.g
## Unit: microseconds
        expr min lq mean median uq
##
                                                  max neval
## cyq2.g(x) 10.886 11.3015 11.99085 11.599 12.057 32.397 100
# These are very slow
tcyq.ag <- microbenchmark(cyq.ag(x), unit="us")</pre>
# microseconds
tcyq.ag
## Unit: microseconds
             min lq
        expr
                              mean median
                                                        max neval
                                                   uq
## cyq.ag(x) 104337.8 106343.8 112178.1 108440.7 117579.6 157148
tcyq2.ag <- microbenchmark(cyq2.ag(x), unit="us" )</pre>
# microseconds
tcyq2.ag
## Unit: microseconds
##
       expr min lq mean median uq
                                                         max neval
```

```
## cyq2.ag(x) 26536.71 27265.48 28312.42 27558.83 28022.19 45087.62
# These are quicker, but still slow
cyq.optimized ag <- autodiffr::grad(cyq.f, xsize = runif(length(x)), use tape = TRUE)
cyq2.optimized_ag <- autodiffr::grad(cyq2.f, xsize = runif(length(x)), use_tape = TRUE)</pre>
tcyq.optimized ag <- microbenchmark(cyq.optimized ag(x), unit="us")
# microseconds
tcyq.optimized_ag
## Unit: microseconds
                             min
                                               mean
                                                      median
                                        lq
   cyq.optimized_ag(x) 3982.806 4215.243 8050.274 4343.735 4523.958 369652.4
   neval
##
##
      100
tcyq2.optimized_ag <- microbenchmark(cyq2.optimized_ag(x), unit="us" )</pre>
# microseconds
tcyq2.optimized_ag
## Unit: microseconds
##
                              min
                                         lq
                                                       median
                                                mean
##
   cyq2.optimized_ag(x) 4048.122 4274.257 6561.349 4398.139 4594.382
##
         max neval
## 213564.4
## The slowness of the optimized method is partly because the overhead of the JuliaCall and autodiffr p
## After interface functions become stable, I will try to carry on some performance optimizations,
## which is a goal of the project at last phase.
## For example, even we only have a very simple function, the timing is high because of the overhead.
foobar <- function(x) sum(x)</pre>
foobar.ag <- autodiffr::grad(foobar, xsize = runif(length(x)), use_tape = TRUE)</pre>
tfoobar.ag <- microbenchmark(foobar.ag(x), unit="us")
tfoobar.ag
## Unit: microseconds
##
            expr
                      min
                               lq
                                       mean
                                              median
                                                                   max neval
## foobar.ag(x) 4046.752 4246.27 4457.229 4400.141 4599.678 5345.813
## Suppose we are dealing with input of larger size, the overhead stays roughly the same,
## so the overhead should matters not that much as in the case n = 4.
## For example, if n = 25, the diffrence of performance in ratio is not that much.
## cyq.setup
n <- 25
lower < -rep(-10.0, n)
upper <-rep(10.0, n)
x<-1:n
x<-x/(n+1.0) # Initial value suggested by Fletcher
tcyq.g <- microbenchmark(cyq.g(x), unit = "us")</pre>
tcyq.g
## Unit: microseconds
                           lq
                                  mean
                                          median
                                                       uq
## cyq.g(x) 6382.439 6581.02 6918.853 6774.551 7106.873 11484.97
tcyq2.g <- microbenchmark(cyq2.g(x), unit="us")
# microseconds
```

```
tcyq2.g
## Unit: microseconds
         expr
               min
                                 mean median
                                                         max neval
                          lq
                                                  uq
## cyq2.g(x) 73.636 75.8015 78.49865 76.771 78.726 118.443
## The bad thing is that if the input size changes, we need to make an optimized gradient again.
cyq.optimized_ag <- autodiffr::grad(cyq.f, xsize = runif(length(x)), use_tape = TRUE)</pre>
cyq2.optimized_ag <- autodiffr::grad(cyq2.f, xsize = runif(length(x)), use_tape = TRUE)</pre>
tcyq.optimized_ag <- microbenchmark(cyq.optimized_ag(x), unit="us" )</pre>
# microseconds
tcyq.optimized_ag
## Unit: microseconds
##
                                        lq
                                               mean
                                                      median
                   expr
                             min
                                                                            max
## cyq.optimized_ag(x) 16889.74 17369.67 17752.42 17683.96 18074.09 19619.28
## neval
      100
tcyq2.optimized_ag <- microbenchmark(cyq2.optimized_ag(x), unit="us" )</pre>
# microseconds
tcyq2.optimized_ag
## Unit: microseconds
                    expr
                                                      median
                             min
                                        lq
                                               mean
## cyq2.optimized ag(x) 4307.37 4541.206 4749.065 4701.762 4859.06 5987.145
## neval
##
      100
## Also note it is better to check the correctness when generating optimized gradient,
all.equal(cyq.g(x), cyq.optimized_ag(x))
## [1] TRUE
all.equal(cyq.g(x), cyq2.optimized_ag(x))
## [1] TRUE
## Benchmarking times without user interface wrappers
tape1 <- reverse.grad.tape(cyq.f, runif(length(x)))</pre>
microbenchmark(reverse.grad(tape1, x), unit="us")
## Unit: microseconds
                                min
                                           lq
                                                  mean
                                                         median
  reverse.grad(tape1, x) 17255.68 17885.36 18446.76 18176.75 18503.74
##
##
         max neval
## 40425.25
tape2 <- reverse.grad.tape(cyq2.f, runif(length(x)))</pre>
microbenchmark(reverse.grad(tape2, x), unit="us")
## Unit: microseconds
                      expr
                                min
                                           lq
                                                  mean
                                                         median
  reverse.grad(tape2, x) 4385.389 4614.903 4935.386 4871.044 5135.894
##
         max neval
## 6221.859
               100
## Benchmarking times without autodiffr wrappers
JuliaCall::julia_command("using ReverseDiff")
```

```
tape1 <- reverse.grad.tape(cyq.f, runif(length(x)))</pre>
microbenchmark(JuliaCall::julia_call("ReverseDiff.gradient!", tape1, x), unit="us")
## Unit: microseconds
##
                                                          expr
                                                                    min
##
    JuliaCall::julia_call("ReverseDiff.gradient!", tape1, x) 11247.86
##
                mean
                       median
                                             max neval
          lq
                                     uq
  11636.34 12232.4 11897.51 12200.57 37916.07
tape2 <- reverse.grad.tape(cyq2.f, runif(length(x)))</pre>
microbenchmark(JuliaCall::julia call("ReverseDiff.gradient!", tape2, x), unit="us" )
## Unit: microseconds
##
                                                          expr
    JuliaCall::julia_call("ReverseDiff.gradient!", tape2, x) 545.651 551.974
##
        mean
               median
                           uq
                                    max neval
  594.6466 589.7005 610.664 1201.589
```

Test problem – Hobbs weed infestation

This nonlinear estimation problem was brought to one of the authors (JN) in the mid-1970s (See Nash (1979)). It has just 12 data points and asks for the estimation of a 3-parameter logistic growth curve. The present example does not provide for scaling.

```
hobbs.f<- function(x){ # # Hobbs weeds problem -- function
    if (abs(12*x[3]) > 500) { # check computability
       fbad<-.Machine$double.xmax
       return(fbad)
    }
    res<-hobbs.res(x)
    f<-sum(res*res)
}
hobbs.res<-function(x){ # Hobbs weeds problem -- residual
# This variant uses looping
    if(length(x) != 3) stop("hobbs.res -- parameter vector n!=3")
    y < -c(5.308, 7.24, 9.638, 12.866, 17.069, 23.192, 31.443, 38.558, 50.156, 62.948,
         75.995, 91.972)
    t<-1:12
    if(abs(12*x[3])>50) {
       res<-rep(Inf,12)
    } else {
       res < -x[1]/(1+x[2]*exp(-x[3]*t)) - y
    }
}
hobbs.jac<-function(x){ # Jacobian of Hobbs weeds problem
   jj<-matrix(0.0, 12, 3)
   t<-1:12
    yy < -exp(-x[3]*t)
    zz<-1.0/(1+x[2]*yy)
     jj[t,1] <- zz
     jj[t,2] \leftarrow -x[1]*zz*zz*yy
```

```
jj[t,3] \leftarrow x[1]*zz*zz*yy*x[2]*t
     jjret <- jj
     attr(jjret, "gradient") <- jj</pre>
   return(jjret)
}
hobbs.g<-function(x){ # gradient of Hobbs weeds problem
    # NOT EFFICIENT TO CALL AGAIN
    jj<-hobbs.jac(x)</pre>
    res<-hobbs.res(x)
    gg<-as.vector(2.*t(jj) %*% res)</pre>
    return(gg)
}
hobbs.rsd<-function(x) { # Jacobian second derivative
    rsd < -array(0.0, c(12,3,3))
    t<-1:12
    yy < -exp(-x[3]*t)
    zz<-1.0/(1+x[2]*yy)
    rsd[t,1,1] \leftarrow 0.0
    rsd[t,2,1] \leftarrow -yy*zz*zz
    rsd[t,1,2] \leftarrow -yy*zz*zz
    rsd[t,2,2]<- 2.0*x[1]*yy*yy*zz*zz*zz
    rsd[t,3,1] \leftarrow t*x[2]*yy*zz*zz
    rsd[t,1,3] \leftarrow t*x[2]*yy*zz*zz
    rsd[t,3,2] \leftarrow t*x[1]*yy*zz*zz*(1-2*x[2]*yy*zz)
    rsd[t,2,3] \leftarrow t*x[1]*yy*zz*zz*(1-2*x[2]*yy*zz)
##
      rsd[t,3,3]<- 2*t*t*x[1]*x[2]*x[2]*yy*yy*zz*zz*zz
    rsd[t,3,3]<- -t*t*x[1]*x[2]*yy*zz*zz*(1-2*yy*zz*x[2])
    return(rsd)
}
hobbs.h <- function(x) { ## compute Hessian
   cat("Hessian not yet available\n")
   return(NULL)
    H<-matrix(0,3,3)</pre>
    res<-hobbs.res(x)
    jj<-hobbs.jac(x)</pre>
    rsd<-hobbs.rsd(x)
      H<-2.0*(t(res) %*% rsd + t(jj) %*% jj)</pre>
##
    for (j in 1:3) {
        for (k in 1:3) {
           for (i in 1:12) {
              H[j,k] \leftarrow H[j,k] + res[i] * rsd[i,j,k]
        }
    H<-2*(H + t(jj) %*% jj)
    return(H)
}
x0good \leftarrow c(200, 50, 0.3)
```

```
x0bad <- c(1,1,1)
f0good <- hobbs.f(x0good)</pre>
cat("Sum of squares at the GOOD starting point:",f0good,"\n")
## Sum of squares at the GOOD starting point: 158.2324
f0bad <- hobbs.f(x0bad)
cat("Sum of squares at the BAD starting point:",f0bad,"\n")
## Sum of squares at the BAD starting point: 23520.58
res0good <- hobbs.res(x0good)
## Residuals -- good starting point
res0good
## [1] -0.05050236 -0.20779506 -0.26086802 -0.41247546 -0.61690702
## [6] -1.60525418 -3.36423901 -2.43016453 -4.28734016 -5.63079076
## [11] -5.67542775 -7.44779537
res0bad <- hobbs.res(x0bad)
## Residuals -- bad starting point
res0bad
## [1] -4.576941 -6.359203 -8.685426 -11.883986 -16.075693 -22.194473
## [7] -30.443911 -37.558335 -49.156123 -61.948045 -74.995017 -90.972006
require(autodiffr)
ad_setup()
hobbs.ag <- autodiffr::grad(hobbs.f)</pre>
hobbsag0good <- hobbs.ag(x0good)
## Gradient by AD -- good starting point
hobbsag0good
## [1]
          -17.8438
                       48.2491 -24559.8187
## Compare hand coded function
hobbsggood <- hobbs.g(x0good)</pre>
hobbsggood
                       48.2491 -24559.8187
## [1]
          -17.8438
## Gradient by AD -- bad starting point
hobbsag0bad <- hobbs.ag(x0bad)</pre>
hobbsag0bad
## [1] -824.042084
                      4.764888 -11.025384
## Compare hand coded function
hobbsgbad <- hobbs.g(x0bad)
hobbsgbad
## [1] -824.042084
                      4.764888 -11.025384
## Interestingly, the magnitude of gradient elements greater for "good"
hobbs.aj <- autodiffr::jacobian(hobbs.res)</pre>
## Gradient by AD -- good starting point
hobbsaj0good <- hobbs.aj(x0good)
hobbsaj0good
```

```
##
               [,1]
                          [,2]
                                     [,3]
    [1,] 0.02628749 -0.1023858
                                 5.119291
    [2,] 0.03516102 -0.1356989
##
                                13.569891
##
    [3,] 0.04688566 -0.1787496
                                26.812437
   [4,] 0.06226762 -0.2335615 46.712293
   [5,] 0.08226046 -0.3019747 75.493681
##
   [6,] 0.10793373 -0.3851362 115.540847
##
  [7,] 0.14039380 -0.4827335 168.956738
## [8,] 0.18063918 -0.5920347 236.813864
## [9,] 0.22934330 -0.7069798 318.140911
## [10,] 0.28658605 -0.8178179 408.908969
## [11,] 0.35159786 -0.9119072 501.548971
## [12,] 0.42262102 -0.9760500 585.629985
## Compare hand coded function
hobbsjgood <- hobbs.jac(x0good)
hobbsjgood
##
                          [,2]
                                     [,3]
               [,1]
   [1,] 0.02628749 -0.1023858
                                 5.119291
##
   [2,] 0.03516102 -0.1356989
                                13.569891
##
   [3,] 0.04688566 -0.1787496
                                26.812437
##
  [4,] 0.06226762 -0.2335615 46.712293
  [5,] 0.08226046 -0.3019747 75.493681
##
   [6,] 0.10793373 -0.3851362 115.540847
   [7,] 0.14039380 -0.4827335 168.956738
  [8,] 0.18063918 -0.5920347 236.813864
   [9,] 0.22934330 -0.7069798 318.140911
## [10,] 0.28658605 -0.8178179 408.908969
  [11,] 0.35159786 -0.9119072 501.548971
   [12,] 0.42262102 -0.9760500 585.629985
## attr(,"gradient")
                          [,2]
##
               [,1]
                                     [,3]
    [1,] 0.02628749 -0.1023858
##
                                 5.119291
   [2,] 0.03516102 -0.1356989 13.569891
##
  [3,] 0.04688566 -0.1787496
                                26.812437
   [4,] 0.06226762 -0.2335615
##
                                46.712293
##
  [5,] 0.08226046 -0.3019747 75.493681
## [6,] 0.10793373 -0.3851362 115.540847
## [7,] 0.14039380 -0.4827335 168.956738
   [8,] 0.18063918 -0.5920347 236.813864
  [9,] 0.22934330 -0.7069798 318.140911
## [10,] 0.28658605 -0.8178179 408.908969
## [11,] 0.35159786 -0.9119072 501.548971
## [12,] 0.42262102 -0.9760500 585.629985
## Gradient by AD -- bad starting point
hobbsajObad <- hobbs.aj(xObad)
hobbsaj0bad
##
              [,1]
                            [,2]
                                         [,3]
##
   [1,] 0.7310586 -1.966119e-01 1.966119e-01
  [2,] 0.8807971 -1.049936e-01 2.099872e-01
## [3,] 0.9525741 -4.517666e-02 1.355300e-01
```

```
[4,] 0.9820138 -1.766271e-02 7.065082e-02
##
   [5,] 0.9933071 -6.648057e-03 3.324028e-02
   [6,] 0.9975274 -2.466509e-03 1.479906e-02
##
   [7,] 0.9990889 -9.102212e-04 6.371548e-03
##
   [8,] 0.9996646 -3.352377e-04 2.681901e-03
##
  [9,] 0.9998766 -1.233793e-04 1.110414e-03
## [10,] 0.9999546 -4.539581e-05 4.539581e-04
## [11,] 0.9999833 -1.670114e-05 1.837126e-04
## [12,] 0.9999939 -6.144137e-06 7.372964e-05
## Compare hand coded function
hobbsjbad <- hobbs.jac(x0bad)
hobbsjbad
##
                             [,2]
              [,1]
                                          [,3]
##
    [1,] 0.7310586 -1.966119e-01 1.966119e-01
    [2,] 0.8807971 -1.049936e-01 2.099872e-01
##
##
    [3,] 0.9525741 -4.517666e-02 1.355300e-01
##
    [4,] 0.9820138 -1.766271e-02 7.065082e-02
##
   [5,] 0.9933071 -6.648057e-03 3.324028e-02
##
   [6,] 0.9975274 -2.466509e-03 1.479906e-02
   [7,] 0.9990889 -9.102212e-04 6.371548e-03
##
##
    [8,] 0.9996646 -3.352377e-04 2.681901e-03
   [9,] 0.9998766 -1.233793e-04 1.110414e-03
##
## [10,] 0.9999546 -4.539581e-05 4.539581e-04
  [11,] 0.9999833 -1.670114e-05 1.837126e-04
   [12,] 0.9999939 -6.144137e-06 7.372964e-05
  attr(, "gradient")
##
              [,1]
                             [,2]
##
                                          [,3]
    [1,] 0.7310586 -1.966119e-01 1.966119e-01
##
    [2,] 0.8807971 -1.049936e-01 2.099872e-01
##
   [3,] 0.9525741 -4.517666e-02 1.355300e-01
   [4,] 0.9820138 -1.766271e-02 7.065082e-02
##
   [5,] 0.9933071 -6.648057e-03 3.324028e-02
##
   [6,] 0.9975274 -2.466509e-03 1.479906e-02
##
   [7,] 0.9990889 -9.102212e-04 6.371548e-03
   [8,] 0.9996646 -3.352377e-04 2.681901e-03
##
   [9,] 0.9998766 -1.233793e-04 1.110414e-03
## [10,] 0.9999546 -4.539581e-05 4.539581e-04
## [11,] 0.9999833 -1.670114e-05 1.837126e-04
## [12,] 0.9999939 -6.144137e-06 7.372964e-05
```

Now let us try this in a solution of nonlinear least squares.

WARNING: Because of some compatibility issues with other **R** software, the jacobian must be available in the "gradient" attribute returned by the jacobian function. The purpose of this is to allow the function nlsr::nlfb to have the same name for the residual and jacobian function. This is used in generating a symbolic jacobian function in nlsr::nlxb. However, it can catch unwary users (including us!).

```
# try in a function
require(nlsr)

## Loading required package: nlsr

## manual
smgood <- nlfb(x0good, hobbs.res, hobbs.jac, trace=TRUE)</pre>
```

```
## no weights
## lower:[1] -Inf -Inf -Inf
## upper:[1] Inf Inf Inf
## Start:lamda: 1e-04 SS= 158.2324 at = 200 = 50 = 0.3 1 / 0
\#\# <<lamda: 4e-05 SS= 2.61779 at = 194.3011 = 48.56497 = 0.313994 2 / 1
\#\# <<lamda: 1.6e-05 SS= 2.587325 at = 196.0825 = 49.07632 = 0.313616 3 / 2
## <<lamda: 6.4e-06 SS= 2.587277 at = 196.1839 = 49.09134 = 0.313571 4 / 3
## <<lamda: 2.56e-06 SS= 2.587277 at = 196.1862 = 49.09164 = 0.3135697 5 / 4
## <<lamda: 1.024e-06 SS= 2.587277 at = 196.1863 = 49.09164 = 0.3135697 6 / 5
## WARNING: we need the jacobian in the "gradient" attribute
hobbs.ajx <- function(x){
    jj <- hobbs.aj(x)</pre>
    jjr <- jj
    attr(jjr, "gradient")<- jj # !!! IMPORTANT</pre>
    jjr
}
sagood <- nlfb(x0good, hobbs.res, hobbs.ajx, trace=TRUE)</pre>
## no weights
## lower:[1] -Inf -Inf -Inf
## upper:[1] Inf Inf Inf
## Start:lamda: 1e-04 SS= 158.2324 at = 200 = 50 = 0.3 1 / 0
\#\# <<lamda: 4e-05 SS= 2.61779 at = 194.3011 = 48.56497 = 0.313994 2 / 1
## <<lamda: 1.6e-05 SS= 2.587325 at = 196.0825 = 49.07632 = 0.313616 3 / 2
\#\# << lamda: 6.4e-06 SS= 2.587277 at = 196.1839 = 49.09134 = 0.313571 4 / 3
## <<lamda: 2.56e-06 SS= 2.587277 at = 196.1862 = 49.09164 = 0.3135697 5 / 4
## <<lamda: 1.024e-06 SS= 2.587277 at = 196.1863 = 49.09164 = 0.3135697 6 / 5
```

Test problem – Candlestick

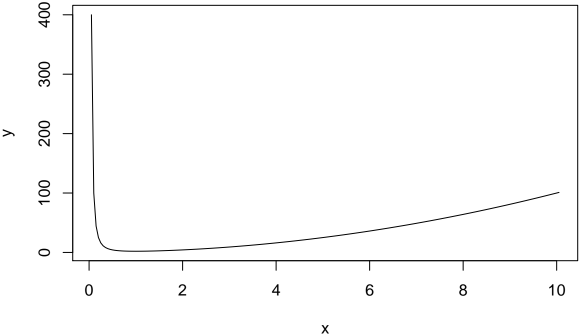
This function was developed by one of us to provide a simple but (for n equal 1 or 2) graphic example of a function with an infinity of solutions for n >= 2. The function can be seen by graphing it to have a spike in the "middle" of a dish, much like some older candlesticks or candle holders. The multiplicity of solutions should make the hessian of a solution singular. For n = 2, for example, the minimum lies on a circular locus at the deepest point of the "saucer".

```
# candlestick function
# J C Nash 2011-2-3
cstick.f<-function(x,alpha=1){
    x<-as.vector(x)
    r2<-sum(x*x)
    f<-as.double(r2+alpha/r2)
    return(f)
}

cstick.g<-function(x,alpha=1){
    x<-as.vector(x)
    r2<-sum(x*x)
    g1<-2*x
    g2 <- (-alpha)*2*x/(r2*r2)
    g<-as.double(g1+g2)
    return(g)</pre>
```

```
x <- seq(-100:100)/20.0
y <- x

for (ii in 1:length(x)){
    y[ii] <- cstick.f(x[ii])
}
plot(x, y, type='l') # ?? does not plot from console??
</pre>
```



```
x0 < -c(1,2)
require(optimx)
## Loading required package: optimx
sdef0 <- optimr(x0, cstick.f, cstick.g, method="Rvmmin", control=list(trace=1))</pre>
## Parameter scaling:[1] 1 1
                                            fval= 5.2
## gradient test tolerance = 6.055454e-06
## compare to max(abs(gn-ga))/(1+abs(fval)) = 6.066115e-12
## Rvmminu -- J C Nash 2009-2015 - an R implementation of Alg 21
## Problem of size n= 2 Dot arguments:
## list()
## Initial fn= 5.2
## ig= 1
          gnorm= 4.293251
                                  1
                                      5.2
## ig= 2
          gnorm= 3.884638
                              2
                                  2
                                      4.468295
                                  3
## *ig= 3
          gnorm= 2.852834
                              4
                                       3.087837
## *ig= 4
           gnorm= 2.154552
                              6
                                  4
                                       2.520416
## ig= 5
          gnorm= 3.731383
                              7
                                  5
                                      2.417429
## ig= 6
          gnorm= 1.086155
                              8
                                  6
                                      2.098346
## ig= 7
          gnorm= 0.504457
                              9
                                  7
                                      2.018112
## ig= 8
          gnorm= 0.1371067
                               10
                                    8
                                        2.001136
                                     9 2.000012
## ig= 9
          gnorm= 0.01377774
                               11
```

```
## ig= 10 gnorm= 0.0003470064 12 10 2
## ig= 11 gnorm= 8.985444e-07 13 11 2
## ig= 12 gnorm= 5.846616e-11 14 12 2
## *****No acceptable point
## Converged
## Seem to be done Rvmminu
sdef0
## $par
## [1] -0.4472136 -0.8944272
## $value
## [1] 2
##
## $counts
## function gradient
##
          19
##
## $convergence
## [1] 0
##
## $message
## [1] "Rvmminu appears to have converged"
xstar <- sdef0$par
gstar <- cstick.g(xstar)</pre>
cat("Gradient at proposed solution:")
## Gradient at proposed solution:
print(gstar)
## [1] -2.614686e-11 -5.229372e-11
## FIXED??
## This doesn't seem to work well??
require(autodiffr)
ad_setup()
hc <- autodiffr::hessian(cstick.f)</pre>
hstar<-hc(xstar)
cat("Hessian at proposed solution:\n")
## Hessian at proposed solution:
print(hstar)
##
      [,1] [,2]
## [1,]
## [2,]
            0
print(eigen(hstar)$values)
## [1] 0 0
## ?? doesn't seem right
hc(x0)
##
      [,1] [,2]
## [1,] 0 0
```

```
## [2,]
require(numDeriv)
hcn0 <- numDeriv::hessian(cstick.f, x0)</pre>
##
         [,1] [,2]
## [1,] 1.984 0.128
## [2,] 0.128 2.176
hcnstar <- numDeriv::hessian(cstick.f, xstar)</pre>
hcnstar
##
        [,1] [,2]
## [1,] 1.6 3.2
## [2,] 3.2 6.4
hcnj0 <- numDeriv::jacobian(cstick.g, x0)</pre>
hcnj0
##
         [,1] [,2]
## [1,] 1.984 0.128
## [2,] 0.128 2.176
hcnjstar <- numDeriv::jacobian(cstick.g, xstar)</pre>
hcnjstar
        [,1] [,2]
##
## [1,] 1.6 3.2
## [2,] 3.2 6.4
eigen(hcnstar)$values
## [1] 8.000000e+00 1.137876e-10
```

Test problem – Wood 4 parameter function

This is reported by Moré, Garbow, and Hillstrom (1980) as coming from Colville (1968). The problem in 4 parameters seems to have a false solution far from the accepted one. Is there a good description of this function and the issues it presents?

```
#hessian:
wood.h <- function(x){
 h11 \leftarrow 1200*x[1]^2-400*x[2]+2; h12 \leftarrow -400*x[1]; h13 \leftarrow h14 \leftarrow 0
 h22 <- 220.2; h23 <- 0; h24 <- 19.8
 h33 \leftarrow 1080*x[3]^2-360*x[4]+2; h34 \leftarrow -360*x[3]
 h44 <- 200.2
 H <- matrix(c(h11,h12,h13,h14,h12,h22,h23,h24,</pre>
                h13,h23,h33,h34,h14,h24,h34,h44),ncol=4)
 return(H)
}
x0 \leftarrow c(-3,-1,-3,-1) \# Wood standard start
cat("Function value at x0=",wood.f(x0),"\n")
## Function value at x0= 19192
wood.ag <- autodiffr::grad(wood.f)</pre>
cat("Autodiffr gradient value:")
## Autodiffr gradient value:
vwag0 < -wood.ag(x0)
print(vwag0)
## [1] -12008 -2080 -10808 -1880
cat("Manually coded:")
## Manually coded:
vwg0 \leftarrow wood.g(x0)
print(vwg0)
## [1] -12008 -2080 -10808 -1880
cat("Differences:\n")
## Differences:
print(vwag0-vwg0)
## [1] 0 0 0 0
cat("Autodiffr hessian of function value:")
## Autodiffr hessian of function value:
wood.ah <- autodiffr::hessian(wood.f)</pre>
vwah0 <- wood.ah(x0)</pre>
print(vwah0)
##
         [,1]
              [,2] [,3]
                              [,4]
## [1,] 11202 1200.0 0
                              0.0
## [2,] 1200 220.2
                         0 19.8
## [3,]
                0.0 10082 1080.0
            0
## [4,]
            0
              19.8 1080 200.2
cat("Autodiffr hessian via jacobian of autodiff gradient value:")
```

Autodiffr hessian via jacobian of autodiff gradient value:

```
wood.ahjag <- autodiffr::jacobian(wood.ag)</pre>
vwahjag0<-wood.ahjag(x0)</pre>
print(vwahjag0)
         [,1]
                [,2] [,3]
                              [, 4]
## [1,] 11202 1200.0
                        0
                              0.0
## [2,] 1200 220.2
                         0
                             19.8
## [3,]
            0
                 0.0 10082 1080.0
              19.8 1080 200.2
## [4,]
cat("Autodiffr hessian via jacobian of manual gradient value:")
## Autodiffr hessian via jacobian of manual gradient value:
wood.ahj <- autodiffr::jacobian(wood.g)</pre>
vwahj0 <- wood.ah(x0)</pre>
print(vwahj0)
                              [,4]
         [,1]
                [,2] [,3]
## [1,] 11202 1200.0
                              0.0
## [2,] 1200 220.2
                             19.8
## [3,]
            0
                 0.0 10082 1080.0
## [4,]
               19.8 1080 200.2
cat("Manually coded:")
## Manually coded:
vwh0 < -wood.h(x0)
print(vwh0)
         [,1]
                [,2] [,3]
                              [,4]
## [1,] 11202 1200.0
                              0.0
                         0
## [2,] 1200 220.2
                            19.8
                         0
## [3,]
            0
                 0.0 10082 1080.0
               19.8 1080 200.2
## [4,]
cat("Differences from vwh0\n")
## Differences from vwh0
cat("vwah0\n")
## vwah0
print(vwah0-vwh0)
        [,1] [,2] [,3] [,4]
## [1,]
## [2,]
           0
                0
                     0
                          0
## [3,]
           0
                     0
## [4,]
                     0
                          0
                0
cat("\n")
cat("vwahj0\n")
```

vwahj0

```
print(vwahj0-vwh0)
       [,1] [,2] [,3] [,4]
## [1,]
            0 0
                        0
         0
## [2,]
          0
              0
                   0
                        0
## [3,]
          0
              0
                   0
                        0
## [4,]
          0
                   0
                        0
cat("\n")
cat("vwahjag0\n")
## vwahjag0
print(vwahjag0-vwh0)
       [,1] [,2] [,3] [,4]
## [1,]
       0
             0
                   0
## [2,]
          0
              0
                   0
                        0
## [3,]
          0
                        0
## [4,]
        0
            0
                   0
                        0
cat("\n")
## d <- c(1,1,1,1)
require(optimx)
meths <- c("snewton", "snewtonm", "nlm")</pre>
wdefault <- opm(x0, fn=wood.f, gr=wood.g, hess=wood.h, method=meths, control=list(trace=0))</pre>
print(wdefault)
##
                            value fevals gevals convergence kkt1 kkt2
           p1 p2 p3 p4
## snewton 1 1 1 1.142616e-29 119 70
                                                92 TRUE TRUE
## snewtonm 1 1 1 1.399599e-28 88
                                            50
                                                       O TRUE TRUE
           1 1 1 1.004943e-16 NA
                                                       O TRUE TRUE
## nlm
                                           335
##
           xtime
## snewton 0.008
## snewtonm 0.004
           0.008
## nlm
wagah <- opm(x0, fn=wood.f, gr=wood.ag, hess=wood.ah, method=meths, control=list(trace=0))</pre>
## Small gradient
print(wagah)
                            value fevals gevals convergence kkt1 kkt2
           p1 p2 p3 p4
                                                 O TRUE TRUE
          1 1 1 1 0.000000e+00
## snewton
                                     116 67
## snewtonm 1 1 1 1 0.000000e+00
                                     81
                                            50
                                                        O TRUE TRUE
                                                        O TRUE TRUE
           1 1 1 1.004942e-16
## nlm
                                    NA
                                           335
##
           xtime
## snewton 4.804
## snewtonm 3.456
## nlm
       26.800
```

Performance issues

##

0.024

0.000

0.022

Optimization is, by its very nature, about improving things. Thus it is of prime interest to seek faster and better ways to optimize functions. In this section we look at some issues that may influence the speed, reliability and correctness of optimization calculations.

First, it is critical to note that **R** almost always offers several ways to accomplish the same computational result. However, the speed with which the different approaches return a result can be wildly different. (?? can JN find the 800% scale factor example??).

Second, there are many parts of the autodiffr wrapper of Julia's automatic differentiation that may use up computing cycles:

- We must translate from one programming language to another in some sense in order to call the appropriate functions in Julia based on \mathbf{R} functions.
- Results must be properly structured on return to R.
- Hand coded derivative expressions, especially hand-optimized ones, can be expected to out-perform automatic differentiation results.

NOTE: Performance is interesting, but it is far from the complete picture. We can use results from autodiffr to validate hand-coded functions. We can get results that are efficient of human time and effort that may be otherwise unavailable. Moreover, the results of computing gradients and hessians allow us to conclude that a solution has been achieved.

A small performance comparison using autodiffr

```
rm(list=ls())
require(autodiffr)
autodiffr::ad_setup() # to ensure it is established
ores <- function(x){</pre>
    \boldsymbol{x} # Function will be the parameters. ofn is sum of squares
logit <- function(x) exp(x) / (1 + exp(x))
ofn <- function(x){
    res <- ores(x) # returns a vector of residual values
    sum(logit(res) ^ 2)
}
## Now try to generate the gradient function
ogr <- autodiffr::grad(ofn)</pre>
system.time(ogr(runif(100)))
##
      user system elapsed
##
      0.24
              0.00
                       0.24
system.time(ogr(runif(100)))
##
      user system elapsed
```

```
ogr1 <- autodiffr::grad(ofn, xsize = runif(100))</pre>
system.time(ogr1(runif(100)))
      user system elapsed
##
     0.016
           0.000
                   0.014
system.time(ogr1(runif(100)))
     user system elapsed
##
##
     0.016
           0.000
                   0.015
ogr2 <- autodiffr::grad(ofn, xsize = runif(100), use_tape = TRUE)
system.time(ogr2(runif(100)))
##
     user system elapsed
##
    0.132
           0.000
                    0.129
system.time(ogr2(runif(100)))
##
      user system elapsed
##
     0.008
            0.000
                    0.007
```

A problem with discontinuous gradient

Restarts for stagnation =0

Problems with discontinuous gradient may give gradient methods difficulty.

Here is a problem where the gradient is discontinuous, but not at the minimum.

```
## discontin.R, a test function with discontinuous gradient
disc.f <- function(x){</pre>
   nn <- length(x)
   val <- 0.0
   for (ii in 1:nn){
       tt <- (x[ii] - ii)
       if (abs(tt) < ii) {</pre>
            ff <- tt*tt
       } else {
            ff <- abs(tt)
       }
       val \leftarrow val + ff*ff
    }
    val
}
require(optimx)
x0 <- runif(4)
x0
## [1] 0.8135548 0.7440748 0.6562804 0.7350514
sol0 <- optimr(x0, disc.f, method="nmkb")</pre>
## Warning in optimr(x0, disc.f, method = "nmkb"): Successful convergence
```

```
sol0
## [1] 1.002540 1.991503 2.987462 4.022389
## $value
## [1] 2.812157e-07
##
## $convergence
## [1] 0
##
## $message
## [1] "Successful convergence"
##
## $counts
## [1] 69 NA
##
## $nitns
## [1] NA
require(autodiffr)
ad_setup()
disc.ag <- autodiffr::grad(disc.f)</pre>
sol1 <- optimr(x0, disc.f, disc.ag, method="Rvmmin", control=list(trace=1))</pre>
## Parameter scaling:[1] 1 1 1 1
## gradient test tolerance = 6.055454e-06 fval= 146.2958
## compare to max(abs(gn-ga))/(1+abs(fval)) = 1.512723e-11
## Rvmminu -- J C Nash 2009-2015 - an R implementation of Alg 21
## Problem of size n= 4 Dot arguments:
## list()
## Initial fn= 146.2958
## ig= 1
          gnorm= 148.6464
                              1 1 146.2958
## **ig= 2 gnorm= 49.01469
                                       28.94909
## *ig= 3 gnorm= 40.28683
                               6
                                   3
                                       24.09259
                              7 4
## ig= 4
          gnorm= 8.289792
                                      13.97521
## ig= 5
          gnorm= 19.37912
                              8
                                  5
                                      9.100838
## ig= 6
          gnorm= 4.644537
                              9 6
                                      4.993596
## ig= 7
          gnorm= 12.02315 UPDATE NOT POSSIBLE: ilast, ig 1 7
   10
         7 4.621053
## *ig= 8
           gnorm= 3.431155
                               12
                                    8
                                        0.8742065
## ig= 9
          gnorm= 0.2316705
                               13
                                    9
                                        0.02258458
## ig= 10
          gnorm= 0.1679991
                               14
                                     10
                                          0.014737
## ig= 11
           gnorm= 0.05793144
                                 15
                                      11
                                           0.003592968
## ig= 12
           gnorm= 0.02703047
                                 16
                                      12
                                           0.001319294
                                17
                                     13
                                          0.000436323
## ig= 13
           gnorm= 0.0113772
## ig= 14
           gnorm= 0.005236783
                                  18
                                      14 0.0001722984
## ig= 15
           gnorm= 0.002785145
                                       15
                                            8.423379e-05
                                  19
           gnorm= 0.001955969
                                       16
## ig= 16
                                  20
                                           5.345518e-05
                                  21
                                       17
                                            3.957719e-05
## ig= 17
           gnorm= 0.001632266
                                  22
                                       18 3.052032e-05
## ig= 18
           gnorm= 0.001392964
## ig= 19
                                  23
                                       19
                                            2.251137e-05
           gnorm= 0.001113926
                                       20
## ig= 20
           gnorm= 0.000795105
                                  24
                                           1.469809e-05
## ig= 21
           gnorm= 0.000601506
                                  25
                                       21
                                            1.094901e-05
## ig= 22
           gnorm= 0.0005719681
                                  26 22 9.81399e-06
```

```
gnorm= 0.0005016775
                                     27
                                           23
                                                8.114558e-06
## ig= 23
            gnorm= 0.0002166367
                                     28
                                           24
                                                2.575389e-06
## ig= 24
## ig= 25
            gnorm= 0.0001107834
                                                1.062676e-06
## ig= 26
            gnorm= 5.603816e-05
                                                4.429216e-07
                                     30
                                           26
## ig= 27
            gnorm= 3.600459e-05
                                     31
                                           27
                                                2.485985e-07
            gnorm= 2.958579e-05
## ig= 28
                                     32
                                           28
                                                1.864565e-07
## ig= 29
            gnorm= 2.663565e-05
                                     33
                                                1.570322e-07
## ig= 30
            gnorm= 2.282095e-05
                                     34
                                           30
                                                1.220478e-07
            gnorm= 1.68282e-05
## ig= 31
                                     35
                                          31
                                               7.931622e-08
## ig= 32
            gnorm= 1.433972e-05
                                     36
                                           32
                                                6.810163e-08
## ig= 33
            gnorm= 1.261174e-05
                                     37
                                           33
                                                5.550548e-08
## ig= 34
            gnorm= 9.008336e-06
                                     38
                                           34
                                                3.568127e-08
            gnorm= 6.592002e-06
                                     39
                                           35
                                                2.649039e-08
## ig= 35
## ig= 36
            gnorm= 6.00894e-06
                                     40
                                          36
                                               2.606665e-08
## ig= 37
            gnorm= 6.130287e-06
                                     41
                                           37
                                                2.55239e-08
## ig= 38
            gnorm= 6.095345e-06
                                     42
                                           38
                                                2.547318e-08
            gnorm= 5.967945e-06
                                     43
                                           39
                                                2.523985e-08
## ig= 39
            gnorm= 5.818798e-06
                                                2.48371e-08
## ig= 40
                                                2.350164e-08
## ig= 41
            gnorm= 5.459854e-06
                                     45
                                           41
## ig= 42
            gnorm= 4.84135e-06
                                     46
                                          42
                                               2.04056e-08
## ig= 43
            gnorm= 3.67076e-06
                                     47
                                          43
                                               1.371316e-08
                                                5.981338e-09
## ig= 44
            gnorm= 2.072426e-06
                                     48
            gnorm= 9.934191e-07
                                                2.128672e-09
## ig= 45
                                     49
                                           45
## ig= 46
            gnorm= 4.625029e-07
                                     50
                                           46
                                                7.390688e-10
## ig= 47
            gnorm= 2.118326e-07
                                     51
                                           47
                                                2.535775e-10
            gnorm= 9.601958e-08
## ig= 48
                                     52
                                           48
                                                8.657756e-11
                                                2.912793e-11
## ig= 49
            gnorm= 4.284391e-08
                                     53
                                           49
## ig= 50
            gnorm= 1.902241e-08
                                     54
                                           50
                                                9.993603e-12
## ig= 51
            gnorm= 9.725521e-09
                                           51
                                                4.491789e-12
                                                3.449057e-12
## ig= 52
            gnorm= 7.962036e-09
                                     56
                                           52
## ig= 53
            gnorm= 7.962845e-09
                                     57
                                           53
                                                3.318037e-12
## ig= 54
            gnorm= 7.960273e-09
                                     58
                                           54
                                                3.23399e-12
## ig= 55
            gnorm= 7.803206e-09
                                                2.976418e-12
                                                2.469253e-12
## ig= 56
            gnorm= 7.162424e-09
                                     60
                                           56
            gnorm= 5.438652e-09
                                     61
                                           57
                                                1.602454e-12
## ig= 57
## ig= 58
            gnorm= 3.141714e-09
                                     62
                                          58
                                                7.572872e-13
## ig= 59
            gnorm= 1.42477e-09
                                     63
                                          59
                                               2.69848e-13
                                                1.712267e-13
## ig= 60
            gnorm= 8.819687e-10
                                     64
                                           60
                                                1.482642e-13
## ig= 61
            gnorm= 7.963541e-10
                                     65
                                           61
            gnorm= 3.858799e-10
                                     66
                                           62
                                                6.156006e-14
## ig= 62
## ig= 63
            gnorm= 3.35396e-10
                                     67
                                          63
                                               5.127169e-14
                                                4.925589e-14
## ig= 64
            gnorm= 3.318282e-10
                                     68
                                           64
            gnorm= 3.301397e-10
## ig= 65
                                     69
                                           65
                                                4.854075e-14
                                     70
## ig= 66
            gnorm= 3.282529e-10
                                           66
                                                4.83081e-14
## ig= 67
            gnorm= 3.268454e-10
                                     71
                                           67
                                                4.818537e-14
## ig= 68
            gnorm= 3.231113e-10
                                     72
                                           68
                                                4.778396e-14
            gnorm= 3.170085e-10
## ig= 69
                                     73
                                           69
                                                4.68832e-14
## ig= 70
            gnorm= 3.048951e-10
                                     74
                                           70
                                                4.445086e-14
## ig= 71
            gnorm= 2.813282e-10
                                     75
                                           71
                                                3.868174e-14
## ig= 72
            gnorm= 2.291848e-10
                                     76
                                           72
                                                2.716048e-14
## ig= 73
            gnorm= 1.492394e-10
                                     77
                                           73
                                                1.537597e-14
## ig= 74
            gnorm= 1.105431e-10
                                     78
                                           74
                                                1.139537e-14
## ig= 75
            gnorm= 8.616058e-11
                                     79
                                           75
                                                8.874215e-15
            gnorm= 8.099898e-11
                                     80
                                           76
                                                8.249677e-15
## ig= 76
```

```
## ig= 77
            gnorm= 7.976152e-11
                                         77
                                              8.077251e-15
                                   81
## ig= 78
            gnorm= 7.766164e-11
                                    82
                                         78
                                              7.881404e-15
## ig= 79
            gnorm= 7.70462e-11
                                   83
                                        79
                                             7.859951e-15
## ig= 80
            gnorm= 7.675405e-11
                                   84
                                        80
                                              7.855574e-15
## ig= 81
            gnorm= 7.66912e-11
                                   85
                                        81
                                             7.855309e-15
## ig= 82
            gnorm= 7.66865e-11
                                   86
                                        82
                                             7.855299e-15
## ig= 83
            gnorm= 7.668288e-11
                                    87
                                        83
                                              7.855286e-15
                                        84
## ig= 84
            gnorm= 7.667674e-11
                                    88
                                              7.85525e-15
            gnorm= 7.666696e-11
## ig= 85
                                    89
                                        85
                                              7.855158e-15
## ig= 86
            gnorm= 7.665079e-11
                                    90
                                        86
                                             7.854915e-15
## ig= 87
            gnorm= 7.66242e-11
                                   91
                                        87
                                             7.854281e-15
## ig= 88
            gnorm= 7.657974e-11
                                    92
                                        88
                                              7.852621e-15
            gnorm= 7.650429e-11
## ig= 89
                                    93
                                        89
                                              7.848278e-15
## ig= 90
            gnorm= 7.637298e-11
                                    94
                                        90
                                              7.836918e-15
## ig= 91
            gnorm= 7.613642e-11
                                    95
                                        91
                                              7.807274e-15
## ig= 92
            gnorm= 7.568786e-11
                                    96
                                         92
                                              7.730285e-15
## ig= 93
            gnorm= 7.47625e-11
                                   97
                                        93
                                             7.532918e-15
## ig= 94
            gnorm= 7.256457e-11
                                    98
                                             7.044139e-15
                                         95
## ig= 95
            gnorm= 6.656402e-11
                                    99
                                              5.941821e-15
            gnorm= 5.190274e-11
## ig= 96
                                    100
                                         96
                                              3.998643e-15
## ig= 97
            gnorm= 3.075193e-11
                                    101
                                          97
                                               1.930519e-15
                                              6.957847e-16
## ig= 98
            gnorm= 1.42663e-11
                                   102
                                         98
## ig= 99
            gnorm= 8.913349e-12
                                    103
                                          99
                                               4.164659e-16
## ig= 100
             gnorm= 7.819694e-12
                                     104
                                           100
                                                 3.507575e-16
## ig= 101
             gnorm= 2.62496e-12
                                    105
                                          101
                                                8.293119e-17
## ig= 102
             gnorm= 1.244356e-12
                                     106
                                           102
                                                 3.12269e-17
## ig= 103
                                     107
                                           103
             gnorm= 5.255914e-13
                                                 1.02155e-17
## ig= 104
             gnorm= 2.448042e-13
                                     108
                                           104
                                                 3.777248e-18
                                     109
## ig= 105
             gnorm= 1.272943e-13
                                           105
                                                 1.521749e-18
## ig= 106
                                     110
                                           106
                                                 7.280223e-19
             gnorm= 8.110499e-14
## ig= 107
             gnorm= 5.991518e-14
                                     111
                                           107
                                                 4.155754e-19
## ig= 108
             gnorm= 4.6291e-14
                                   112
                                         108
                                               2.68634e-19
## ig= 109
             gnorm= 3.383621e-14
                                     113
                                           109
                                                 1.725135e-19
## ig= 110
                                                 9.072173e-20
             gnorm= 1.924103e-14
                                     114
                                           110
## ig= 111
                                     115
                                                 8.996756e-20
             gnorm= 1.777681e-14
                                           111
## ig= 112
             gnorm= 1.809286e-14
                                     116
                                           112
                                                 8.813619e-20
## ig= 113
             gnorm= 1.807931e-14
                                     117
                                           113
                                                 8.533112e-20
## ig= 114
             gnorm= 1.71003e-14
                                    118
                                          114
                                                7.402365e-20
                                                 5.448504e-20
## ig= 115
             gnorm= 1.410577e-14
                                     119
                                           115
## ig= 116
                                   Seem to be done Rvmminu
             gnorm= 8.747156e-15
sol1
## $par
## [1] 0.9999996 2.0000006 2.9999941 4.0000130
## $value
## [1] 2.941554e-20
##
## $counts
```

function gradient

116

120

\$convergence

[1] 2

##

##

```
##
```

- ## \$message
- ## [1] "Rvmminu appears to have converged"
- ?? Do we want to try discontinuity at solution? Discontinuous function value?

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