nlmrt-vignette

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November 16, 2024

Background

This vignette discusses the R package nlmrt, that aims to provide computationally robust tools for nonlinear least squares problems. Note that R already has the nls() function to solve nonlinear least squares problems, and this function has a large repertoire of tools for such problems. However, it is specifically NOT indicated for problems where the residuals are small or zero. Furthermore, it frequently fails to find a solution if starting parameters are provided that are not close enough to a solution. The tools of nlmrt are very much intended to cope with both these issues.

The functions are also intended to provide stronger support for bounds constraints and to introduce the capability for masks, that is, parameters that are xed for a given run of the function.

nlmrt tools generally do not return the large nls-style object. However, we do provide a tool wrapnls that will run either nlxb followed by a call to nls. The call to nls is adjusted to use the port algorithm if there are bounds constraints.

1 An example problem and its solution

Let us try an example initially presented by [5] and developed by [2]. This is a model for the regrowth of pasture. We set up the computation by putting the data for the problem in a data frame, and specifying the formula for the model. This can be as a formula object, but I have found that saving it as a character string seems to give fewer difficulties. Note the " " that implies "is modeled by". There must be such an element in the formula for this package (and for $nls()$). We also specify two sets of starting parameters, that is, the ones which is a trivial (but possibly unsuitable) start with all parameters set to 1, and huetstart which was suggested in [2]. Finally we load the routines in the package nlmrt.

```
> options(width=60)
```
- > pastured <- data.frame(
- + time=c(9, 14, 21, 28, 42, 57, 63, 70, 79),

```
+ yield= c(8.93, 10.8, 18.59, 22.33, 39.35,
+ 56.11, 61.73, 64.62, 67.08))
> regmod <- "yield \tilde{ } t1 - t2*exp(-exp(t3+t4*log(time)))"
> ones <- c(t1=1, t2=1, t3=1, t4=1) # all ones start
> huetstart <- c(t1=70, t2=60, t3=0, t4=1)
> require(nlmrt)
```
Let us now call the routine nlsmnqb (even though we are not specifying bounds). We try both starts.

```
> anmrt <- nlxb(regmod, start=ones, trace=FALSE, data=pastured)
> print(anmrt)
nlmrt class object: x
residual sumsquares = 4648.1 on 9 observations
   after 3 Jacobian and 4 function evaluations
 name coeff SE tstat pval gradient JSingval
t1 38.8378 NA NA NA -2.95e-11 3
t2 1.00007 NA NA NA -7.748e-10 1.437e-09
t3 0.998202 NA NA NA 1.889e-08 2.275e-16
t4 0.996049 NA NA NA 4.15e-08 5.091e-26
> anmrtx <- try(nlxb(regmod, start=huetstart, trace=FALSE, data=pastured))
> print(strwrap(anmrtx))
 [1] "c(0.480575683702355, 0.669264006079278,"
 [2] "-2.28426563497321, 0.843862687207526,"
 [3] "0.734652618487651, 0.0665106492952035,"
 [4] "-0.985862291968047, -0.0250879549066241,"
 [5] "0.500350456693468)"
 [6] "c(1, 1, 1, 1, 1, 1, 1, 1, 1, -0.981556726091093,"
 [7] "-0.948171282599528, -0.869750270888725,"
 [8] "-0.758399834057041, -0.484261107837458,"
 [9] "-0.223408951427347, -0.149363030476155,"
[10] "-0.0869332933121748, -0.0385258954067767,"
[11] "1.12712321032758, 3.11275223693948, 7.48692917929384,"
[12] "12.9373484175605, 21.6609765596451, 20.6543768151933,"
[13] "17.5183401160426, 13.0985419560087, 7.73883739451391,"
[14] "2.47654281941786, 8.21473160617142, 22.7941238760063,"
[15] "43.1098907467032, 80.9615739893338, 83.5067043689998,"
[16] "72.5808432835138, 55.649093177887, 33.8144464340503)"
[17] "44"
[18] "32"
[19] "c(t1 = 69.9553722026373, t2 = 61.6818319271118, t3 ="
[20] "-9.20880204813341, t4 = 2.37778402563408)"
[21] "8.37588360361963"
[22] "c(-Inf, -Inf, -Inf, -Inf)"
```

```
[23] "c(Inf, Inf, Inf, Inf)"
[24] "integer(0)"
```
Note that the standard $nls()$ of R fails to find a solution from either start.

```
> anls <- try(nls(regmod, start=ones, trace=FALSE, data=pastured))
  > print(strwrap(anls))
[1] "Error in nlsModel(formula, mf, start, wts,"
[2] "scaleOffset = scOff, nDcentral = nDcntr) : singular"
[3] "gradient matrix at initial parameter estimates'
  > anlsx <- try(nls(regmod, start=huetstart, trace=FALSE, data=pastured))
  > print(strwrap(anlsx))
```

```
[1] "Error in nls(regmod, start = huetstart, trace ="
[2] "FALSE, data = pastured) : singular gradient"
```
In both cases, the nls() failed with a 'singular gradient'. This implies the Jacobian is effectively singular at some point. The Levenberg-Marquardt stabilization used in nlxb avoids this particular issue by augmenting the Jacobian until it is non-singular. The details of this common approach may be found elsewhere [4, Algorithm 23].

There are some other tools for R that aim to solve nonlinear least squares problems. We have not yet been able to successfully use the INRA package nls2. This is a quite complicated package and is not installable as a regular R package using install.packages(). Note that there is a very different package by the same name on CRAN by Gabor Grothendieck.

2 The nls solution

We can call nls after getting a potential nonlinear least squares solution using nlxb. Package nlmrt has function wrapnls to allow this to be carried out automatically. Thus,

```
> awnls <- wrapnls(regmod, start=ones, data=pastured, control=list(rofftest=FALSE))
  > print(awnls)
Nonlinear regression model
 model: yield ~ t1 - t2 * exp(-exp(t3 + t4 * log(time)))
  data: data
t1 t2 t3 t4
69.96 61.68 -9.21 2.38
residual sum-of-squares: 8.38
Number of iterations to convergence: 0
Achieved convergence tolerance: 7.91e-08
```
> cat("Note that the above is just the nls() summary result.\n") Note that the above is just the nls() summary result.

3 Problems specified by residual functions

The model expressions in R , such as

yield \sim t1 - t2*exp(-exp(t3+t4*log(time)))

are an extremely helpful feature of the language. Moreover, they are used to compute symbolic or automatic derivatives, so we do not have to rely on numerical approximations for the Jacobian of the nonlinar least squares problem. However, there are many situations where the expression structure is not flexible enough to allow us to define our residuals, or where the construction of the residuals is simply too complicated. In such cases it is helpful to have tools that work with R functions.

Once we have an R function for the residuals, we can use the safeguarded Marquardt routine nlfb from package nlmrt or else the routine nls.lm from package minpack.lm [1]. The latter is built on the Minpack Fortran codes of [3] implemented by Kate Mullen. nlfb is written entirely in R , and is intended to be quite aggessive in ensuring it finds a good minimum. Thus these two approaches have somewhat different characteristics.

Let us consider a slightly different problem, called WEEDS. Here the objective is to model a set of 12 data points (density y of weeds at annual time points tt) versus the time index. (A minor note: use of t rather than tt in R may encourage confusion with the transpose function $t()$, so I tend to avoid plain t.) The model suggested was a 3-parameter logistic function,

 $y_{model} = b_1/(1 + b_2 exp(-b_3 t))$

and while it is possible to use this formulation, a scaled version gives slightly better results

 $y_{model} = 100b_1/(1+10b_2exp(-0.1b_3tt))$

The residuals for this latter model (in form "model" minus "data") are coded in R in the following code chunk in the function shobbs.res. We have also coded the Jacobian for this model as shobbs.jac

```
> shobbs.res \leq function(x){ # scaled 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)
+ tt <- 1:12
+ res <- 100.0*x[1]/(1+x[2]*10.*exp(-0.1*x[3]*tt)) - y
+ }
> shobbs.jac \le function(x) { # scaled Hobbs weeds problem -- Jacobian
+ jj <- matrix(0.0, 12, 3)
+ tt <- 1:12
+ yy <- exp(-0.1*x[3]*tt) # We don't need data for the Jacobian
+ zz <- 100.0/(1+10.*x[2]*yy)
+ jj[tt,1] <- zz
+ jj[tt,2] <- -0.1*x[1]*zz*zz*yy
+ jj[tt,3] <- 0.01*x[1]*zz*zz*yy*x[2]*tt
```
+ return(jj) + }

With package nlmrt, function nlfb can be used to estimate the parameters of the WEEDS problem as follows, where we use the naive starting point where all parameters are 1.

```
> st <- c(b1=1, b2=1, b3=1)
  > ans1 <- nlfb(st, shobbs.res, shobbs.jac, trace=FALSE)
  > print(ans1)
nlmrt class object: x
residual sumsquares = 2.5873 on 12 observations
            Jacobian and 14 function evaluations<br>coeff 5E tstat p
 name coeff SE tstat pval gradient JSingval
b1 1.96186 0.1131 17.35 3.166e-08 -7.327e-06 130.1
                                 29.08 3.282e-10
b3 3.1357 0.06863 45.69 5.768e-12 1.717e-06 2.735
```
This works very well, with almost identical iterates as given by nlxb. (Since the algorithms are the same, this should be the case.) Note that we turn o the trace output. There is also the possibility of interrupting the iterations to watch the progress. Changing the value of watch in the call to nlfb below allows this. In this code chunk, we use an internal numerical approximation to the Jacobian.

```
> cat("No jacobian function -- use internal approximation\n")
No jacobian function -- use internal approximation
  > ans1n <- nlfb(st, shobbs.res, trace=FALSE, control=list(watch=FALSE)) # NO jacfn
  > print(ans1n)
nlmrt class object: x
residual sumsquares = 2.5873 on 12 observations
             Jacobian and 14 function evaluations<br>coeff SE tstat pval
name coeff SE tstat pval gradient JSingval<br>b1 1.96186 0.1131 17.35 3.166e-08 -7.327e-06 130.1
b1 1.96186 0.1131 17.35 3.166e-08 -7.327e-06 130.1
b2 4.90916 0.1688 29.08 3.282e-10 1.429e-07 6.165
b3 3.1357 0.06863 45.69 5.768e-12 1.719e-06 2.735
```
Note that we could also form the sum of squares function and the gradient and use a function minimization code. The next code block shows how this is done, creating the sum of squares function and its gradient, then using the optimx package to call a number of minimizers simultaneously.

```
> shobbs.f \leq function(x){
+ res <- shobbs.res(x)
+ as.numeric(crossprod(res))
+ }
> shobbs.g \le function(x){
+ res <- shobbs.res(x) # This is NOT efficient -- we generally have res already calculations.
+ JJ <- shobbs.jac(x)
+ 2.0*as.vector(crossprod(JJ,res))
+ }
```


> cat("\nNow with numerical gradient approximation or derivative free methods\n") Now with numerical gradient approximation or derivative free methods

> aopxn <- optimx(st, shobbs.f, control=list(all.methods=TRUE))

nmkb NA 9999 NA NA 0.000 NA 0.000

We see that most of the minimizers work with either the analytic or approximated gradient. The 'CG' option of function optim() does not do very well in either case. As the author of the original step and description and then Turbo Pascal code, I can say I was never very happy with this method and replaced it recently with Rcgmin from the package of the same name, in the process adding the possibility of bounds or masks constraints.

4 Converting an expression to a function

Clearly if we have an expression, it would be nice to be able to automatically convert this to a function, if possible also getting the derivatives. Indeed, it is possible to convert an expression to a function, and there are several ways to do this (references??). In package nlmrt we provide the tools model2grfun.R, model2jacfun.R, model2resfun.R, and model2ssfun.R to convert a model expression to a function to compute the gradient, Jacobian, residuals or sum of squares functions respectively. We do not provide any tool for converting a function for the residuals back to an expression, as functions can use structures that are not easily expressed as R expressions.

Below are code chunks to illustrate the generation of the residual, sum of squares, Jacobian and gradient code for the Ratkowsky problem used earlier in the vignette. The commented-out first line shows how we would use one of these function generators to output the function to a file named "testresfn. R ". However, it is not necessary to generate the file.

First, let us generate the residuals. We must supply the names of the parameters, and do this via the starting vector of parameters ones. The actual values are not needed by model2resfun, just the names. Other names are drawn from the variables used in the model expression regmod.

```
> # jres <- model2resfun(regmod, ones, funname="myxres", file="testresfn.R")
  > jres <- model2resfun(regmod, ones)
  > print(jres)
function (prm, yield = NULL, time = NULL)
{
   t1 <- prm[[1]]
   t2 <- prm[[2]]
   t3 <- prm[[3]]
   t4 <- prm[[4]]
   resids \leftarrow as.numeric(eval(t1 - t2 * exp(-exp(t3 + t4 * log(time))) -
      yield))
}
<environment: 0x55ec8891fef8>
  > valjres <- jres(ones, yield=pastured$yield, time=pastured$time)
  > cat("valjres:")
valjres:
  > print(valjres)
[1] -7.93 -9.80 -17.59 -21.33 -38.35 -55.11 -60.73 -63.62
[9] -66.08
```
Now let us also generate the Jacobian and test it using the numerical approximations from package numDeriv.

```
> jjac <- model2jacfun(regmod, ones)
   > print(jjac)
function (prm, yield = NULL, time = NULL)
{
    t1 <- prm[[1]]
    t2 <- prm[[2]]
     t3 <- prm[[3]]
t4 <- prm[[4]]
    localdf <- data.frame(yield, time)
     jstruc <- with(localdf, eval({
.expr1 <- log(time)
         .expr4 <- exp(t3 + t4 * .expr1)
.expr6 <- exp(-.expr4)
         value < -t1 - t2 * .expr6 - yield
         .grad <- array(0, c(length(.value), 4), list(NULL, c("t1",<br>"t2", "t3", "t4")))<br>.grad[, "t1"] <- 1<br>.grad[, "t2"] <- -.expr6
         .grad[, "t3"] <- t2 * (.expr6 * .expr4)
         .grad[, "t4"] <- t2 * (.expr6 * (.expr4 * .expr1))
        attr(.value, "gradient") <- .grad
        .value
    }))
    jacmat <- attr(jstruc, "gradient")
    return(jacmat)
}
<environment: 0x55ec887e6f50>
   > # Note that we now need some data!
   > valjjac <- jjac(ones, yield=pastured$yield, time=pastured$time)
   > cat("valjac:")
valjac:
   > print(valjjac)
 t1 t2 t3 t4
[1,] 1 -2.3724e-11 5.8040e-10 1.2753e-09
 [2,] 1 -2.9683e-17 1.1296e-15 2.9812e-15
 [3,] 1 -1.6172e-25 9.2317e-24 2.8106e-23
[4,] 1 -8.8110e-34 6.7062e-32 2.2347e-31
 [5,] 1 -2.6154e-50 2.9859e-48 1.1160e-47
[6,] 1 -5.1229e-68 7.9375e-66 3.2092e-65
 [7,] 1 -4.2297e-75 7.2434e-73 3.0010e-72
 [8,] 1 -2.3044e-83 4.3849e-81 1.8629e-80
[9,] 1 -5.4670e-94 1.1740e-91 5.1298e-91
   > # Now compute the numerical approximation
   > require(numDeriv)
   > Jn <- jacobian(jres, ones, , yield=pastured$yield, time=pastured$time)
   > cat("maxabsdiff=",max(abs(Jn-valjjac)),"\n")
maxabsdiff= 3.7744e-10
```
As with the WEEDS problem, we can compute the sum of squares function and the gradient.

```
> ssfn <- model2ssfun(regmod, ones) # problem getting the data attached!
  > print(ssfn)
function (prm, yield = NULL, time = NULL)
{
```

```
t1 <- prm[[1]]
    t2 < - \text{prm} [[2]]
    t3 <- prm[[3]]
    t4 < - \text{prm}[[4]]
    resids \leftarrow as.numeric(eval(t1 - t2 * exp(-exp(t3 + t4 * log(time))) -
    yield))
ss <- as.numeric(crossprod(resids))
}
<environment: 0x55ec882689f8>
   > valss <- ssfn(ones, yield=pastured$yield, time=pastured$time)
   > cat("valss: ",valss,"\n")
valss: 17533
   > grfn <- model2grfun(regmod, ones) # problem getting the data attached!
   > print(grfn)
function (prm, yield = NULL, time = NULL)
{
    t1 <- prm[[1]]
    t2 <- prm[[2]]
    t3 <- prm[[3]]
t4 <- prm[[4]]
    localdf <- data.frame(yield, time)
    jstruc <- with(localdf, eval({
        .expr1 <- log(time)
        .expr4 \leftarrow exp(t3 + t4 * .expr1)
        .expr6 <- exp(-.expr4)
        .value <- t1 - t2 * .expr6 - yield
        .grad <- array(0, c(length(.value), 4), list(NULL, c("t1",
"t2", "t3", "t4")))
.grad[, "t1"] <- 1
        .grad[, "t2"] <- -.expr6
.grad[, "t3"] <- t2 * (.expr6 * .expr4)
        .grad[, "t4"] <- t2 * (.expr6 * (.expr4 * .expr1))
        attr(.value, "gradient") <- .grad
        .value
    }))
    jacmat <- attr(jstruc, "gradient")
    resids \leftarrow as.numeric(eval(t1 - t2 * exp(-exp(t3 + t4 * log(time))) -
       yield))
    grj <- as.vector(2 * crossprod(jacmat, resids))
}
<environment: 0x55ec882326a8>
   > valgr <- grfn(ones, yield=pastured$yield, time=pastured$time)
   > cat("valgr:")
valgr:
   > print(valgr)
[1] -6.8108e+02 3.7626e-10 -9.2051e-09 -2.0226e-08
   > gn <- grad(ssfn, ones, yield=pastured$yield, time=pastured$time)
   > cat("maxabsdiff=",max(abs(gn-valgr)),"\n")
maxabsdiff= 1.1034e-07
```
Moreover, we can use the Huet starting parameters as a double check on our conversion of the expression to various optimization-style functions.

```
> cat("\n\nHuetstart:")
Huetstart:
  > print(huetstart)
t1 t2 t3 t4
70 60 0 1
```

```
> valjres <- jres(huetstart, yield=pastured$yield, time=pastured$time)
  > cat("valjres:")
valjres:
  > print(valjres)
[1] 61.063 59.200 51.410 47.670 30.650 13.890 8.270 5.380
[9] 2.920
  > valss <- ssfn(huetstart, yield=pastured$yield, time=pastured$time)
  > cat("valss:", valss, "\n\times")
valss: 13387
  > valjjac <- jjac(huetstart, yield=pastured$yield, time=pastured$time)
  > cat("valjac:")
valjac:
  > print(valjjac)
 t1 t2 t3 t4
[1,] 1 -1.2341e-04 6.6641e-02 1.4643e-01
 [2,] 1 -8.3153e-07 6.9848e-04 1.8433e-03
 [3,] 1 -7.5826e-10 9.5540e-07 2.9087e-06
 [4,] 1 -6.9144e-13 1.1616e-09 3.8708e-09
 [5,] 1 -5.7495e-19 1.4489e-15 5.4154e-15
 [6,] 1 -1.7588e-25 6.0151e-22 2.4319e-21
[7,] 1 -4.3596e-28 1.6479e-24 6.8276e-24
 [8,] 1 -3.9754e-31 1.6697e-27 7.0937e-27
 [9,] 1 -4.9061e-35 2.3255e-31 1.0161e-30
  > Jn <- jacobian(jres, huetstart, , yield=pastured$yield, time=pastured$time)
  > cat("maxabsdiff=",max(abs(Jn-valjjac)),"\n")
maxabsdiff= 5.3945e-10
  > valgr <- grfn(huetstart, yield=pastured$yield, time=pastured$time)
  > cat("valgr:")
valgr:
  > print(valgr)
[1] 560.90509 -0.01517 8.22138 18.10084
  > gn <- grad(ssfn, huetstart, yield=pastured$yield, time=pastured$time)
  > cat("maxabsdiff=",max(abs(gn-valgr)),"\n")
maxabsdiff= 6.0068e-08
```
Now that we have these functions, let us apply them with nlfb.

```
> cat("All ones to start\n")
All ones to start
  > anlfb <- nlfb(ones, jres, jjac, trace=FALSE, yield=pastured$yield, time=pastured$time)
  > print(strwrap(anlfb))
 [1] "c(29.9077777777472, 28.0377777777778,"
 [2] "20.2477777777778, 16.5077777777778,"
 [3] "-0.512222222222185, -17.2722222222222,"
 [4] "-22.8922222222222, -25.7822222222222,"
 [5] "-28.2422222222222)"
 [6] "c(1, 1, 1, 1, 1, 1, 1, 1, 1, -2.5904803198541e-11,"
 [7] "-3.48177832682637e-17, -2.11977026263411e-25,"
 [8] "-1.30186504324169e-33, -5.00031799754126e-50,"
 [9] "-1.31593194786314e-67, -1.22799624106577e-74,"
[10] "-7.73807145709702e-83, -2.22164181475742e-93,"
[11] "6.31486798422103e-10, 1.31950211229137e-15,"
[12] "1.20434229540344e-23, 9.8581678893932e-32,
[13] "5.67649212681333e-48, 2.02656873886828e-65,"
```

```
[14] "2.0899294411951e-72, 1.46306184917541e-80,"
[15] "4.73981859866247e-91, 1.38751831375641e-09,"
[16] "3.48224172088535e-15, 3.66664714105511e-23,"
[17] "3.28494315031004e-31, 2.12168521608149e-47,"
[18] "8.19352130903529e-65, 8.65885924352346e-72,"
[19] "6.21581130504567e-80, 2.0710390197009e-90)"
[20] "4"
[21] "3"
[22] "c(t1 = 38.837777777778, t2 = 1.00007369903283, t3 ="
\begin{bmatrix} 23 \\ 23 \end{bmatrix} "0.998201661261902, t4 = 0.996048644398237)'
[24] "4648.06335555373"
[25] "c(-Inf, -Inf, -Inf, -Inf)'[26] "c(Inf, Inf, Inf, Inf)"
[27] "NULL"
  > cat("Huet start\n")
Huet start
  > anlfbh <- nlfb(huetstart, jres, jjac, trace=FALSE, yield=pastured$yield, time=pastured$t
  > print(strwrap(anlfbh))
 [1] "c(0.480575683702348, 0.669264006079271,"
 [2] "-2.28426563497322, 0.843862687207512,"
[3] "0.734652618487637, 0.0665106492951892,"
 [4] "-0.985862291968061, -0.0250879549066383,"
 [5] "0.500350456693454)"
 [6] "c(1, 1, 1, 1, 1, 1, 1, 1, 1, -0.981556726091093,"
 [7] "-0.948171282599528, -0.869750270888724,"
 [8] "-0.758399834057038, -0.484261107837453,"
[9] "-0.223408951427342, -0.14936303047615,"
[10] "-0.0869332933121714, -0.0385258954067745,"
[11] "1.12712321032758, 3.1127522369395, 7.4869291792939,"
[12] "12.9373484175606, 21.6609765596452, 20.6543768151931,"
[13] "17.5183401160423, 13.0985419560084, 7.73883739451359,"
[14] "2.47654281941787, 8.21473160617148, 22.7941238760065,"
[15] "43.1098907467036, 80.9615739893341, 83.5067043689991,"
[16] "72.5808432835127, 55.6490931778857, 33.8144464340489)"
[17] "44"
[18] "32"
[19] "c(t1 = 69.9553722026373, t2 = 61.6818319271118, t3 ="
[20] "-9.20880204813341, t4 = 2.37778402563408)"
[21] "8.37588360361962"
[22] "c(-Inf, -Inf, -Inf, -Inf)'
[23] "c(Inf, Inf, Inf, Inf)"
[24] "NULL"
```
5 Using bounds and masks

The manual for nls() tells us that bounds are restricted to the 'port' algorithm.

lower, upper: vectors of lower and upper bounds, replicated to be as long as 'start'. If unspecified, all parameters are assumed to be unconstrained. Bounds can only be used with the '"port"' algorithm. They are ignored, with a warning, if given for other algorithms.

Later in the manual, there is the discomforting warning:

The 'algorithm = "port"' code appears unfinished, and does not even check that the starting value is within the bounds. Use with caution, especially where bounds are supplied.

We will base the rest of this discussion on the examples in man/nlmrtpackage.Rd, and use an unscaled version of the WEEDS problem.

First, let us estimate the model with no constraints.

```
> require(nlmrt)
  > # Data for Hobbs problem
  > ydat <- 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)
  > tdat <- 1:length(ydat)
  > weeddata1 <- data.frame(y=ydat, tt=tdat)
  > start1 <- c(b1=1, b2=1, b3=1) # name parameters for nlxb, nls, wrapnls.
  > eunsc <- y \sim b1/(1+b2*exp(-b3*tt))> anlxb1 <- try(nlxb(eunsc, start=start1, data=weeddata1))
  > print(anlxb1)
nlmrt class object: x
residual sumsquares = 2.5873 on 12 observations
   after 18 Jacobian and 25 function evaluations<br>ame coeff SE tstat pro
name coeff SE tstat pval gradient JSingval
المستوفى السبب المستوفى المست<br>196.186 1.688 29.08 3.281e-10 1.59e-07 0.4605<br>197.08 3.281e-10 1.59e-07 0.4605
                                      29.08 3.281e-10b3 0.31357 0.006863 45.69 5.768e-12 -5.531e-05 0.04715
```
Now let us see if we can apply bounds. Note that we name the parameters in the vectors for the bounds. First we apply bounds that are NOT active at the unconstrained solution.

```
> # WITH BOUNDS
  > startf1 <- c(b1=1, b2=1, b3=.1) # a feasible start when b3 <= 0.25> anlxb1 <- try(nlxb(eunsc, start=startf1, lower=c(b1=0, b2=0, b3=0),
  + upper=c(b1=500, b2=100, b3=5), data=weeddata1))
  > print(anlxb1)
nlmrt class object: x
residual sumsquares = 2.5873 on 12 observations
 after 13 Jacobian and 18 function evaluations<br>name coeff SE tstat p
name coeff SE tstat pval gradient JSingval<br>b1 196.186 11.31 17.35 3.164e-08 -2.662e-07 1011
b1 196.186 11.31 17.35 3.164e-08 -2.662e-07 1011
b2 49.0916 1.688 29.08 3.281e-10 1.648e-07 0.4605
                                     b3 0.31357 0.006863 45.69 5.768e-12 -5.872e-05 0.04715
```
We note that $nls()$ also solves this case.

```
> anlsb1 \leq try(nls(eunsc, start=startf1, lower=c(b1=0, b2=0, b3=0),
          upper=c(b1=500, b2=100, b3=5), data=weeddata1, algorithm='port'))
  > print(anlsb1)
Nonlinear regression model
 model: y \sim b1/(1 + b2 * exp(-b3 * tt))
  data: weeddata1<br>b1 b2
          b2 b3<br>092 0.314196.186 49.092
residual sum-of-squares: 2.59
```
Algorithm "port", convergence message: relative convergence (4)

Now we will change the bounds so the start is infeasible.

```
> ## Uncon solution has bounds ACTIVE. Infeasible start
  > anlxb2i <- try(nlxb(eunsc, start=start1, lower=c(b1=0, b2=0, b3=0),
  + upper=c(b1=500, b2=100, b3=.25), data=weeddata1))
  > print(anlxb2i)
[1] "Error in nlxb(eunsc, start = start1, lower = c(b1 = 0, b2 = 0, b3 = 0), : \n Infeasible start\n"
attr(,"class")
[1] "try-error"
attr(,"condition")
\sinh\theta = \sinh\theta (sunsc, start = start1, lower = c(b1 = 0, b2 = 0, b3 = 0), upper = c(b1 = 500, b2 = 100, b3 = 0.25), data
  > anlsb2i <- try(nls(eunsc, start=start1, lower=c(b1=0, b2=0, b3=0),
  + upper=c(b1=500, b2=100, b3=.25), data=weeddata1, algorithm='port'))
  > print(anlsb2i)
[1] "Error in nls(eunsc, start = start1, lower = c(b1 = 0, b2 = 0, b3 = 0), : \n Convergence failure: initial par violates cons
attr(,"class")
[1] "try-error"
attr(,"condition")
\langlesimpleError in nls(eunsc, start = start1, lower = c(b1 = 0, b2 = 0, b3 = 0), upper = c(b1 = 500, b2 = 100, b3 = 0.25), data
```
Both nlxb() and nls() (with 'port') do the right thing and refuse to proceed. There is a minor "glitch" in the output processing of both knitR and Sweave here. Let us start them off properly and see what they accomplish.

```
> ## Uncon solution has bounds ACTIVE. Feasible start
  > anlxb2f <- try(nlxb(eunsc, start=startf1, lower=c(b1=0, b2=0, b3=0),
 + upper=c(b1=500, b2=100, b3=.25), data=weeddata1))
  > print(anlxb2f)
nlmrt class object: x
residual sumsquares = 29.993 on 12 observations
            Jacobian and 18 function evaluations<br>coeff 5E tstat pval
name coeff SE tstat pval gradient JSingval<br>b1 500U NA NA NA O 1.529
b1 500U NA NA NA 0 1.529
b2 87.9425 NA NA NA -1.808e-10 0
b3 0.25U NA NA NA 0 0
 > anlsb2f \leq try(nls(eunsc, start=startf1, lower=c(b1=0, b2=0, b3=0),
 + upper=c(b1=500, b2=100, b3=.25), data=weeddata1, algorithm='port'))
  > print(anlsb2f)
Nonlinear regression model<br>model: y \approx b1/(1 + b2) * cb1/(1 + b2 * exp(-b3 * tt))data: weeddata1<br>b1 b2 b3
b1 b2 b3
500.00 87.94 0.25
residual sum-of-squares: 30
```
Algorithm "port", convergence message: both X-convergence and relative convergence (5)

Both methods get essentially the same answer for the bounded problem, and this solution has parameters b1 and b3 at their upper bounds. The Jacobian elements for these parameters are zero as returned by nlxb().

Let us now turn to **masks**, which functions from nlmrt are designed to handle. Masks are also available with packages Rcgmin and Rvmmin. I would like to hear if other packages offer this capability.

```
> ## TEST MASKS
> anlsmnqm <- try(nlxb(eunsc, start=start1, lower=c(b1=0, b2=0, b3=0),
```

```
+ upper=c(b1=500, b2=100, b3=5), masked=c("b2"), data=weeddata1))
  > print(anlsmnqm) # b2 masked
nlmrt class object: x
residual sumsquares = 6181.2 on 12 observations
            Jacobian and 35 function evaluations
name coeff SE tstat pval gradient JSingval<br>b1 50.4013 NA NA NA -0.001511 162.1
b1 50.4013 NA NA NA -0.001511 162.1
b2 1 M NA NA NA 0 0.4918
b3 0.19862 NA NA NA -0.0468 0
  > an1qm3 <- try(nlxb(eunsc, start=start1, data=weeddata1, masked=c("b3")))
  > print(an1qm3) # b3 masked
nlmrt class object: x
residual sumsquares = 1031 on 12 observations
  after 17 Jacobian and 18 function evaluations<br>ame coeff SE tstat pval
 name coeff SE tstat pval gradient JSingval
b1 78.5698 NA NA NA 8.489e-08 1.944
b2 2293.71 NA NA NA -1.757e-09 0.01097
b3 1 M NA NA NA 0 0
  > # Note that the parameters are put in out of order to test code.
  > an1qm123 <- try(nlxb(eunsc, start=start1, data=weeddata1, masked=c("b2","b1","b3")))
  > print(an1qm123) # ALL masked - fails!!
[1] "Error in nlxb(eunsc, start = start1, data = weeddata1, masked = c(\"b2\", : \n All parameters are masked\n"
attr(,"class")
[1] "try-error"
attr(,"condition")
<simpleError in nlxb(eunsc, start = start1, data = weeddata1, masked = c("b2", "b1", "b3")): All parameters are masked>
```
Finally (for nlxb) we combine the bounds and mask.

```
> ## BOUNDS and MASK
 > an1qbm2 <- try(nlxb(eunsc, start=startf1, data=weeddata1,
 + lower=c(0,0,0), upper=c(200, 60, .3), masked=c("b2")))
 > print(an1qbm2)
nlmrt class object: x
residual sumsquares = 6181.2 on 12 observations
          Jacobian and 28 function evaluations<br>coeff 5E tstat pval
name coeff SE tstat pval gradient JSingval<br>b1 50.4016 NA NA NA 0.0004618 162.2
b1 50.4016 NA NA NA 0.0004618 162.2
b2 1 M NA NA NA 0 0.4918
b3 0.198618 NA NA NA -0.0746 0
 > an1qbm2x <- try(nlxb(eunsc, start=startf1, data=weeddata1,
 + lower=c(0,0,0), upper=c(48, 60, .3), masked=c("b2")))
 > print(an1qbm2x)
nlmrt class object: x
residual sumsquares = 6206.1 on 12 observations
 after 11 Jacobian and 20 function evaluations<br>name coeff SE tstat pval
name coeff SE tstat pval gradient JSingval
b1 48U NA NA NA 0 141.2
b2 1 M NA NA NA 0 0
b3 0.215971 NA NA NA -0.1502 0
```
Turning to the function-based nlfb,

> hobbs.res <- function(x){ # Hobbs weeds problem -- residual

- + 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)
  + tt <- 1:12
  + res <- x[1]/(1+x[2]*exp(-x[3]*tt)) - y
  + }
  > hobbs.jac <- function(x) { # Hobbs weeds problem -- Jacobian
  + jj <- matrix(0.0, 12, 3)
  + tt <- 1:12
  + yy <- exp(-x[3]*tt)
  + zz <- 1.0/(1+x[2]*yy)+ jj[tt,1] <- zz
  + jj[tt,2] <- -x[1]*zz*zz*yy
  + jj[tt,3] <- x[1]*zz*zz*yy*x[2]*tt
  + return(jj)
  + }
  > # Check unconstrained
  > ans1 <- nlfb(start1, hobbs.res, hobbs.jac)
  > ans1
nlmrt class object: x
residual sumsquares = 2.5873 on 12 observations
            Jacobian and 25 function evaluations<br>coeff 5E tstat p
name coeff SE tstat pval gradient JSingval
b1 196.186 11.31 17.35 3.164e-08 -2.663e-07 1011
b2 49.0916 1.688 29.08 3.281e-10 1.59e-07 0.4605
b3 0.31357 0.006863 45.69 5.768e-12 -5.531e-05
  > ## No jacobian - use internal approximation
  > ans1n <- nlfb(start1, hobbs.res)
  > ans1n
nlmrt class object: x
residual sumsquares = 2.5873 on 12 observations
            Jacobian and 25 function evaluations<br>coeff 5E tstat pval
name coeff SE tstat pval gradient JSingval<br>b1 196.186 11.31 17.35 3.164e-08 -2.663e-07 1011
b1 196.186 11.31 17.35 3.164e-08 -2.663e-07 1011
b2 49.0916 1.688 29.08 3.281e-10 1.589e-07 0.4605
                                 45.69 5.768e-12 -5.528e-05
  > # Bounds -- infeasible start
  > ans2i <- try(nlfb(start1, hobbs.res, hobbs.jac,
  + lower=c(b1=0, b2=0, b3=0), upper=c(b1=500, b2=100, b3=.25)))
  > ans2i
[1] "Error in nlfb(start1, hobbs.res, hobbs.jac, lower = c(b1 = 0, b2 = 0, : \n Infeasible start\n"
attr(,"class")
[1] "try-error"
attr(,"condition")
\leimpleError in nlfb(start1, hobbs.res, hobbs.jac, lower = c(b1 = 0, b2 = 0, b3 = 0), upper = c(b1 = 500, b2 = 100, b3 = 0.25
  > # Bounds -- feasible start
  > ans2f <- nlfb(startf1, hobbs.res, hobbs.jac,
  + lower=c(b1=0, b2=0, b3=0), upper=c(b1=500, b2=100, b3=.25))
  > ans2f
nlmrt class object: x
residual sumsquares = 29.993 on 12 observations
 after 13 Jacobian and 18 function evaluations<br>name coeff SE tstat pval
name coeff SE tstat pval gradient JSingval
b1 500U NA NA NA 0 1.529
b2 87.9425 NA NA NA -1.809e-10 0
b3 0.25U NA NA NA 0 0
```

```
> # Mask b2
 > ansm2 <- nlfb(start1, hobbs.res, hobbs.jac, maskidx=c(2))
 > ansm2
nlmrt class object: x
residual sumsquares = 6181.2 on 12 observations
          Jacobian and 38 function evaluations<br>coeff 5E tstat p
 name coeff SE tstat pval gradient JSingval
b1 50.4022 NA NA NA 0.001528 162.2
b2 1 M NA NA NA 0 0.4918
b3 0.198611 NA NA NA 0.04544 0
 > # Mask b3
 > ansm3 <- nlfb(start1, hobbs.res, hobbs.jac, maskidx=c(3))
 > ansm3
nlmrt class object: x
residual sumsquares = 1031 on 12 observations
  after 17 Jacobian and 18 function evaluations<br>ame coeff SE tstat pval
 name coeff SE tstat pval gradient JSingval<br>1 78.5698 MA NA NA 8.489e-08 1.94
b1 78.5698 NA NA NA 8.489e-08 1.944
b2 2293.71 NA NA NA -1.757e-09 0.01097
b3 1 M NA NA NA O O
 > # Mask all -- should fail
 > ansma <- try(n1fb(start1, hobbs.res, hobbs.jac, maskidx=c(3,1,2)))> ansma
[1] "Error in nlfb(start1, hobbs.res, hobbs.jac, maskidx = c(3, 1, 2)) : \n All parameters are masked\n"
attr(,"class")
[1] "try-error"
attr(,"condition")
<simpleError in nlfb(start1, hobbs.res, hobbs.jac, maskidx = c(3, 1, 2)): All parameters are masked>
 > # Bounds and mask
  > ansmbm2 <- nlfb(startf1, hobbs.res, hobbs.jac, maskidx=c(2),
 + lower=c(0,0,0), upper=c(200, 60, .3))
  > ansmbm2
nlmrt class object: x
residual sumsquares = 6181.2 on 12 observations
           Jacobian and 28 function evaluations<br>coeff 5E tstat p
 name coeff SE tstat pval gradient JSingval
b1 50.4016 NA NA NA 0.0004618 162.2
b2 1 M NA NA NA 0 0.4918
b3 0.198618 NA NA NA -0.0746 0
 > # Active bound
  > ansmbm2x <- nlfb(startf1, hobbs.res, hobbs.jac, maskidx=c(2),
 + lower=c(0,0,0), upper=c(48, 60, .3))
 > ansmbm2x
nlmrt class object: x
residual sumsquares = 6206.1 on 12 observations
 after 11 Jacobian and 20 function evaluations<br>name coeff SE tstat pval
name coeff SE tstat pval gradient JSingval
b1 48U NA NA NA 0 141.2
b2 1 M NA NA NA O O
b3 0.215971 NA NA NA -0.1502 0
```
The results match those of nlxb()

Finally, let us check the results above with Rvmmin and Rcgmin. Note that this vignette cannot be created on systems that lack these codes.

```
> require(Rcgmin)
  > require(Rvmmin)
  > hobbs.f \leq function(x) {
  + res<-hobbs.res(x)
  + as.numeric(crossprod(res))
  + }
  > hobbs.g \leq function(x) {
  + res <- hobbs.res(x) # Probably already available
  + JJ <- hobbs.jac(x)
  + 2.0*as.numeric(crossprod(JJ, res))
  + }
  > # Check unconstrained
  > a1cg <- Rcgmin(start1, hobbs.f, hobbs.g)
  > a1cg
$par
b1 b2 b3
196.17510 49.09041 0.31358
$value
[1] 2.5873
$counts
[1] 573 195
$convergence
[1] 0
$message
[1] "Rcgmin seems to have converged"
  > a1vm <- Rvmmin(start1, hobbs.f, hobbs.g)
  > a1vm
$par
b1 b2 b3
196.18626 49.09164 0.31357
$value
[1] 2.5873
$counts
function gradient
            81 41
$convergence
[1] 3
$message
[1] "Rvmminu appears to have converged"
  > ## No jacobian - use internal approximation
  > a1cgn <- try(Rcgmin(start1, hobbs.f))
  > a1cgn
[1] "Error in Rcgmin(start1, hobbs.f) : \n Rcgmin must have gradient function provided. Call via optimr() to use approximations.
attr(,"class")
[1] "try-error"
attr(,"condition")
<simpleError in Rcgmin(start1, hobbs.f): Rcgmin must have gradient function provided. Call via optimr() to use approximations.>
  > a1vmn <- try(Rvmmin(start1, hobbs.f))
```

```
> a1vmn
```

```
$par
b1 b2 b3
35.532 35.785 35.204
$value
[1] 9205.4
$counts
function gradient
26 6
$convergence
[1] 0
$message
[1] "Rvmminu appears to have converged"
  > # But
  > grfwd <- function(par, userfn, fbase=NULL, eps=1.0e-7, ...) {
  + # Forward different gradient approximation
  + if (is.null(fbase)) fbase <- userfn(par, ...) # ensure we function value at par
  + df <- rep(NA, length(par))
  + teps <- eps * (abs(par) + eps)
  + for (i in 1:length(par)) {
  + dx <- par
  + dx[i] <- dx[i] + teps[i]
  + df[i] <- (userfn(dx, ...) - fbase)/teps[i]
  + }
  + df
  + }
  > a1vmn <- try(Rvmmin(start1, hobbs.f, gr="grfwd"))
  > a1vmn
$par \b1 b2 b3
35.532 35.785 35.204
$value
[1] 9205.4
$counts
function gradient
   26
$convergence
[1] 0
$message
[1] "Rvmminu appears to have converged"
  > # Bounds -- infeasible start
  > # Note: These codes move start to nearest bound
  > a1cg2i <- Rcgmin(start1, hobbs.f, hobbs.g,
  + lower=c(b1=0, b2=0, b3=0), upper=c(b1=500, b2=100, b3=.25))
Rcgminb: maxfeval set to 10000
  > a1cg2i
$ $\rm{bar}$ $\rm{b1}$b2 b3<br>942 0.250
500.000 87.942
attr(,"status")
```

```
[1] " " " " "U"
$value
[1] 29.993
$counts
[1] 88 45
$convergence
[1] 0
$message
[1] "Rcgmin seems to have converged"
$bdmsk
[1] -1 1 -1
  > a1vm2i <- Rvmmin(start1, hobbs.f, hobbs.g,
  + lower=c(b1=0, b2=0, b3=0), upper=c(b1=500, b2=100, b3=.25))
  > a1vm2i # Fails to get to solution!
\sup<sub>b1</sub>
b1 b2 b3
500.000 87.942 0.250
attr(,"status")
[1] " " " " "U"
$value
[1] 29.993
$counts
function gradient<br>54 24
            54 24
$convergence
[1] 2
$message
[1] "Rvmminb appears to have converged"
$bdmsk
[1] -1 1 -1
  > # Bounds -- feasible start
  > a1cg2f <- Rcgmin(startf1, hobbs.f, hobbs.g,
  + lower=c(b1=0, b2=0, b3=0), upper=c(b1=500, b2=100, b3=.25))
Rcgminb: maxfeval set to 10000
  > a1cg2f
$par \ b1b1 b2 b3
500.000 87.942 0.250
$value
[1] 29.993
$counts
[1] 65 37
$convergence
[1] 0
$message
[1] "Rcgmin seems to have converged"
$bdmsk
\begin{bmatrix} 1 \end{bmatrix} -1 1 -1
```

```
19
```

```
> a1vm2f <- Rvmmin(startf1, hobbs.f, hobbs.g,
  + lower=c(b1=0, b2=0, b3=0), upper=c(b1=500, b2=100, b3=.25))
  > a1vm2f # Gets there, but only just!
$par
b1 b2 b3
500.000 87.942 0.250
$value
[1] 29.993
$counts
function gradient
    86
$convergence
[1] 2
$message
[1] "Rvmminb appears to have converged"
$bdmsk
[1] -1 1 -1
  > # Mask b2
 > a1cgm2 <- Rcgmin(start1, hobbs.f, hobbs.g, bdmsk=c(1,0,1))
Rcgminb: maxfeval set to 10000
  > a1cgm2
$par
b1 b2 b3
50.40178 1.00000 0.19861
$value
[1] 6181.2
$counts
[1] 89 29
$convergence
[1] 0
$message
[1] "Rcgmin seems to have converged"
$bdmsk
[1] 1 0 1
  > a1vmm2 <- Rvmmin(start1, hobbs.f, hobbs.g, bdmsk=c(1,0,1))
  > a1vmm2
\frac{1}{2} \frac{1}{2}b2 b3
50.40179 1.00000 0.19861
$value
[1] 6181.2
$counts
function gradient
             43 14
$convergence
[1] 3
$message
[1] "Rvmminb appears to have converged"
```

```
$bdmsk
[1] 1 0 1
   > # Mask b3
   > a1cgm3 <- Rcgmin(start1, hobbs.f, hobbs.g, bdmsk=c(1,1,0))
Rcgminb: maxfeval set to 10000
   > a1cgm3
\frac{1}{2} \frac{1}{2}b2 b3<br>.947 1.000
  78.571 2293.947
$value
[1] 1031
$counts
[1] 182 75
$convergence
[1] 0
$message
[1] "Rcgmin seems to have converged"
$bdmsk
[1] 1 1 0
   > a1vmm3 <- Rvmmin(start1, hobbs.f, hobbs.g, bdmsk=c(1,1,0))
   > a1vmm3
$par
    b1 b2 b3<br>.571 2293.947 1.00078.571 2293.947
$value
[1] 1031
$counts
function gradient
             39 29
$convergence
[1] 0
$message
[1] "Rvmminb appears to have converged"
$bdmsk
[1] 1 1 0
   > # Mask all -- should fail
   > a1cgma \leftarrow Rcgmin(start1, hobbs.f, hobbs.g, bdmsk=c(0,0,0))
Rcgminb: maxfeval set to 10000
   > a1cgma
$par
b1 b2 b3
1 1 1
$value
[1] 23521
$counts
[1] 1 1
$convergence
[1] 0
```

```
$message
[1] "Rcgmin seems to have converged"
$bdmsk
[1] 0 0 0
  > a1vmma <- Rvmmin(start1, hobbs.f, hobbs.g, bdmsk=c(0,0,0))
  > a1vmma
$par
b1 b2 b3
1 1 1
$value
[1] 23521
$counts
function gradient
    1 1
$convergence
[1] 0
$message
[1] "Rvmminb appears to have converged"
$bdmsk
[1] 0 0 0
  > # Bounds and mask
  > ansmbm2 <- nlfb(startf1, hobbs.res, hobbs.jac, maskidx=c(2),
  + lower=c(0,0,0), upper=c(200, 60, .3))
  > ansmbm2
nlmrt class object: x
residual sumsquares = 6181.2 on 12 observations
 after 17 Jacobian and 28 function evaluations<br>name coeff SE tstat p
name coeff SE tstat pval gradient JSingval<br>b1 50.4016 WA NA 0.0004618 162.2<br>b2
b1 50.4016 NA NA NA 0.0004618 162.2
b2 1 M NA NA NA 0 0.4918
b3 0.198618 NA NA NA -0.0746 0
  > a1cgbm2 <- Rcgmin(start1, hobbs.f, hobbs.g, bdmsk=c(1,0,1),
  + lower=c(0,0,0), upper=c(200, 60, .3))
Rcgminb: maxfeval set to 10000
  > a1cgbm2
$par
b1 b2 b3
50.40179 1.00000 0.19861
attr(,"status")
[1] " " "M" "U"
$value
[1] 6181.2
$counts
[1] 71 28
$convergence
[1] 0
$message
[1] "Rcgmin seems to have converged"
$bdmsk
[1] 1 0 1
```

```
> a1vmbm2 <- Rvmmin(start1, hobbs.f, hobbs.g, bdmsk=c(1,0,1),
  + lower=c(0,0,0), upper=c(200, 60, .3))
  > a1vmbm2
\frac{1}{2} \frac{1}{2}b1 b2 b3
50.40179 1.00000 0.19861
 attr(,"status")
[1] " " "M" "U"
$value
[1] 6181.2
$counts
function gradient
     20 11
$convergence
[1] 0
$message
[1] "Rvmminb appears to have converged"
$bdmsk
[1] 1 0 1
  > # Active bound
  > a1cgm2x <- Rcgmin(start1, hobbs.f, hobbs.g, bdmsk=c(1,0,1),
  + lower=c(0,0,0), upper=c(48, 60, .3))
Rcgminb: maxfeval set to 10000
  > a1cgm2x
$par
 b1 b2 b3
48.00000 1.00000 0.21597
 attr(,"status")
[1] " " "M" "U"
$value
[1] 6206.1
$counts
[1] 36 14
$convergence
[1] 0
$message
[1] "Rcgmin seems to have converged"
$bdmsk
[1] -1 0 1
  > a1vmm2x <- Rvmmin(start1, hobbs.f, hobbs.g, bdmsk=c(1,0,1),
  + lower=c(0,0,0), upper=c(48, 60, .3))
  > a1vmm2x
\frac{1}{2} \frac{1}{2}b1 b2 b3
48.00000 1.00000 0.21597
 attr(,"status")
[1] " " "M" "U"
$value
[1] 6206.1
$counts
```

```
function gradient<br>15 10
                  1<sub>C</sub>
$convergence
[1] 0
$message
[1] "Rvmminb appears to have converged"
$bdmsk
[1] 1 0 1
```
6 Brief example of minpack.lm

Recently Kate Mullen provided some capability for the package minpack.lm to include bounds constraints. I am particularly happy that this effort is proceeding, as there are significant differences in how minpack. Im and nlmrt are built and implemented. They can be expected to have different performance characteristics on different problems. A lively dialogue between developers, and the opportunity to compare and check results can only improve the tools.

The examples below are a very quick attempt to show how to run the Ratkowsky-Huet problem with nls.lm from minpack.lm.

```
> require(minpack.lm)
  > anlslm <- nls.lm(ones, lower=rep(-1000,4), upper=rep(1000,4), jres, jjac, yield=pastured
   > cat("anlslm from ones\n")
anlslm from ones
  > print(strwrap(anlslm))
 [1] "c(t1 = NaN, t2 = NaN, t3 = NaN, t4 = NaN)"
 [2] "c(NaN, NaN, NaN, NaN, NaN, NaN, NaN, NaN, NaN, NaN,"
 [3] "NaN, NaN, NaN, NaN, NaN, NaN)"
 [4] "c(NaN, NaN, NaN, NaN, NaN, NaN, NaN, NaN, NaN)"
 [5] "-1"
 [6] "Number of iterations has reached `maxiter' == 50."
 [7] "list(t1 = 3, t2 = NaN, t3 = NaN, t4 = NaN)"
 [8] "50"
 [9] "c(17533.3402000004, 16864.5616372991, NaN, NaN, NaN,"
[10] "NaN, NaN, NaN, NaN, NaN, NaN, NaN, NaN, NaN, NaN,"
[11] "NaN, NaN, NaN, NaN, NaN, NaN, NaN, NaN, NaN, NaN,"
[12] "NaN, NaN, NaN, NaN, NaN, NaN, NaN, NaN, NaN, NaN,"
[13] "NaN, NaN, NaN, NaN, NaN, NaN, NaN, NaN, NaN, NaN,"
[14] "NaN, NaN, NaN, NaN, NaN, 4.24399161290846e-313)"
[15] "NaN"
  > anlslmh <- nls.lm(huetstart, lower=rep(-1000,4), upper=rep(1000,4), jres, jjac, yield=pa
  > cat("anlslmh from huetstart\n")
anlslmh from huetstart
  > print(strwrap(anlslmh))
 [1] "c(t1 = 69.9551973916736, t2 = 61.6814877170941, t3 ="
 [2] "-9.20891880263442, t4 = 2.37781455978467)"
[3] "c(9, -4.54037977686007, 105.318033221555,"
 [4] "403.043210394646, -4.54037977686007,
 [5] "3.51002837648689, -39.5314537948582,"
 [6] "-137.559566823765, 105.318033221555,"
 [7] "-39.5314537948582, 1668.11894086463, 6495.6770219983,"
 [8] "403.043210394646, -137.559566823765, 6495.6770219983,"
```

```
[9] "25481.4530263827)"
[10] "c(0.480682793156284, 0.669303022602282,"
[11] "-2.28431914156848, 0.843754801653773,"
[12] "0.734587578832205, 0.0665510313004276,"
[13] "-0.985814877917534, -0.0250630130722698,"
[14] "0.500317790294616)"
[15] "1"
[16] "Relative error in the sum of squares is at most"
[17] "`ftol'."
[18] "list(t1 = 3, t2 = 2.35105755434962, t3 ="
[19] "231.250186433367, t4 = 834.778914353851)'
[20] "42"
[21] "c(13386.9099465603, 13365.3097414383,"
[22] "13351.1970260154, 13321.6478455192, 13260.1135652244,"
[23] "13133.6391318145, 12877.8542053848, 12373.5432344283,"
[24] "11428.8257706578, 9832.87890178625, 7138.12187613237,"
[25] "3904.51162830831, 2286.64875980737, 1978.18149980306,"
[26] "1620.89081508973, 1140.58638304326, 775.173148616758,"
[27] "635.256627921479, 383.73614705125, 309.341249993346,"
[28] "219.735856060244, 177.398738179149, 156.718991828473,"
[29] "135.51359456819, 93.4016394568234, 72.8219383036211,"
[30] "66.3315609834918, 56.2809616213409, 54.9453021619835,"
[31] "53.6227655715768, 51.9760950696957, 50.1418078879664,"
[32] "48.1307021647523, 44.709775710931, 42.8838792615121,"
[33] "32.3474231559242, 26.5253835687495, 15.3528215541072,"
[34] "14.721550701286, 8.37980617628203, 8.37589765770218,"
[35] "8.37588365348105, 8.37588355972584)"
[36] "8.37588355972584"
```
References

- [1] Timur V. Elzhov, Katharine M. Mullen, Andrej-Nikolai Spiess, and Ben Bolker, minpack.lm: R interface to the levenberg-marquardt nonlinear leastsquares algorithm found in minpack, plus support for bounds, R Project for Statistical Computing, 2012, R package version 1.1-6.
- [2] S. (Sylvie) Huet et al., Statistical tools for nonlinear regression: a practical guide with S-PLUS examples, Springer series in statistics, 1996.
- [3] J. J. Moré, B. S. Garbow, and K. E. Hillstrom, $ANL-80-74$, User Guide for MINPACK-1, Tech. report, 1980.
- [4] J. C. Nash, Compact numerical methods for computers : linear algebra and function minimisation, Hilger, Bristol :, 1979 (English).
- [5] David A. Ratkowsky, Nonlinear regression modeling: A unified practical approach, Marcel Dekker Inc., New York and Basel, 1983.