# Refactoring the nls() function in R

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# **Abstract**

This article considers the features and limitations of the R function nls(). It arose from a Google Summer of Code project "Improvements to nls()", a tool intended for the estimation of models written as a formula that has at least one parameter that is "nonlinear". That is, the model is not estimable via solving a finite set of linear equations. A companion document Variety in Nonlinear Least Squares Codes presents an

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overview of methods for the problem which takes a much wider view of the problem of minimizing a function that can be written as a sum of squared terms, and of the even more general case of nonlinear regression.

An important overall consideration in our work has been the maintainability of the code base that supports the nls() functionality, as we believe that the existing code makes maintenance and improvement very difficult. Moreover, legacy applications and examples mean some improvements are blocked. Thus we have not addressed all the issues raised, but we believe that the discussion and trials so far are relevant to improving both nls() and R generally.

# The nls() function: strengths and shortcomings

nls() is the tool in base R (the distributed software package from CRAN for estimating nonlinear statistical models. The function was developed mainly in the 1980s and 1990s by Doug Bates et al., initially for S (see <a href="https://en.wikipedia.org/wiki/S\_%28programming\_language%29">https://en.wikipedia.org/wiki/S\_%28programming\_language%29</a>). The ideas spring primarily from the book by Bates and Watts (1988).

The nls() function has a remarkable and quite comprehensive set of capabilities for estimating nonlinear models that are expressed as formulas. In particular, we note that it

- handles formulas that include R functions
- allows data to be subset
- permits parameters to be indexed over a set of related data
- produces measures of variability (i.e., standard error estimates) for the estimated parameters
- has related profiling capabilities for exploring the likelihood surface as parameters are changed

With such a range of features and a long history, it is not surprising that the code has become untidy and overly patched. It is, to our mind, essentially unmaintainable. Moreover, its underlying methods can and should be improved. Let us review some of the issues. We will then propose corrective actions, some of which we have carried out either in demonstrations or in the distributed coed.

#### Feature: Convergence and termination tests (FIXED)

A previous issue with nls() that prevented it from providing parameter estimates for zero-residual (i.e., perfect fit) data was corrected thanks to suggestions by one of us.

In the manual page for nls() in R 4.0.0 there is the warning

#### Do not use nls on artificial "zero-residual" data.

The nls function uses a relative-offset convergence criterion that compares the numerical imprecision at the current parameter estimates to the residual sum-of-squares. This performs well on data of the form

$$y = f(x, \theta) + eps$$

(with var(eps) > 0). It fails to indicate convergence on data of the form

$$y = f(x, \theta)$$

because the criterion amounts to comparing two components of the round-off error.

If you wish to test nls on artificial data please add a noise component, as shown in the example below.

This amounts to admitting R cannot solve perfectly well-posed problems. The suggestion that one needs to pollute data with errors is one that should be offensive to good science.

It turns out that this issue can be easily resolved. The key "convergence test" – more properly a "termination test" for the **program** rather than for convergence of the underlying **algorithm** – is the Relative Offset Convergence Criterion (see Bates, Douglas M. and Watts, Donald G. (1981)). This works by projecting the

proposed step in the parameter vector on the gradient and estimating how much the sum of squares loss function will decrease. This is divide by the current size of the loss function to avoid scale issues. When we have "converged", the estimated decrease is very small, as usually is its ratio to the sum of squares. However, with small residuals, the sum of squares loss function is (almost) zero and we get the possibility of a zero-divide failure.

Adding a small quantity to the loss function before dividing avoids trouble. In 2021, one of us (J. Nash) proposed that nls.control() have an additional parameter scaleOffset with a default value of zero. Setting it to a small number – 1.0 is a reasonable choice – allows small-residual problems (i.e., near-exact fits) to be dealt with easily. We call this the safeguarded relative offset convergence criterion. The default value gives the legacy behaviour. We are pleased to note that this improvement is now in the R distributed code as of version 4.1.0.

### Example of a small-residual problem

```
t <- -10:10
y <- 100/(1+.1*exp(-0.51*t))
lform < -y \sim a/(1+b*exp(-c*t))
ldata<-data.frame(t=t, y=y)</pre>
plot(t,y)
lstartbad < -c(a=1, b=1, c=1)
lstart2<-c(a=100, b=10, c=1)
nlsr::nlxb(lform, data=ldata, start=lstart2)
nls(lform, data=ldata, start=lstart2, trace=TRUE)
# Fix with scaleOffset
nls(lform, data=ldata, start=lstart2, trace=TRUE, control=list(scaleOffset=1.0))
sessionInfo()
Edited output of running this function follows:
> t <- -10:10
> y <- 100/(1+.1*exp(-0.51*t))
> 1form < -y^a/(1+b*exp(-c*t))
> ldata<-data.frame(t=t, y=y)</pre>
> plot(t,y)
> lstartbad<-c(a=1, b=1, c=1)
> lstart2<-c(a=100, b=10, c=1)
> nlsr::nlxb(lform, data=ldata, start=lstart2)
nlsr object: x
residual sumsquares = 1.007e-19 on 21 observations
                  Jacobian and 19 function evaluations
  name
                   coeff
                                  SE
                                                                               JSingval
                                            tstat
                                                       pval
                                                                  gradient
                      100
                              2.679e-11
                                          3.732e+12
                                                     1.863e-216
                                                                  -6.425e-11
                                                                                    626.6
а
                      0.1
                               3.78e-13 2.646e+11 9.125e-196
                                                                  -3.393e-08
                                                                                    112.3
b
                     0.51
                                6.9e-13 7.391e+11 8.494e-204
                                                                   1.503e-08
                                                                                    2.791
> nls(lform, data=ldata, start=lstart2, trace=TRUE)
          (1.08e+00): par = (100 10 1)
40346.
11622.
          (2.93e+00): par = (101.47 \ 0.49449 \ 0.71685)
5638.0
          (1.08e+01): par = (102.23 \ 0.38062 \ 0.52792)
          (1.04e+01): par = (102.16 \ 0.22422 \ 0.41935)
642.08
97.712
          (1.79e+01): par = (100.7 \ 0.14774 \ 0.45239)
22.250
          (1.78e+02): par = (99.803 \ 0.093868 \ 0.50492)
0.025789 (1.33e+03): par = (100.01 0.10017 0.50916)
6.0571e-08 (7.96e+05): par = (100 0.1 0.51)
4.7017e-19 (1.86e+04): par = (100 0.1 0.51)
1.2440e-27 (5.71e-01): par = (100 0.1 0.51)
```

```
1.2440e-27 (5.71e-01): par = (100 0.1 0.51)
... (approx 40 lines omitted)
1.2440e-27 (5.71e-01): par = (100 0.1 0.51)
Error in nls(lform, data = ldata, start = lstart2, trace = TRUE) :
  number of iterations exceeded maximum of 50
> # Fix with scaleOffset
> nls(lform, data=ldata, start=lstart2, trace=TRUE, control=list(scaleOffset=1.0))
            (1.08e+00): par = (100 10 1)
40346.
            (2.91e+00): par = (101.47 \ 0.49449 \ 0.71685)
11622.
5638.0
            (9.23e+00): par = (102.23 \ 0.38062 \ 0.52792)
642.08
            (5.17e+00): par = (102.16 \ 0.22422 \ 0.41935)
97.712
            (2.31e+00): par = (100.7 \ 0.14774 \ 0.45239)
22,250
            (1.11e+00): par = (99.803 \ 0.093868 \ 0.50492)
0.025789
            (3.79e-02): par = (100.01 \ 0.10017 \ 0.50916)
6.0571e-08
            (5.80e-05): par = (100 \ 0.1 \ 0.51)
4.7017e-19
            (1.62e-10): par = (100 \ 0.1 \ 0.51)
Nonlinear regression model
  model: v \sim a/(1 + b * exp(-c * t))
   data: ldata
         0.10
100.00
                0.51
residual sum-of-squares: 4.7e-19
Number of iterations to convergence: 8
Achieved convergence tolerance: 1.62e-10
```

Running under: Linux Mint 20.3

### More general termination tests

The single convergence criterion of nls() leaves out some possibilities that could be useful for some problems. The package nlsr (John C Nash and Duncan Murdoch (2019)) already offers both the safeguarded relative offset test (roffset) as well as a small sum of squares test (smallsstest) that compares the latest evaluated sum of squared (weighted) residuals to e4 times the initial sum of squares. The value of e4 is computed as

```
epstol<-100*.Machine$double.eps
e4 <- epstol^4
e4</pre>
```

```
## [1] 2.430865e-55
```

We note that nls() uses a termination test to stop after maxiter "iterations". Unfortunately, for almost all iterative algorithms, the meaning of "iteration" requires careful examination of the code. we prefer to express such tests using the number of times the residuals or the jacobian have been computed and put upper limits on these. Our codes exit (terminate) when these limits are reached. Generally we prefer larger limits than the default maxiter=50 of nls(), but that may simply reflect our history of dealing with more difficult problems as we are the tool-makers; users consult us when things go wrong.

### Feature: Failure when Jacobian is computationally singular

This is the infamous "singular gradient" termination message of nls(). A Google search of

```
R nls "singular gradient"
```

gets over 4000 hits that are spread over the years. In some cases this is due to the failure of the simple finite difference approximation of the Jacobian in the numericDeriv() function that is a part of nls(). nlsr can use analytic derivatives, and we could import this functionality to the nls() code as an improvement. See below in the section Jacobian computation.

A common source of the issue is that the Jacobian is very close to singular for some values of the model parameters. In such cases we need to find an alternative algorithm to the Gauss-Newton iteration of nls(). The most common work-around is the Levenberg-Marquardt stabilization (see Marquardt (1963), Levenberg (1944), Nash (1977)). Versions of this have been implemented in packages minpack.lm and nlsr. and we have prepared experimental versions of nls replacements that can incorporate stabilizations. Integration of such ideas with other code structures so that all the features of nls() work properly has unfortunately proved difficult.

Let us consider an example which presents the infamous 'singular gradient' error message. This was posted on StackExchange. https://stats.stackexchange.com/questions/13053/singular-gradient-error-in-nls-with-correct-starting-values

```
reala=-3
realb=5
realc=0.5
realr=0.7
realm=1
x=1:11 #x values 11 timepoint data
#linear+exponential function
y=reala + realb*realr^(x-realm) + realc*x
testdat=data.frame(x,y)
strt<-list(a=-3, b=5, c=0.5, r=0.7, m=1)
jform<-y~a+b*r^(x-m)+c*x # Formula
linexp=try(nls(jform, data=testdat, start=strt, trace=F))
## Error in nlsModel(formula, mf, start, wts, scaleOffset = scOff, nDcentral = nDcntr) :
##
     singular gradient matrix at initial parameter estimates
library(nlsr)
# Note singular values of Jacobian in rightmost column
linexp2<-try(nlxb(jform, data=testdat, start=strt, trace=F))</pre>
linexp2
## nlsr object: x
## residual sumsquares = 0
                                  11 observations
                              on
##
                    Jacobian and
                                  1 function evaluations
                      coeff
                                     SE
##
     name
                                               tstat
                                                          pval
                                                                     gradient
                                                                                  JSingval
## a
                          -3
                                         NA
                                                                NA
                                                                              0
                                                                                      26.49
                                                    NA
## b
                           5
                                         NA
                                                    NA
                                                                NA
                                                                              0
                                                                                      9.615
## c
                         0.5
                                         NA
                                                    NA
                                                                NA
                                                                              0
                                                                                      2.466
                         0.7
                                         NA
                                                                              0
                                                                                     0.1098
## r
                                                    NA
                                                                NA
## m
                                         NA
                                                    NA
                                                                NA
                                                                                  1.398e-16
```

In the above, we see that the Jacobian is essentially singular as shown by its singular values. Note that we DISPLAY the singular values in a single column for convenience in the output. The values do NOT correspond to individual parameters, but are a property of the whole problem.

### Feature: Jacobian computation

nls(), with the numericDeriv() function, computes the Jacobian as the "gradient" attribute of the residual vector. This is implemented as a mix of R and C code, but we have created a rather more compact version entirely in R. Code project. See the companion document **Jacobian Calculations for nls()** (?? need citation – currently **DerivsNLS.pdf**).

The Jacobian used by nlsr::nlxb() is computed from analytic expressions, while that of nls() (and, by extension, minpack.lm::nlsLM) will use a numerical approximation. This can lead to different behaviour:

- A pasture regrowth problem (Huet et al. (2004), page 1, based on Ratkowsky (1983)) has a poorly conditioned Jacobian and nls() fails with "singular gradient". Worse, numerical approximation to the Jacobian give the error "singular gradient matrix at initial parameter estimates" for minpack.lm::nlsLM so that the Marquardt stabilization is unable to take effect, while the analytic derivatives of nlsr::nlxb give a solution.
- Karl Schilling (private communication) provided an example where a model specified with the formula  $y \sim a * (x \hat{ } b)$  causes nlsr::nlxb to fail because the partial derivative w.r.t. b is  $a * (x\hat{ } b * log(x))$ . If there is data for which x = 0, this is undefined. In such cases, we observed that nls() and minpack.lm::nlsLM found a solution, though it can be debated whether such lucky accidents can be taken as an advantage.

Note that the selfStart models in the ./src/library/stats/R/zzModels.R file provide the Jacobian in the "gradient" attribute of the "one-sided" formula that defines each model, and these Jacobians may be the analytic forms. The nls() function, after computing the "right hand side" or rhs of the residual, then checks to see if the "gradient" attribute is defined, and, if not, uses numericDeriv to compute a Jacobian into that attribute. This code is within the nlsModel() or nlsModel.pliner() functions. The use of analytic Jacobians almost certainly contributes to the performance of nls() on selfStart models.

### Feature: Subsetting

nls() accepts an argument subset. Unfortunately, this acts through the mediation of model.frame and is not clearly obvious in the source code files /src/library/stats/R/nls.R and /src/library/stats/src/nls.C.

While the implementation of subset at the level of the call to nls() has a certain attractiveness, it does mean that the programmer of the solver needs to be aware of the source (and value) of objects such as the data, residuals and Jacobian. By preference, we would implement subsetting by means of zero-value weights, with observation counts (and degrees of freedom) computed via the numbers of non-zero weights. Alternatively, we would extract a working dataframe from the relevant elements in the original.

#### Feature: na.action

na.action is an argument to the nls() function, but it does not appear obviously in the source code, often being handled behind the scenes after referencing the option na.action. A useful, but possibly dated, description is given in: https://stats.idre.ucla.edu/r/faq/how-does-r-handle-missing-values/

The typical default action, which can be seen by using the command getOption("na.action") is na.omit. This option essentially presents computations with data with all observations containing any missing values (i.e. any row of a data frame with an NA) omitted. na.exclude does much the same for computations, but keeps the rows with NA elements so that predictions are in the correct row position. We recommend that workers actually test output to verify behaviour is as wanted.

A succinct description of the issue is given in: https://stats.stackexchange.com/questions/492955/should-i-use-na-omit-or-na-exclude-in-a-linear-model-in-r

The only benefit of na.exclude over na.omit is that the former will retain the original number of rows in the data. This may be useful where you need to retain the original size of the dataset - for example it is useful when you want to compare predicted values to original values. With na.omit you will end up with fewer rows so you won't as easily be able to compare.

na.pass simply passes on data "as is", while na.fail will essentially stop if any missing values are present.

### Feature: model frame

model is an argument to the nls() function, which is documented

model logical. If true, the model frame is returned as part of the object. Default is FALSE.

Indeed, the argument only gets used when nls() is about to return its result object, and the element model is NULL unless the calling argument model is TRUE. (Using the same name could be confusing.) However, the model frame is used within the function code in the form of the object mf.

#### Feature: sources of data

nls() can be called without specifying the data argument. In this case, it will search in the available environments (i.e., workspaces) for suitable data objects. We do NOT like this approach, but it is "the R way". R allows users to leave many objects in the default (.GlobalEnv) workspace. Moreover, users have to actively suppress saving this workspace (.RData) on exit, and any such file in the path when R is launched will be loaded. The overwhelming proportion of R users in our acquaintance avoid saving the workspace because of the danger of lurking data and functions which may cause unwanted results.

Nevertheless, to provide compatible behaviour with nls(), competing programs need to ensure equivalent behaviour, but users should test that the operation is correct.

### Feature: missing start vector and self-starting models

Nonlinear estimation algorithms are almost all iterative and need a set of starting parameters. nls() offers a special class of modeling functions called selfStart models. There are a number of these in base R (see list below) and others in R packages such as CRAN package nlraa (Miguez (2021)), as well as the now-archived package NRAIA. Unfortunately, the structure of selfStart codes is such that the methods by which initial parameters are computed is entangled with the particulars of the nls() code. Though there is a getInitial() function, this is not easy to use to simply compute the initial parameter estimates, in part because it may call nls(). Such circular references are, in our view, dangerous.

In the example below, we show how the SSlogis selfStart function can generate a set of initial parameters for a 3-parameter logistic curve. The form used by SSlogis is

$$y \sim Asym/(1 + exp((xmid - tt)/scal))$$

, but we show how the starting parameters for this model can be transformed to those of another form of the model, namely,

$$y \sim b1/(1 + b2 * exp(-b3 * t))$$

Let us look at the actual code for SSlogis() in R-devel/src/library/stats/R/zzModels.R:

```
SSlogis <- selfStart(~ Asym/(1 + exp((xmid - input)/scal)),
        function(input, Asym, xmid, scal)
        {
               .expr1 <- xmid - input
               .expr3 \leftarrow exp(.e2 \leftarrow .expr1/scal)
               .expr4 <- 1 + .expr3
               .value <- Asym/.expr4
               .actualArgs <- as.list(match.call()[c("Asym", "xmid", "scal")])</pre>
               if(all(vapply(.actualArgs, is.name, NA)))
                        .expr10 <- .expr4^2
                    .grad <- array(0, c(length(.value), 3L), list(NULL, c("Asym", "xmid", "scal")))</pre>
                    .grad[, "Asym"] <- 1/.expr4
                        .grad[, "xmid"] <- - (xm <- Asym * .expr3/scal/.expr10)</pre>
                        .grad[, "scal"] \leftarrow xm * .e2
                   dimnames(.grad) <- list(NULL, .actualArgs)</pre>
                   attr(.value, "gradient") <- .grad
```

```
}
       .value
},
initial = function(mCall, data, LHS, ...) {
      xy <- sortedXyData(mCall[["input"]], LHS, data)</pre>
      if(nrow(xy) < 4) {
           stop("too few distinct input values to fit a logistic model")
      z \leftarrow xy[["y"]]
      ## transform to proportion, i.e. in (0,1):
      rng <- range(z); dz <- diff(rng)</pre>
      z \leftarrow (z - rng[1L] + 0.05 * dz)/(1.1 * dz)
      xy[["z"]] \leftarrow log(z/(1 - z))
                                            # logit transformation
      aux \leftarrow coef(lm(x \sim z, xy))
      pars <- coef(nls(y \sim 1/(1 + exp((xmid - x)/scal))),
                         data = xy,
                         start = list(xmid = aux[[1L]], scal = aux[[2L]]),
                         algorithm = "plinear", ...))
      setNames(pars [c(".lin", "xmid", "scal")],
                mCall[c("Asym", "xmid", "scal")])
},
parameters = c("Asym", "xmid", "scal"))
```

This R function includes analytic expressions for the Jacobian ("gradient"). These could be be useful to R users, especially if documented. Moreover, we wonder why the programmers have chosen to save so many quantities in "hidden" variables, i.e., with names preceded by ".". These are then not displayed by the ls() command, making them difficult to access.

In the event that a selfStart model is not available, nls() sets all the starting parameters to 1. This is, in our view, tolerable, but could possibly be improved by using a set of values that are slightly different e.g., in the case of a model

$$y \sim a * exp(-b * x) + c * exp(-d * x)$$

it would be useful to have b and d values different so the Jacobian is not singular. Thus, some sort of sequence like 1.0, 1.1, 1.2, 1.3 for the four parameters might be better and it can be provided quite simply instead of all 1's.

```
weed \leftarrow 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
NLSformula0 \leftarrow y \sim b1/(1+b2*exp(-b3*tt))
NLSformula <- y ~ SSlogis(tt, Asym, xmid, scal)
NLSformulax <- y ~ Asym/(1+exp((xmid-tt)/scal))</pre>
NLStestdata <- data.frame(y=weed, tt=tt)</pre>
s0 <- getInitial(NLSformula, NLStestdata)</pre>
print(s0)
                 xmid
                           scal
       Asym
## 196.1862 12.4173
                         3.1891
# We transform the parameters for the NLS formula 0 model of original specification.
s1<-list(b1=s0[1], b2=exp(s0[2]/s0[3]), b3=1/s0[3])
print(as.numeric(s1))
## [1] 196.18624 49.09163
                               0.31357
```

```
# No actual improvement because nls() has been already used to get the starting values,
# but we do get SEs
hobblog<-nls(NLSformula0, data=NLStestdata, start=s1)
summary(hobblog)
##
## Formula: y \sim b1/(1 + b2 * exp(-b3 * tt))
##
## Parameters:
          Estimate Std. Error t value Pr(>|t|)
## b1.Asym 1.96e+02 1.13e+01 17.4 3.2e-08 ***
## b2.xmid 4.91e+01 1.69e+00 29.1 3.3e-10 ***
## b3.scal 3.14e-01 6.86e-03 45.7 5.8e-12 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.536 on 9 degrees of freedom
## Number of iterations to convergence: 0
## Achieved convergence tolerance: 1.11e-06
deviance(hobblog)
## [1] 2.5873
# nls fails without selfStart -- singular gradient, even on stabilized formula
try(hobblogx<-nls(NLSformulax, data=NLStestdata))</pre>
## Warning in nls(NLSformulax, data = NLStestdata): No starting values specified for some parameters.
## Initializing 'Asym', 'xmid', 'scal' to '1.'.
## Consider specifying 'start' or using a selfStart model
## Error in nls(NLSformulax, data = NLStestdata) : singular gradient
# But Marquardt is able to get a solution easily
library(nlsr)
hobblogxx<-nlxb(NLSformulax, data=NLStestdata, start=c(Asym=1, xmid=1, scal=1))
hobblogxx
## nlsr object: x
## residual sumsquares = 2.5873 on 12 observations
##
      after 23
                   Jacobian and 31 function evaluations
##
    name
                                                                  gradient
                     coeff
                                   SE
                                           tstat
                                                       pval
                                                                              JSingval
## Asym
                   196.186
                                   11.31
                                             17.35 3.167e-08
                                                                 4.136e-12
                                                                                  44.93
## xmid
                   12.4173
                                  0.3346
                                              37.11 3.716e-11
                                                                 7.492e-11
                                                                                   15.6
                                              45.69 5.768e-12 1.606e-11
## scal
                                  0.0698
                    3.18908
                                                                                 0.0474
selfStart models in base R
The following models are provided (in file ./src/library/stats/R/zzModels.R)
SSasymp
                - asymptotic regression model
SSasympOff
               - alternate formulation of asymptotic regression model with offset
               - exponential curve through the origin to an asymptote
SSasympOrig
               -y \sim A1 * \exp(-\exp(1rc1)*input) + A2 * \exp(-\exp(1rc2) * input)
SSbiexp
               - y ~ Dose * (exp(1Ke + 1Ka - 1C1) * (exp(-exp(1Ke) * input) -
SSfol
                          exp(-exp(lKa) * input))/(exp(lKa) - exp(lKe)))
```

```
SSfpl - four parameter logistic model
SSlogis - three parameter logistic model
```

SSmicmen - Michaelis-Menten model for enzyme kinetics

SSgompertz2 - Gompertz model for growth curve data
SSweibull - Weibull model for growth curve data

#### Strategic issues in selfStart models

Because the Gauss-Newton algorithm is unreliable from many starting sets of parameters, selfStart models are more than an accessory to nls() but a part of the infrastructure. However, creating such functions is a lot of work, and their documentation (file ./src/library/stats/man/selfStart.Rd) is quite complicated. We believe that the focus would better be placed on getting good initial parameters, possibly with some interactive tools. That is, the emphasis should be on the getInitial() function, though avoiding the current calls back to nls().

### Issue: documentation of the results of running nls()

The output of nls() is an object of class "nls" which has the following structure:

### nls() result output according to the documentation

```
A list of:
```

```
an nlsModel object incorporating the model.
            the expression that was passed to nls as the data argument. The actual data values
data
              are present in the environment of the m components, e.g., environment(m$conv).
call
            the matched call with several components, notably algorithm.
na.action
           the "na.action" attribute (if any) of the model frame.
dataClasses the "dataClasses" attribute (if any) of the "terms" attribute of the model frame.
            if model = TRUE, the model frame.
model
            if weights is supplied, the weights.
weights
            a list with convergence information.
convInfo
control
            the control list used, see the control argument.
```

There are also two deprecated items if algorithm = "port" fit only. These are convergence (a code = 0 for convergence) and message. These are available from convInfo.

#### Example output

##

deviance.nls

To illustrate, let us run the Croucher example.

stats

```
# Croucher example
xdata \leftarrow c(-2,-1.64,-1.33,-0.7,0,0.45,1.2,1.64,2.32,2.9)
ydata \leftarrow c(0.699369, 0.700462, 0.695354, 1.03905, 1.97389, 2.41143, 1.91091, 0.919576, -0.730975, -1.42001)
p1<- 1; p2<-0.2; NLSstart<-list(p1=p1,p2=p2)
NLSformula <- ydata ~ p1*cos(p2*xdata) + p2*sin(p1*xdata)</pre>
NLSdata<-data.frame(xdata, ydata)</pre>
# Try full output version of nls
library(nlspkg) # use the packaged version of nls()
## Registered S3 methods overwritten by 'nlspkg':
##
     method
                        from
##
     anova.nls
                         stats
##
     coef.nls
                         stats
##
     confint.nls
                         stats
```

```
##
     df.residual.nls
##
     fitted.nls
                       stats
##
     formula.nls
##
     logLik.nls
                       stats
##
     nobs.nls
                       stats
##
     plot.profile.nls stats
     predict.nls
##
                       stats
##
     print.nls
                       stats
##
     print.summary.nls stats
##
     profile.nls
                       stats
##
     residuals.nls
                       stats
##
     summary.nls
                       stats
##
     vcov.nls
                       stats
##
     weights.nls
                       stats
##
## Attaching package: 'nlspkg'
## The following objects are masked from 'package:stats':
##
       nls, nls.control, numericDeriv
##
result <- nls (NLS formula, data=NLS data, start=NLS start, model=TRUE)
# We can display the result with several commands
# str(result) -- displays large amount of material - suppressed here
     as it is too wide for the page
# result # displays the object
# ls(result) # to list the elements of the output
# ls(result$m) # and in particular the "m" object
```

#### Concerns with content of the nls result object

The nls object contains some elements that are awkward to produce by other algorithms, but some information that would be useful is not presented obviously.

In the following, we use result as the returned object from nls().

The data return element is an R symbol. To actually access the data from this element, we need to use the syntax

```
eval(parse(text=result$data))
```

However, if the call is made with model=TRUE, then there is a returned element model which contains the data, and we can list its contents using

```
ls(result$model)
```

and if there is an element called xdata, it can be accessed as result\$model\$xdata.

#### Information that is NOT in the nls result object

```
The jacobian matrix (partial derivatives of residuals w.r.t. the parameters) at the returned parameters.

The number of residual evaluations (sum of squares computations) used.

The number of Jacobian evaluations used.

However, actually looking at the structure of a returned result gives a list of 11 items:

$ resid : num [1:12] 0.0119 -0.0328 0.092 0.2088 0.3926 ...

- attr(* "gradient")= num [1:12 1:3] 0.0271 0.0367 0.0496 0.0666 0.089
```

```
..- attr(*, "gradient")= num [1:12, 1:3] 0.0271 0.0367 0.0496 0.0666 0.089 ...
 ... - attr(*, "dimnames")=List of 2
 .. .. ..$ : NULL
 .. .. ..$ : chr [1:3] "Asym" "xmid" "scal"
$ jacobian : num [1:12, 1:3] 0.0271 0.0367 0.0496 0.0666 0.089 ...
 ..- attr(*, "dimnames")=List of 2
 .. ..$ : NULL
 ....$ : chr [1:3] "Asym" "xmid" "scal"
              : num 31
              : num 23
$ jeval
$ coefficients: Named num [1:3] 196.19 12.42 3.19
 ..- attr(*, "names")= chr [1:3] "Asym" "xmid" "scal"
$ ssquares
             : num 2.59
$ lower
              : num [1:3] -Inf -Inf -Inf
$ upper
             : num [1:3] Inf Inf Inf
             : int(0)
$ maskidx
$ weights
              : NULL
$ formula
              :Class 'formula' language y ~ Asym/(1 + exp((xmid - tt)/scal))
 ....- attr(*, ".Environment")=<environment: R_GlobalEnv>
- attr(*, "class")= chr "nlsr"
```

This is still a smaller result object than the one nls() returns. Moreover, nlxb explicitly returns the sum of squares as well as the residual vector and Jacobian. The counts of evaluations are also returned. Working on this project showed several potential updates to the nlsr documentation. Note that the singular values of the Jacobian are actually computed in the print and summary methods for the result.

#### Weights in returned functions from nls()

The functions resid() (an alias for residuals()) and fitted() and lhs() are UNWEIGHTED. But if we return ans from nls() or minpack.lm::nlsLM or our new nlsj (interim package), then ans\$m\$resid() is WEIGHTED.

### Interim output from the "port" algorithm

As the nls() man page states, when the "port" algorithm is used with the trace argument TRUE, the iterations display the objective function value which is 1/2 the sum of squares (or deviance). It is likely that the trace display is embedded in the Fortran of the nlminb routine that is called to execute the "port" algorithm, but the discrepancy is nonetheless unfortunate for users.

#### Failure to return best result achieved

If nls() reaches a point where it cannot continue but has not found a point where the relative offset convergence criterion is met, it may simply exit, especially if a "singular gradient" (singular Jacobian) is found. However, this may occur AFTER the function has made considerable progress in reducing the sum of squared residuals. An example is to be found in the Tetra\_1.R example from the nlsCompare package. Here is an abbreviated version of that problem and the nls() output:

```
time=c(1, 2, 3, 4, 6, 8, 10, 12, 16)
conc = c(0.7, 1.2, 1.4, 1.4, 1.1, 0.8, 0.6, 0.5, 0.3)
NLSdata <- data.frame(time,conc)</pre>
NLSstart <-c(lrc1=-2,lrc2=0.25,A1=150,A2=50) # a starting vector (named!)
NLSformula <-conc ~ A1*exp(-exp(lrc1)*time)+A2*exp(-exp(lrc2)*time)
tryit <- try(nls(NLSformula, data=NLSdata, start=NLSstart, trace=TRUE))</pre>
## 61216.
             (3.56e+03): par = (-2 \ 0.25 \ 150 \ 50)
## 2.1757
             (2.23e+01): par = (-1.9991 \ 0.31711 \ 2.6182 \ -1.3668)
             (7.14e+00): par = (-1.9605 -2.6203 2.5753 -0.55599)
## 1.6211
## Error in nls(NLSformula, data = NLSdata, start = NLSstart, trace = TRUE) :
     singular gradient
print(tryit)
## [1] "Error in nls(NLSformula, data = NLSdata, start = NLSstart, trace = TRUE) : \n singular gradien
## attr(,"class")
## [1] "try-error"
## attr(,"condition")
## <simpleError in nls(NLSformula, data = NLSdata, start = NLSstart, trace = TRUE): singular gradient>
```

Note that the sum of squares has been reduced from 61216 to 1.6211, but unless trace is invoked, the user will not get any information about this. This would be an almost trivial change to the 'nls() function and could be useful to R users.

#### Feature: partially linear models and their specification

Specifying a model to a solver should, ideally, use the same syntax across solver tools. Unfortunately, R allows multiple approaches within different modelling tools, and within nls() itself.

One obvious case is that nonlinear modeling tools are a superset of linear ones. Yet the explicit model

```
y \sim a*x + b
```

does not work with the linear modeling function 1m(), which requires this model to be specified as

```
y ~ x
```

Within nls(), consider the following FOUR different calling sequences for the same problem, though the fm2a – an intuitive choice – will not work. In this failed attempt, putting the Asym parameter in the model causes the plinear algorithm to try to add another term to the model. We believe this is unfortunate, and would like to see a consistent syntax. At the time of writing we do not foresee a resolution for this issue. In the example we have NOT evaluated the commands to save space.

```
algorithm = "plinear")
summary(fm2DNase1)
## using conditional linearity AND Asym does NOT work
\label{local_sym} fm2aDNase1 <- try(nls(density ~ Asym/(1 + exp((xmid - log(conc))/scal)),
                  data = DNase1,
                  start = list(Asym=3, xmid = 0, scal = 1),
                  algorithm = "plinear",
                  trace = TRUE))
summary(fm2aDNase1)
## without conditional linearity
fm3DNase1 <- nls(density ~ Asym/(1 + exp((xmid - log(conc))/scal)),</pre>
                  data = DNase1,
                  start = list(Asym = 3, xmid = 0, scal = 1))
summary(fm3DNase1)
## using Port's nl2sol algorithm
fm4DNase1 <- try(nls(density ~ Asym/(1 + exp((xmid - log(conc))/scal)),</pre>
                  data = DNase1,
                  start = list(Asym = 3, xmid = 0, scal = 1),
                  algorithm = "port"))
summary(fm4DNase1)
```

### Issue: code structure and documentation for maintenance

The nls() code is structured in a way that inhibits both maintenance and improvement. In particular, the iterative setup is such that introduction of Marquardt stabilization is not easily available. To obtain performance a lot of the code was in C with consequent calls and returns that complicate the code. Over time, R has become much more efficient on modern computers, and the need to use compiled C and Fortran is less critical. Moreover, the burden for maintenance could be reduced by moving code entirely to R.

While R and its packages are generally very well documented for usage, that documentation rarely extends as far as it might in directions helpful for code maintenance. The paucity of such documentation is exacerbated by the mixed R/C/Fortran code base, where seemingly trivial differences in things like representation of numbers or base index for arrays lead to traps for unwary programmers.

For nls() the concern is demonstrated by a short email query from John Nash to Doug Bates and his reply. This is NOT a criticism of Prof. Bates (or any other) work, but a reflection on how difficult it is to develop code in this subject area and to keep it maintainable. We have experienced similar loss of understanding for some of our own codes.

. . .

https://gitlab.com/nashjc/improvenls/-/blob/master/Croucher-expandednlsnoc.R ... has the test problem and the expanded code. Around line 367 is where we are scratching our heads. The function code (from nlsModel()) is in the commented lines below the call. This is

```
# > setPars
# function(newPars) {
# setPars(newPars)
# resid <<- .swts * (lhs - (rhs <<- getRHS())) # envir = thisEnv {2 x}
# dev <<- sum(resid^2) # envir = thisEnv
# if(length(gr <- attr(rhs, "gradient")) == 1L) gr <- c(gr)</pre>
```

```
# QR <<- qr(.swts * gr) # envir = thisEnv
# (QR$rank < min(dim(QR$qr))) # to catch the singular gradient matrix
# }</pre>
```

I'm anticipating that we will be able to set up a (possibly inefficient) code with documentation that will be easier to follow and test, then gradually figure out how to make it more efficient.

In both there is the recursive call, which must have a purpose I don't understand.

The reply was

```
I'm afraid that I don't know the purpose of the recursive call either. I know that I wrote the code to use a closure for the response, covariates, etc., but I don't recall anything like a recursive call being necessary.
```

I don't think I will be of much help. My R skills have atrophied to the point where I wouldn't even know how to start exploring what is happening in the first call as opposed to the recursive call.

### Feature: indexed parameters

The man file for nls() includes the following example of a situation in which parameters are indexed. It also uses the "plinear" option as an added complication. Here we use a truncated version of the example to save display space.

```
## The muscle dataset in MASS is from an experiment on muscle
## contraction on 21 animals. The observed variables are Strip
## (identifier of muscle), Conc (Cacl concentration) and Length
## (resulting length of muscle section).
if(! requireNamespace("MASS", quietly = TRUE)) stop("Need MASS pkg")
mm<- MASS::muscle[1:12,] # take only 1st few values of Strip
str(mm)
## 'data.frame':
                   12 obs. of 3 variables:
## $ Strip : Factor w/ 21 levels "S01", "S02", "S03", ...: 1 1 1 1 2 2 2 2 3 3 ...
## $ Conc : num 1 2 3 4 1 2 3 4 0.25 0.5 ...
## $ Length: num 15.8 20.8 22.6 23.8 20.6 26.8 28.4 27 7.2 15.4 ...
mm<-droplevels(mm)
str(mm)
## 'data.frame': 12 obs. of 3 variables:
## $ Strip : Factor w/ 3 levels "S01", "S02", "S03": 1 1 1 1 2 2 2 2 3 3 ...
## $ Conc : num 1 2 3 4 1 2 3 4 0.25 0.5 ...
```

```
## $ Length: num 15.8 20.8 22.6 23.8 20.6 26.8 28.4 27 7.2 15.4 ...
nlev <- nlevels(mm)</pre>
withAutoprint({
  ## The non linear model considered is
           Length = alpha + beta*exp(-Conc/theta) + error
  ## where theta is constant but alpha and beta may vary with Strip.
  with(mm, table(Strip)) # 2, 3 or 4 obs per strip
 nl <- nlevels(mm$Strip)</pre>
  ## We first use the plinear algorithm to fit an overall model,
  ## ignoring that alpha and beta might vary with Strip.
  musc.1 <- nls(Length ~ cbind(1, exp(-Conc/th)), mm,</pre>
                start = list(th = 1), algorithm = "plinear")
  summary(musc.1)
  ## Then we use nls' indexing feature for parameters in non-linear
  ## models to use the conventional algorithm to fit a model in which
  ## alpha and beta vary with Strip. The starting values are provided
  ## by the previously fitted model.
  ## Note that with indexed parameters, the starting values must be
  ## given in a list (with names):
  ## ?? but why use b here AND in the new formula??
  b <- coef(musc.1)</pre>
  musc.2 <- nls(Length ~ a[Strip] + b[Strip]*exp(-Conc/th), data=mm,</pre>
                start = list(a = rep(b[2], nl), b = rep(b[3], nl), th = b[1]))
  summary(musc.2)
## > with(mm, table(Strip))
## Strip
## S01 S02 S03
## 4 4 4
## > nl <- nlevels(mm$Strip)</pre>
## > musc.1 <- nls(Length ~ cbind(1, exp(-Conc/th)), mm, start = list(th = 1),</pre>
## +
         algorithm = "plinear")
## > summary(musc.1)
##
## Formula: Length ~ cbind(1, exp(-Conc/th))
##
## Parameters:
        Estimate Std. Error t value Pr(>|t|)
## th
           0.624
                       0.222 2.81 0.0203 *
                               17.82 2.5e-08 ***
## .lin1 25.684
                       1.441
## .lin2 -26.631
                       6.147
                               -4.33 0.0019 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 2.96 on 9 degrees of freedom
## Number of iterations to convergence: 5
## Achieved convergence tolerance: 4.71e-07
## > b <- coef(musc.1)
## > musc.2 <- nls(Length ~ a[Strip] + b[Strip] * exp(-Conc/th), data = mm,</pre>
```

```
start = list(a = rep(b[2], nl), b = rep(b[3], nl), th = b[1]))
## > summary(musc.2)
##
## Formula: Length ~ a[Strip] + b[Strip] * exp(-Conc/th)
##
## Parameters:
     Estimate Std. Error t value Pr(>|t|)
                            44.49
## a1 22.9277
                   0.5154
                                  1.1e-07 ***
## a2
      27.8606
                   0.5150
                            54.09 4.1e-08 ***
## a3 28.3426
                   1.0771
                            26.31
                                   1.5e-06 ***
## b1 -44.1558
                  12.3900
                            -3.56
                                     0.016 *
## b2 -43.8084
                  12.3148
                            -3.56
                                     0.016 *
## b3 -32.8421
                   2.1553
                           -15.24
                                  2.2e-05 ***
       0.5548
                   0.0808
                             6.86
                                     0.001 **
## th
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.798 on 5 degrees of freedom
## Number of iterations to convergence: 6
## Achieved convergence tolerance: 1.94e-06
```

Note that the answers for the parameters are NOT indexed. e.g. coef(musc.2) is a single level vector of parameters. We do not see a[1], a[2], a[3] but a1, a2, a3. This is because the model must integrate all the parameters because th is a common parameter across the index Strip.

We believe this structure is quite likely to cause confusion and error, and propose an alternative approach below.

# Goals of our effort to improve nls()

### Code rationalization and documentation

We want

- to provide a packaged version of nls() (call it nlsalt) coded entirely in R that matches the version in base R or what is packaged in nlspkg as described in Nash and Bhattacharjee (2022).
- to streamline the overall nls() infrastructure. By this we mean a re-factoring of the routines so they are better suited to maintenance of both the existing nls() methods and features as well as new features we or others would like to add.
- to explain what we do, either in comments or separate maintainer documentation. Since we are complaining about the lack of explanatory material for the current code, we feel it is incumbent on us to provide such material for our own work, and if possible for the existing code.

#### Rationalization of formula specifications

We have seen that nls() uses a different formula specification from the default if the plinear algorithm is used. This is unfortunate, since the user cannot then simply add algorithm="plinear" to the call. This makes errors more likely.

While not consistent with the lm() function and specification of linear models, we believe a complete set of parameters should be specified for nonlinear models. That is, the model formula should be written down as it would be computed, rather than for estimation. Thus  $y \sim a*x + b$  rather than  $y \sim x$ .

Moreover, we want to avoid the partially linear parameters appearing in the result object as .lin1, .lin2, etc., where there actual position in the model is not explicit.

A possible workaround to allow consistency would be to specify the partially linear parameters when the plinear option is specified. For example, we could use algorithm="plinear(Asym)" while keeping the general model formula from the default specification. This would allow for the output of different algorithm options to be consistent. However, we have not yet tried to code such a change.

A further complication of model specification in R is that some formulas require the user to surround a term with I( ) to inhibit interpretation of arithmetic operators. We would prefer that formulas be usable without this complication, which has limited documentation and examples.

It is also of interest to consider code that will detect partial linearity, rather than expecting the user to provide the special structure and settings. Work in this area appears to be quite limited, with "Automatic Linearity Detection, Report NA-13-04" (2013) seemingly a sole contribution we can find. Unfortunately, the mention of a "Linear and Nonlinear Discoverer" in Zhang, Cheng, and Liu (2011) is about a class of models that have almost nothing in common with the present issue. The algorithm in "Automatic Linearity Detection, Report NA-13-04" (2013) will need investigating further, but we suspect that it is complicated enough that it will not easily be adapted to a program like nls(). Possibly a simpler algorithm may be found.

#### Rationalization of indexed models

Indexed models clearly have a place in some areas of research. We need to be able to process models, as the example above, where the model formula is Length ~ a[Strip] + b[Strip] \* exp(-Conc/th) and Strip is the index. Data Length and Conc are available for all observations. Parameter th applies to every fitted point, but a[Strip] and b[Strip] depend on the index, which could be an experimental run, or some other categorization.

Given that [ ] are the R method to index arrays, their presence in a model formula signals indexing. The question is how to perform the correct calculations.

While tedious (??maybe dummies or fastdummies packages make it less so?), we could set up the model as a sum of similar models with dummy variables to turn "on" or "off" the particular case, transforming the model parameters to simple, non-indexed ones (as does nls()).

Wrapper functions to prepare the estimation calculations and present the results in a form that matches the modelling formula would avoid adding complexity to nls() and other tools.

#### Provide tests and use-case examples

We need suitable tests and use-case examples in order:

- to ensure our new nlsalt or related packages work properly, in particular, giving results comparable to or better than the nls() in base R or nlspkg;
- to test individual solver functions to ensure they work across the range of calling mechanisms, that is, different ways of supplying inputs to the solver(s);
- to pose "silly" inputs to nonlinear least squares solvers (in R) to see if these bad input exceptions are caught by the programs.

#### A test runner program ?? NLSCompare

?? Arkajyoti – do you want to expand?

When we have a "new" or trial solver function, we would like to know if it gives acceptable results on a range of sample problems of different types, starting parameters, input conditions, constraints, subsets, weights or other settings. Ideally we want to be able to get a summary that is easy to read and assess. For example, one approach would be to list the names of a set of tests with a red, green or yellow dot beside the name for FAILURE, SUCCESS, or "NOT APPLICABLE". In the last category would be a problem with constraints that the solver is not designed to handle.

To accomplish this, we need a suitable "runner" program that can be supplied with the name of a solver or solvers and a list of test problem cases. Problems generally have a base setup – a specification of the function to fit as a formula, some data and a default starting set of parameters. Other cases can be created by imposing bounds or mask constraints, subsets of the data, and different starts.

How to set up this "runner" and its supporting infrastructure is non-trivial. While the pieces are not as complicated as the inter-related parts of the solvers, especially nls(), the categorization of tests, their documentation, and the structuring to make running them straightforward and easy requires much attention to detail.

Some considerations for our test scripts:

- Is it useful to have a "base" script for each family of test problem, with numbered particular cases? That is, if we run the scripts in order, we can avoid some duplication of code and data.
- While we have developed some tags to document the test problem families and cases, we believe that such tags (essentially summary documentation) will continue to need revision as different tools and problems are included in scope of nlsCompare.
- Similarly, we expect that there will be ongoing review of the structure of the result files.

## Progress so far

Much of our work is reported in https://github.com/nashjc/RNonlinearLS.

### Reports and articles

- The present article is the central report of the Google Summer or Code 2021 project "Improvements to nls()".
- VarietyInNonlinearLeastSquaresCodes.Rmd: a review of the different algorithms and the many choices in their implementation for nonlinear least squares. This is still a DRAFT at 2021-8-20. ?? fix
- TestsDoc.Rmd (??citation) is a survey of testing tools in R. It has more general possibilities and fits into the subject of regression testing, in which case a more extensive literature review will be needed. Note that this document reflects the work in the the "Problem sets and test infrastructure" below.
- DerivsNLS.Rmd ??citation: a document to explain different ways in which Jacobian information is supplied to nonlinear least squares computation in R. File ExDerivs.R is a DRAFT of a script to provide examples.
- The BibTex bibliography ImproveNLS.bib (??citation or reference) was set up for use with all documents in this project, but has wider application to nonlinear least squares projects in general.
- MachineSummary.Rmd (??citation blog post??) is an investigation of ways to report the characteristics and identity of machines running tests. MachID.R offers a suggested concise summary function to identify a particular computational system used for tests.
- Nash and Bhattacharjee (2022) is an explanation of the construction of the nlspkg from the code in R-base.
- As the 2021 Summer of Code period was ending, one of us (JN) was invited to prepare a review of optimization in R. Ideas from the present work have been instrumental in the creation of Nash (2022).

### Problem sets and test infrastructure

?? need to clean up

We have several test problems and variants thereof in the inst/scripts/ directory of the nlsCompare package available on Github (https://github.com/ArkaB-DS/nlsCompare). We direct the reader to that package for

documentation of the test infrastructure, in particular the problems and methods files (problems.csv and methods.csv) and the various functions invoked by run.R to produce an output file in CSV form also.

Towards the end of the project, we have focused our attention on the nlsCompare package, which looks at evaluating and comparing functions for nonlinear least squares problems. We use many of the same tests as checks that are or will be built into our packages for such problems e.g., nlsalt. These use the testthat structure and may include verification of outputs that are specific to a package, such as the upper triangular matrix R of the QR decomposition that has been computed for a Jacobian. Since such an object will not be computed by all methods, testing it in nlsCompare makes no sense, and in that package we concentrate on the minimum sum of squares and the associated model parameters.

#### Code and documentation

A summary of our main results:

- nlspkg: a packaged version of the 'nls() code from R-base. Thanks to Duncan Murdoch
- nlsalt: attempt to mirror nls() behaviour entirely in R. This is UNFINISHED. The effort showed that the structure of the programs was difficult to follow, undocumented, and unsuited to adding improvements like the Marquardt stabilization. We were able to get a pure-R version ofnumericDeriv() and rework most of the functions ofnlsModel(but notnlsModel.plinear). This work may continue after the project formally ends, but collaboration is likely needed with workers who have a deep knowledge of R internals.nls-flowchart.txtwas a start at documentation of the structure ofnls():
- the safeguarded relative offset convergence criterion that allows small-residual problems to be handled with nls() is now in the R base code.
- nlsj is a refactoring of some nls() functionality. ?? What are key features?? This is an interim package for exploration of ideas and will NOT, as it currently is presented, become a distributed package.
- nlsralt: this is intended to be a modified version of Nash and Murdoch package nlsr to examine possible improvements discovered as a result of this project. This is EXPERIMENTAL.

### Strategic choices in a nonlinear model estimation

The key difference in approach between nls() and nlsr::nlxb() is that nls() builds a large infrastructure from which the Gauss-Newton iteration can be executed and other statistical information such as profiles can be computed, while nlxb() returns quite limited information, and in its execution computes what is needed on an "as and when" basis. This follows a path that one of us (JN) established almost 50 years ago with the software that became Nash (1979), using setup, solver, and post-solution analysis phases to computation. Here, that last phase is not part of nlxb(), but is the domain of other functions in the nlsr package. nls() also has some similar functions, but they are much more tied into the infrastructure that is created, mostly in the nlsModel() function.

The nls() approach, as implemented in the base R code and nlspkg leads to considerable entangling of the different functions and capabilities. Even after both the preparation for and carrying out of the present project, we do not feel confident to explain the code completely, nor to maintain it. However, we have made some forays into render parts of the code in R, and to understand where difficulties lie.

#### Opinion for discussion

To advance the stability and maintainability of R, we believe the program objects (R functions) that are created by tools such as nls() should have minimal interactions and side-effects. The aspects of nls() that have given us the most trouble are as follows:

• The functions that compute the residuals and jacobians are frequently computed by presuming the current data and parameters for those functions are available in an environment. As long as the correct

- environment is used, this provides a surprisingly short syntax to invoke the calculations. The danger is that the wrong data is accessed if the internal search finds a valid name that is not the object we want.
- Weights, subsets, and various contextual controls such as that for the na.action that tell our code what to do with undefined or missing numbers. Again, this makes for a very simple invocation of the calculation, but the context is hidden from the user. It can be difficult for those of us trying to maintain or improve the code to be certain we have the context correct.
- The mixing of R and C code made for high performance in the past, but without developer documentation, programmers face a lot of work to fix code. We believe in keeping code at least a runnable reference version in a single programming language. If necessary, by measuring ("profiling") the code, we can find bottlenecks and replace just those slower parts of the reference code.
- We believe that a structure that isolates the setup, solve, and post-solution structures for complicated
  functions reduces the number of objects that must be kept in appropriate alignment at any one stage in
  a set of calculations.
- We feel that all codes should return the best fit they have found so far, even if there are untoward conditions reached, such as a singular Jacobian. This modification could be made to nls() even in the short term.

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