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May 6, 2014

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1 Introduction

1.1 Background

(This section is based on (Jardim, et.al, 2014))

The volume and availability of data useful for fisheries stock assessment is continually increasing. Time series of 'traditional' sources of information, such as surveys and landings data are not only getting longer, but also cover an increasing number of species.

For example, in Europe the 2009 revision of the Data Collection Regulation (EU, 2008a) has changed the focus of fisheries sampling programmes away from providing data for individual assessment of 'key' stocks (i.e. those that are economically important) to documenting fishing trips, thereby shifting the perspective to a large coastal monitoring programme. The result has been that data on growth and reproduction of fish stocks are being collected for more than 300 stocks in waters where the European fleets operate.

Recognizing that the context above required new methodological developments, the European Commission Joint Research Centre (JRC) started its 'Assessment for All' Initiative (a4a), with the aim to develop, test, and distribute methods to assess a large numbers of stocks in an operational time frame, and to build the necessary capacity/expertise on stock assessment and advice provision.

The long-term strategy of a4a is to increase the number of stock assessments by reducing the workload required to run each analysis and by bringing more scientists/analysts into fisheries management advice. The first is achieved by developing a working framework with the methods required to run all the analysis a stock assessment needs, as well as developing methods to deal with recognized bottlenecks, e.g. model averaging to deal with model selection (Colin, et.al, 2014). Such an approach should make the model exploration and selection processes easier, as well as decreasing the burden of moving between software platforms. The second can be achieved by making the analysis more intuitive, thereby attracting more experts to join stock assessment teams.

To achieve these objectives, the Initiative identified a series of tasks, which were or are being carried out, namely:

- define a moderate data stock;
- develop a stock assessment framework;
- develop a forecasting algorithm based on MSE;
- organize training courses for marine scientists.

1.1.1 The moderate data stock

The moderate data stock definition was an important step in the Initiative's development. It clearly focused the initiative on stocks with some information, moving away from the data-poor stocks, but without moving into data rich methodologies. It was recognized that there's a lot of research on both extremes of the data availability spectrum, but comparatively little in the middle 'region'. From this came the idea of the 'moderate data stock'.

The 'moderate data stock' constitutes the entry level of our analysis. It has at least the following datasets, which can be assembled in different ways, using distinct methods.

- in relation to exploitation:
 - volume of catches, which may be split in landings and discards, or not;
 - length frequencies of the catches, landings or discards;
 - nominal effort (optional, needed in case CPUE indices are to be derived);
- in relation to biology:
 - estimate of maturity ogive (e.g. L_{50});

- estimates of growth model and parameters;
- length-weight relationship;
- in relation to abundance:
 - index of abundance.

1.1.2 The stock assessment framework

The stock assessment model framework is a non-linear catch-at-age model implemented in R/FLR/ADMB that can be applied rapidly to a wide range of situations with low parametrization requirements. Later we'll come back to these characteristics and it's application (Section ??).

1.1.3 MSE

The MSE is a sophisticated forecasting algorithm that takes into account structural uncertainty about stock dynamics (growth, recruitment, maturity) and on exploitation by commercial fleets (selectivity), embedding the framework of decision making.

1.1.4 Training

During the last 2 years JRC organized 4 courses of introduction to R and FLR: Varese, January 2012; Varese, June, 2012; Barza, March 2013; FAO / GFCM, Rome, November 2013.

In 2013 a short course about a4a methods was organized in Lisbon. The first full course on FLR and a4a methods was organized in CEFAS, March 2014 and another one is planned for August 2014.

These courses are open to all participants and don't have an attendance fee.

1.2 The a4a approach to stock assessment and management advice

The approach presented here is split in 4 steps: (i) converting length data to age data using a growth model, (ii) modelling natural mortality, (iii) assessing the stock, and (iv) MSE¹.

These steps may be followed in sequence or independently, depending on the user's preferences. All that is needed is to use the objects provided by the previous step and provide the objects required by the next, so that data flows between steps smoothly. One can make the analogy with building with Lego, where for each layer the builder may use the pieces provided by a particular boxset, or make use of pieces from other boxsets. Figure 1 shows the process, including the class of the objects that carry the data (in black).

¹Under development, to be released with version 2.0, scheduled for the fourth quarter of 2014

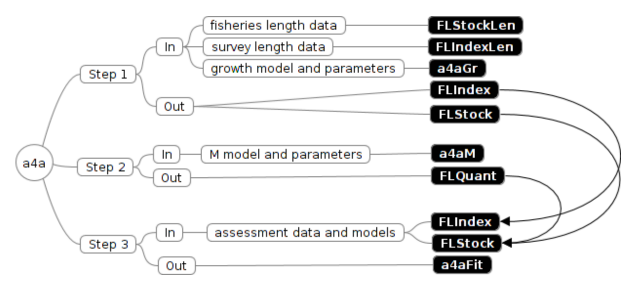


Figure 1: In/out process of the a4a approach. The boxes in black represent the classes of the objects that carry the information for each step and out of each step.

Analysis related to projections and biological reference points are dealt with by the FLR packages FLash and FLBRP. As such the Initiative does not provide specific methods for these analysis.

In Steps 1 and 2 there is no fitting of growth models or natural mortality models. The rationale is to provide tools that allow the uncertainty associated with these processes to be carried on into the stock assessment, e.g. through parameter uncertainty. This approach allows the users to pick up the required information from other sources of information such as papers, PhDs, Fishbase, other stocks, etc. Bearing in mind that the stock under analysis may not have specific information on the growth or natural mortality processes, but generic information about life history invariants can be used, like the generic priors suggested by Bentley, (2014).

Note that an environment like the one distributed by a4a promotes the exploration of different models for each process, giving the analyst a large flexibility, but also opens the possibility to efficiently include distinct models in the analysis. For example a stock assessment using two growth, or several models for natural mortality. Our suggestion to streamline the assessment process is to combine the final outcome using model averaging (Colin, et.al, 2014) but other solutions may be implemented, like scenario analysis, etc. What's important is to keep the data flowing smoothly and the models clear. R (R Core Team, 2014) and FLR (Kell, et.al, 200) are powerful platforms for it.

2 Loading libraries, data and defining some useful functions

```
library(FLa4a)
library(XML)
library(reshape2)
library(diagram)
data(ple4)
data(ple4.indices)
data(rfLen)
```

```
# functions for transforming the data

# quant 2 quant
qt2qt <- function(object, id = 5, split = "-") {
    qt <- object[, id]
    levels(qt) <- unlist(lapply(strsplit(levels(qt), split = split), "[[", 2))
    as.numeric(as.character(qt))</pre>
```

```
}
# check import and massage
cim <- function(object, n, wt, hrv = "missing") {</pre>
    v <- object[sample(1:nrow(object), 1), ]</pre>
    c1 < c(n[as.character(v$V5), as.character(v$V1), 1, as.character(v$V2)] ==
        v$V6)
    c2 \leftarrow c(wt[as.character(v$V5), as.character(v$V1), 1, as.character(v$V2)] ==
        v$V7)
    if (missing(hrv)) {
        c1 + c2 == 2
    } else {
        c3 \leftarrow c(hrv[as.character(v$V5), as.character(v$V1), 1, as.character(v$V2)] ==
             v$V8)
        c1 + c2 + c3 == 3
    }
}
# and a plot for later
plotS4 <- function(object, linktext = "typeof", main = "S4 class", ...) {</pre>
    args <- list(...)</pre>
    obj <- getClass(as.character(object))</pre>
    df0 <- data.frame(names(obj@slots), unlist(lapply(obj@slots, "[[", 1)))</pre>
    nms \leftarrow c(t(df0))
    nslts <- length(nms)/2
    M <- matrix(nrow = length(nms), ncol = length(nms), byrow = TRUE, data = 0)
    for (i in 1:nslts) {
        M[i * 2, i * 2 - 1] \leftarrow linktext
    }
    args$A = M
    args$pos = rep(2, length(nms)/2)
    args$name = nms
    args$main = main
    do.call("plotmat", args)
}
```

3 Reading files and building FLR objects

For this document we'll use the plaice in ICES area IV dataset, provided by FLR, and a length-based simulated dataset based on red fish, using Gadget, provided by Daniel Howell (Institute of Marine Research, Norway).

In this section we read in the Gadget data files, and transform them into FLR objects.

First we read in the files as data frames and recode some variables.

```
# catch
cth.orig <- read.table("data/catch.len", skip = 5)

# stock
stk.orig <- read.table("data/red.len", skip = 4)

# surveys
idx.orig <- read.table("data/survey.len", skip = 5)
idxJmp.orig <- read.table("data/jump.survey.len", skip = 5)
idxTrd.orig <- read.table("data/tend.survey.len", skip = 5)</pre>
```

```
# Recode the length categories into something usable

# catch
cth.orig[, 5] <- qt2qt(cth.orig)

# stock
stk.orig[, 5] <- qt2qt(stk.orig)

# surveys
idx.orig[, 5] <- qt2qt(idx.orig)
idxJmp.orig[, 5] <- qt2qt(idxJmp.orig)
idxTrd.orig[, 5] <- qt2qt(idxTrd.orig)</pre>
```

Then we reshape the data frames into six dimensional arrays using cast from package reshape2.

```
# catch
cth.n \leftarrow acast(V5 \sim V1 \sim 1 \sim V2 \sim 1 \sim 1, value.var = "V6", data = cth.orig)
cth.wt <- acast(V5 ~ V1 ~ 1 ~ V2 ~ 1 ~ 1, value.var = "V7", data = cth.orig)
hrv <- acast(V5 ~ V1 ~ 1 ~ V2 ~ 1 ~ 1, value.var = "V8", data = cth.orig)</pre>
# stock
stk.n \leftarrow acast(V5 \sim V1 \sim 1 \sim V2 \sim 1 \sim 1, value.var = "V6", data = stk.orig)
stk.wt <- acast(V5 ~ V1 ~ 1 ~ V2 ~ 1 ~ 1, value.var = "V7", data = stk.orig)
# surveys
idx.n \leftarrow acast(V5 \sim V1 \sim 1 \sim V2 \sim 1 \sim 1, value.var = "V6", data = idx.orig)
idx.wt <- acast(V5 ~ V1 ~ 1 ~ V2 ~ 1 ~ 1, value.var = "V7", data = idx.orig)
idx.hrv <- acast(V5 ~ V1 ~ 1 ~ V2 ~ 1 ~ 1, value.var = "V8", data = idx.orig)
idxJmp.n <- acast(V5 ~ V1 ~ 1 ~ V2 ~ 1 ~ 1, value.var = "V6", data = idxJmp.orig)</pre>
idxJmp.wt <- acast(V5 ~ V1 ~ 1 ~ V2 ~ 1 ~ 1, value.var = "V7", data = idxJmp.orig)
idxJmp.hrv <- acast(V5 ~ V1 ~ 1 ~ V2 ~ 1 ~ 1, value.var = "V8", data = idxJmp.orig)
idxTrd.n \leftarrow acast(V5 \sim V1 \sim 1 \sim V2 \sim 1 \sim 1, value.var = "V6", data = idxTrd.orig)
idxTrd.wt <- acast(V5 ~ V1 ~ 1 ~ V2 ~ 1 ~ 1, value.var = "V7", data = idxTrd.orig)
idxTrd.hrv <- acast(V5 ~ V1 ~ 1 ~ V2 ~ 1 ~ 1, value.var = "V8", data = idxTrd.orig)
```

We take the arrays and make FLQuant objects from them.

```
# catch
dnms <- dimnames(cth.n)</pre>
names(dnms) <- names(dimnames(FLQuant()))</pre>
names(dnms)[1] <- "len"</pre>
cth.n <- FLQuant(cth.n, dimnames = dnms)</pre>
cth.wt <- FLQuant(cth.wt, dimnames = dnms)</pre>
hrv <- FLQuant(hrv, dimnames = dnms)</pre>
units(hrv) <- "f"
# stock
dnms <- dimnames(stk.n)</pre>
names(dnms) <- names(dimnames(FLQuant()))</pre>
names(dnms)[1] <- "len"</pre>
stk.n <- FLQuant(stk.n, dimnames = dnms)</pre>
stk.wt <- FLQuant(stk.wt, dimnames = dnms)</pre>
# surveys
dnms <- dimnames(idx.n)</pre>
names(dnms) <- names(dimnames(FLQuant()))</pre>
names(dnms)[1] <- "len"</pre>
```

```
idx.n <- FLQuant(idx.n, dimnames = dnms)
idx.wt <- FLQuant(idx.wt, dimnames = dnms)
idx.hrv <- FLQuant(idx.hrv, dimnames = dnms)

dnms <- dimnames(idxJmp.n)
names(dnms) <- names(dimnames(FLQuant()))
names(dnms)[1] <- "len"
idxJmp.n <- FLQuant(idxJmp.n, dimnames = dnms)
idxJmp.wt <- FLQuant(idxJmp.wt, dimnames = dnms)
idxJmp.hrv <- FLQuant(idxJmp.hrv, dimnames = dnms)

dnms <- dimnames(idxTrd.n)
names(dnms) <- names(dimnames(FLQuant()))
names(dnms)[1] <- "len"
idxTrd.n <- FLQuant(idxTrd.n, dimnames = dnms)
idxTrd.wt <- FLQuant(idxTrd.wt, dimnames = dnms)
idxTrd.hrv <- FLQuant(idxTrd.hrv, dimnames = dnms)
idxTrd.hrv <- FLQuant(idxTrd.hrv, dimnames = dnms)</pre>
```

Some sanity checks to check that the resulting objects have matching dimensions.

```
# catch
cim(cth.orig, cth.n, cth.wt, hrv)

## [1] TRUE

# stock
cim(stk.orig, stk.n, stk.wt)

## [1] TRUE

# surveys
cim(idx.orig, idx.n, idx.wt, idx.hrv)

## [1] TRUE

cim(idxJmp.orig, idxJmp.n, idxJmp.wt, idxJmp.hrv)

## [1] TRUE

cim(idxTrd.orig, idxTrd.n, idxTrd.wt, idxTrd.hrv)

## [1] TRUE
```

Finally, we make FLR objects from the data.

4 Converting length data to age

The stock assessment framework is based on age dynamics. Therefore, to use length information it must be processed before it can be used in an assessment. The rationale is that the processing should give the analyst the flexibility to use a range of sources of information, e.g. literature or online databases, to grab information about the species growth model and the uncertainty about the model parameters.

Within the a4a framework this is handled using the a4aGr class. In this section we introduce the a4aGr class and look at the variety of ways that parameter uncertainty can be included.

4.1 a4aGr - The growth class

The conversion of length data to age is performed through the use of a growth model. The implementation is done through the $a \nmid a Gr$ class.

```
showClass("a4aGr")
## Class "a4aGr" [package "FLa4a"]
##
## Slots:
##
## Name:
              grMod grInvMod
                                  params
                                               VCOV
                                                         distr
            formula
                      formula
                                   FLPar
## Class:
                                              array character character
##
## Name:
                desc
                         range
## Class: character
                       numeric
##
## Extends: "FLComp"
```

A simple construction of $a \not a G r$ objects requires the model and parameters to be provided. Check the help file for more information.

Here we show an example using the von Bertalanffy growth model. To create the a4aGr object it's necessary to pass the model equation ($length \sim time$), the inverse model equation ($time \sim length$) and the parameters. Any growth model can be used as long as it's possible to write the equation as an R formula.

```
vb0bj < -a4aGr(grMod = ~linf * (1 - exp(-k * (t - t0))), grInvMod = ~t0 - 1/k *
    log(1 - len/linf), params = FLPar(linf = 58.5, k = 0.086, t0 = 0.001, units = c("cm",
    "ano-1", "ano")))
# Check the model and its inverse
1c = 20
predict(vb0bj, len = lc)
##
      iter
##
##
     1 4.866
predict(vb0bj, t = predict(vb0bj, len = lc)) == lc
##
      iter
##
          1
##
     1 TRUE
```

The predict method allows the transformation between age and lengths.

```
predict(vb0bj, len = 5:10 + 0.5)
##
     iter
##
           1
##
    1 1.149
##
    2 1.371
##
    3 1.596
##
    4 1.827
    5 2.062
##
     6 2.301
predict(vb0bj, t = 5:10 + 0.5)
##
      iter
##
           1
##
     1 22.04
##
     2 25.05
##
    3 27.80
##
    4 30.33
     5 32.66
##
     6 34.78
##
```

4.2 Adding parameter uncertainty with a multivariate normal distribution

Uncertainty in the growth model is introduced through the inclusion of parameter uncertainty. This is done by making use of the parameter variance-covariance matrix (the vcov slot of the a4aGr class) and assuming a distribution. The numbers in the variance-covariance matrix could come from the parameter uncertainty from fitting the growth model parameters.

Here we set the variance-covariance matrix by scaling a correlation matrix, using a cv of 0.2.

```
# Make an empty cor matrix
cm \leftarrow diag(c(1, 1, 1))
# k and linf are negatively correlated while tO is independent
cm[1, 2] \leftarrow cm[2, 1] \leftarrow -0.5
# scale cor to var using CV=0.2
cv <- 0.2
p \leftarrow c(linf = 60, k = 0.09, t0 = -0.01)
vc <- matrix(1, ncol = 3, nrow = 3)</pre>
1 <- vc
1[1, ] \leftarrow 1[, 1] \leftarrow p[1] * cv
k <- vc
k[, 2] \leftarrow k[2, ] \leftarrow p[2] * cv
t <- vc
t[3, ] \leftarrow t[, 3] \leftarrow p[3] * cv
mm \leftarrow t * k * 1
diag(mm) <- diag(mm)^2</pre>
mm <- mm * cm
# check that we have the intended correlation
all.equal(cm, cov2cor(mm))
## [1] TRUE
# Create the a4aGr object as before but now we also include the vcov
# argument for the variance-covariance matrix
```

```
vbObj <- a4aGr(grMod = ~linf * (1 - exp(-k * (t - t0))), grInvMod = ~t0 - 1/k *
        log(1 - len/linf), params = FLPar(linf = p["linf"], k = p["k"], t0 = p["t0"],
        units = c("cm", "ano-1", "ano")), vcov = mm)</pre>
```

First we show a simple example where we assume that the parameters are represented using a multivariate normal distribution.

```
# Note that the object we have just created has a single iteration of each
# parameter
vb0bj@params
## An object of class "FLPar"
## params
## linf
             k
                  t.0
## 60.00 0.09 -0.01
## units: cm ano-1 ano
dim(vbObj@params)
## [1] 3 1
# We simulate 10000 iterations from the a4aGr object by calling mvrnorm()
# using the the variance-covariance matrix we created earlier.
vbNorm <- mvrnorm(10000, vb0bj)</pre>
# Now we have 10000 iterations of each parameter, randomly sampled from the
# multivariate normal distribution
vbNorm@params
## An object of class "FLPar"
## iters: 10000
##
## params
##
                   linf
## 60.0781351(11.96020) 0.0900066( 0.01809) -0.0099972( 0.00198)
## units: cm ano-1 ano
dim(vbNorm@params)
## [1]
           3 10000
```

We can now convert from length to ages data based on the 10000 parameter iterations. This gives us 10000 sets of ages data. For example, here we convert a single length vector:

```
ages <- predict(vbNorm, len = 5:10 + 0.5)</pre>
dim(ages)
## [1]
           6 10000
# We show the first ten iterations only as an illustration
ages[, 1:10]
##
      iter
##
                   2
                          3
                                 4
                                       5
                                               6
                                                     7
                                                           8
                                                                       10
             1
     1 0.9769 1.338 0.9591 1.158 1.497 0.9315 1.121 1.149 0.906 1.585
##
```

```
## 2 1.1661 1.600 1.1459 1.383 1.793 1.1114 1.336 1.372 1.083 1.912

## 3 1.3587 1.868 1.3363 1.613 2.097 1.2941 1.554 1.599 1.262 2.253

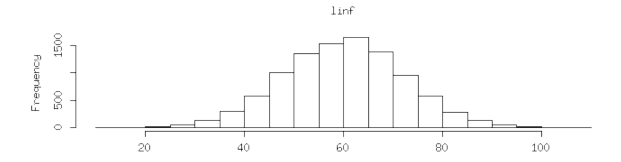
## 4 1.5549 2.141 1.5304 1.846 2.408 1.4795 1.775 1.830 1.446 2.609

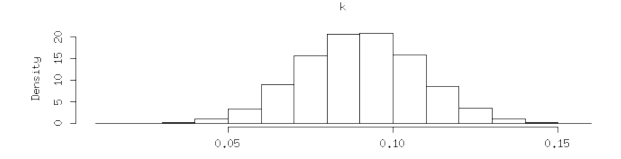
## 5 1.7546 2.421 1.7284 2.085 2.727 1.6678 2.000 2.065 1.633 2.982

## 6 1.9582 2.707 1.9305 2.328 3.056 1.8590 2.228 2.305 1.823 3.372
```

The marginal distributions can be seen in Figure 2.

```
par(mfrow = c(3, 1))
hist(c(params(vbNorm)["linf", ]), main = "linf", xlab = "")
hist(c(params(vbNorm)["k", ]), main = "k", prob = TRUE, xlab = "")
hist(c(params(vbNorm)["t0", ]), main = "t0", xlab = "")
```





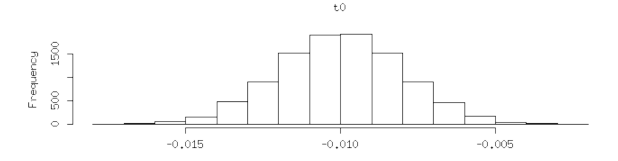


Figure 2: The marginal distributions of each of the parameters from using a multivariate normal distribution.

The shape of the correlation can be seen in Figure 3.

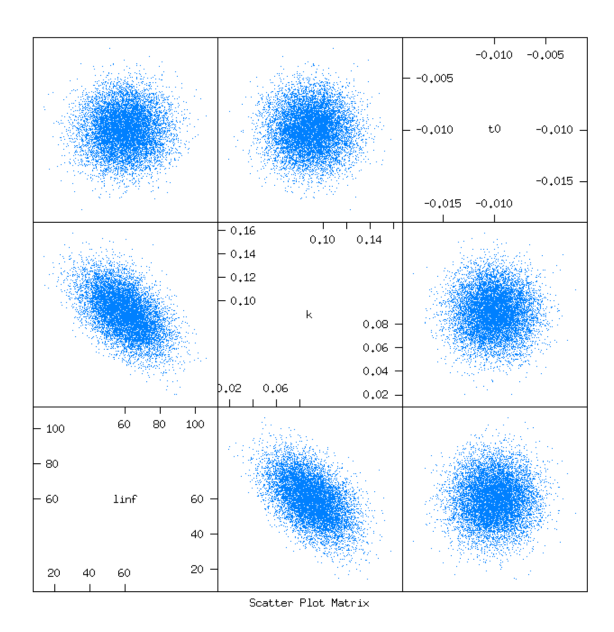


Figure 3: Scatter plot of the 10000 samples parameter from the multivariate normal distribution.

Growth curves for the 1000 iterations can be seen in Figure 4.

boxplot(t(predict(vbNorm, t = 0:50 + 0.5)))

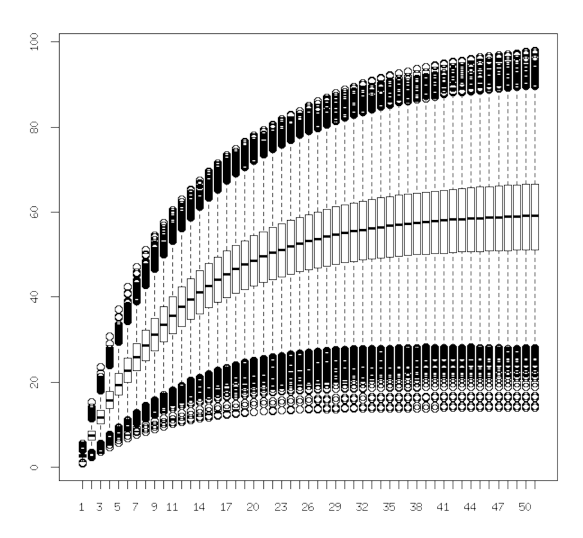


Figure 4: Growth curves using parameters simulated from a multivariate normal distribution.

4.3 Adding parameter uncertainty with a multivariate triangle distribution

One alternative to using normal distributions is to use a triangle distribution. We use the package triangle, where this distribution is parametrized using the minimum, maximum and median values. This can be very attractive if the analyst needs to scrape information from the web or literature and perform some kind of meta-analysis.

Here we show an example of setting a triangle distribution with values taken from Fishbase.

```
# The web address for the growth parameters for redfish (Sebastes
# norvegicus)
addr <- "http://www.fishbase.org/PopDyn/PopGrowthList.php?ID=501"
# Scrape the data
tab <- try(readHTMLTable(addr))
# Interrogate the data table and get vectors of the values
linf <- as.numeric(as.character(tab$dataTable[, 2]))</pre>
```

The marginals will reflect the uncertainty on the parameter values that were scraped from Fishbase but, as we don't really believe the parameters are multivariate normal, here we adopted a distribution based on a t copula with triangle marginals. The marginal distributions can be seen in Figure 5 and the shape of the correlation can be seen in Figure 6.

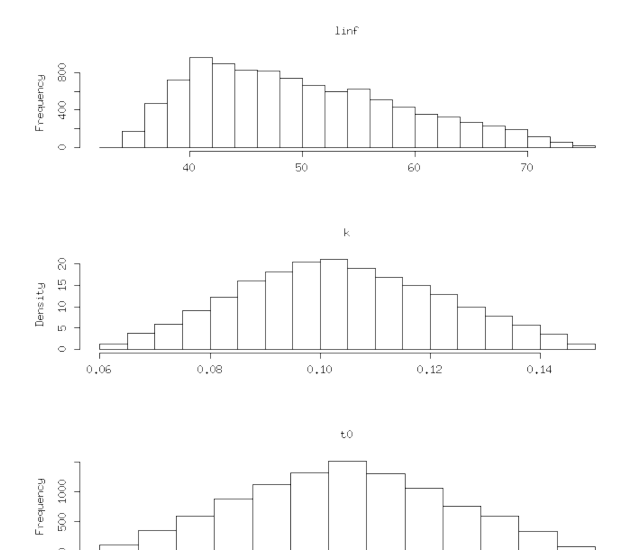


Figure 5: The marginal distributions of each of the parameters from using a multivariate triangle distribution.

-1.2

-1.0

-0.8

-0.6

-1.8

-1.6

-1.4

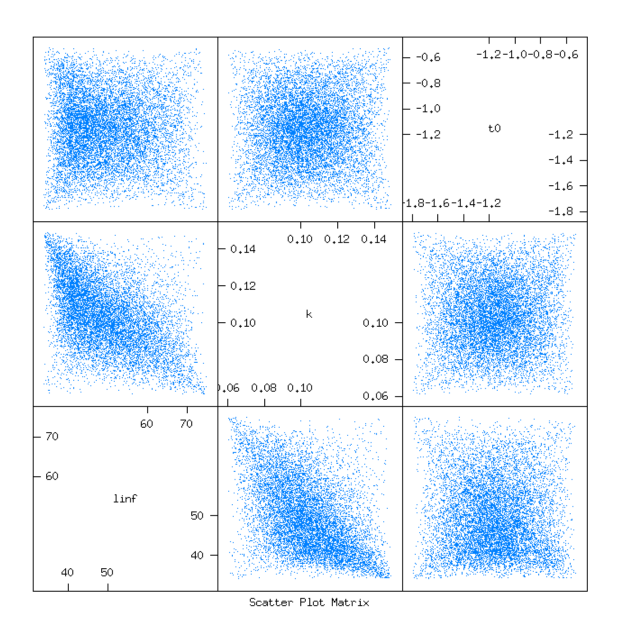


Figure 6: Scatter plot of the 10000 samples parameter from the multivariate triangle distribution.

We can still use predict() to get see the growth model uncertainty (Figure 7).

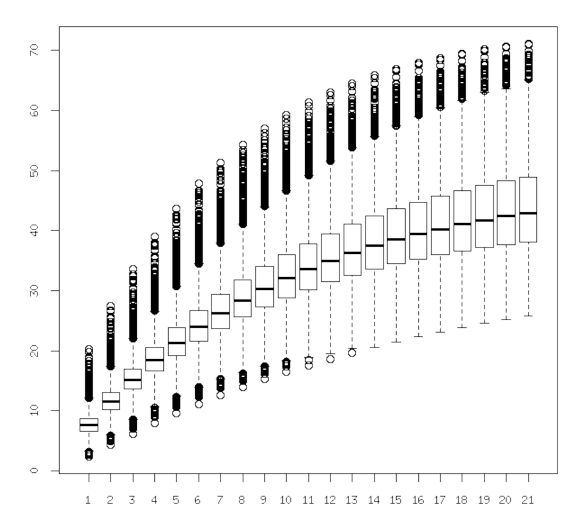


Figure 7: Growth curves using parameters simulated from a multivariate triangle distribution.

Remember that the above examples use a variance-covariance matrix that we essentially made up. An alternative would be to scrape the entire growth parameters dataset from FishBase and compute the shape of the variance-covariance matrix yourself.

4.4 Adding parameter uncertainty with copulas

A more general approach to adding parameter uncertainty is to make use of statistical copulas² and marginal distributions of choice. This is possible with the mvrcop() function borrowed from the package copula. The example below keeps the same parameters and changes only the copula type and family but a lot more can be done. Check the package copula for more.

```
vbCop <- mvrcop(10000, vbObj, copula = "archmCopula", family = "clayton", param = 2,
    margins = "triangle", paramMargins = triPars)</pre>
```

²http://en.wikipedia.org/wiki/Copula_%28probability_theory%29

The shape of the correlation changes (Figure 8) as well as the resulting growth curves (Figure 9).

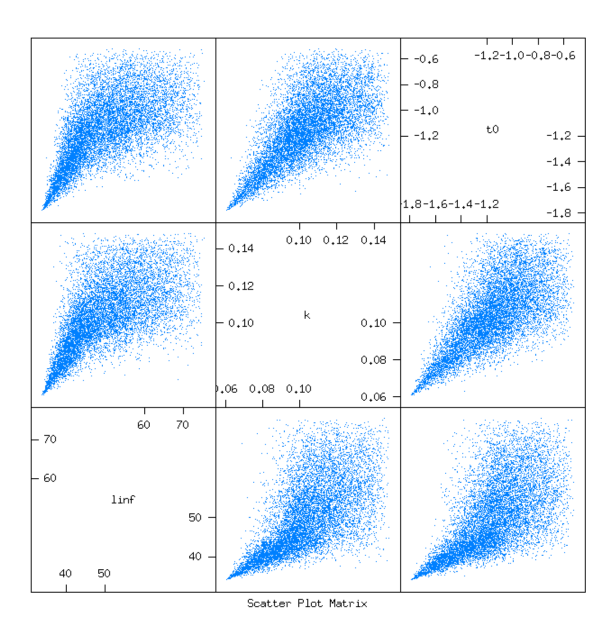


Figure 8: Scatter plot of the 10000 samples parameter from the using an archmCopula copula with triangle margins.

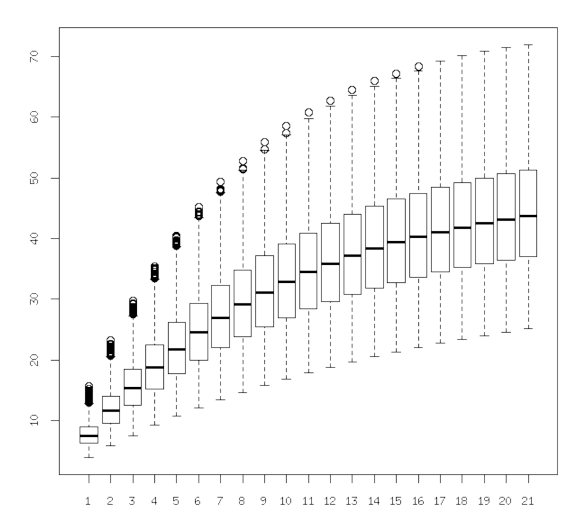


Figure 9: Growth curves from the using an archmCopula copula with triangle margins.

4.5 The 12a() method

After introducing uncertainty in the growth model through the parameters it's time to transform the length-based dataset into an age-based dataset. The method that deals with this process is 12a(). The implementation of this method for the FLQuant class is the main workhorse. There's two other implementations, for the FLStock and FLIndex classes, which are mainly wrappers that call the FLQuant method several times.

When converting from length-based data to age-based data you need to be aware of how the aggregation of length classes is performed. For example, individuals in length classes 1-2, 2-3, and 3-4 cm may all be considered as being of age 1 (obviously depending on the growth model). How should the values in those length classes be combined?

If the values are abundances then the values should be summed. Summing other types of values such as weights does not make sense. Instead these values are averaged over the length classes (weighted by the abundance for weights). This is controlled using the stat argument which can be either mean or sum (the

default). Fishing mortality is not computed to avoid making wrong assumptions about the meaning of F at length.

We demonstrate the method by converting a catch-at-length FLQuant to a catch-at-age FLQuant. First we make an $a \not a Gr$ object with a multivariate triangle distribution. We use 10 iterations as an example. And call 12a() by passing in the length-based FLQuant and the $a \not a Gr$ object.

```
vbTriSmall <- mvrtriangle(10, vb0bj, paramMargins = triPars)
cth.n <- 12a(catch.n(rfLen.stk), vbTriSmall)</pre>
```

```
dim(cth.n)
## [1] 79 26 1 4 1 10
```

In the previous example, the FLQuant object that was sliced (catch.n(rfLen.stk)) had only one iteration. This iteration was sliced by each of the iterations in the growth model. It is possible for the FLQuant object to have the same number of iterations as the growth model, in which case each iteration of the FLQuant and the growth model are used together. It is also possible for the growth model to have only one iteration while the FLQuant object has many iterations. The same growth model is then used for each of the FLQuant iterations. As with all FLR objects, the general rule is one or n iterations.

As well as converting one FLQuant at a time, we can convert entire FLStock and FLIndex objects. In these cases the individual FLQuant slots of those classes are converted from length-based to age-based. As mentioned above, the aggregation method depends on the type of values the slots contain. The abundance slots (*.n, such as stock.n) are summed. The *.wt, m, mat, harvest.spwn and m.spwn slots of an FLStock object are averaged. The index, catch.wt, index.var, sel.pattern and index.q slots of an FLIndex object are averaged³.

The method for FLStock classes takes an additional argument for the plusgroup.

```
aStk <- 12a(rfLen.stk, vbTriSmall, plusgroup = 14)
## [1] "maxfbar has been changed to accomodate new plusgroup"
aIdx <- 12a(rfTrawl.idx, vbTriSmall)</pre>
```

When converting with 12a() all lengths above Linf are converted to the maximum age, as there is no information in the growth model about how to deal with individuals larger than Linf.

³Still working on l2a for index. Not all of these slots can be averaged

5 Natural mortality

Natural mortality is dealt with as an external parameter to the stock assessment model. The rationale is similar to that of growth: one should be able to grab information from a range of sources and feed it into the assessment.

The mechanism used by a4a is to build an interface that makes it transparent, flexible and hopefully easy to explore different options. In relation to natural mortality it means that the analyst should be able to use distinct models like Gislasson's, Charnov's, Pauly's, etc in a coherent framework making it possible to compare the outcomes of the assessment.

Within the a4a framework, the general method for inserting natural mortality in the stock assessment is to:

- Create an object of class a4aM which holds the model and parameters to be used to generate the natural mortality.
- Add uncertainty to the parameters in the a4aM object.
- Apply the m() method to the a4aM object to create an age or length based FLQuant object of the required dimensions.

The resulting FLQuant object can then be directly inserted into an FLStock object to be used for the assessment.

In this section we go through each of the steps in detail using a variety of different models.

5.1 a4aM - The M class

Natural mortality is implemented in a class named $a \not= aM$. This class is made up of three models of the class FLModelSim. Each model represents one effect: an age or length effect, a scaling (level) effect and a time trend, named shape, level and trend, respectively. The impact of the models is multiplicative, i.e. the overal natural mortality is given by $shape \times level \times trend$. Check the help files for more information.

```
showClass("a4aM")
## Class "a4aM" [package "FLa4a"]
##
## Slots:
##
## Name:
                shape
                           level
                                       trend
                                                   name
                                                               desc
                                                                         range
## Class: FLModelSim FLModelSim FLModelSim character
                                                         character
                                                                       numeric
##
## Extends: "FLComp"
```

The $a \nmid aM$ constructor requires that the models and parameters are provided. The default method will build each of these models as a constant value of 1. For example the usual "0.2" guessestimate could be set up by setting the level model to have a single parameter with a fixed value, while the other two models, shape and trend, have a default value of 1 (effectively, they have no effect).

```
mod02 <- FLModelSim(model = ~a, params = FLPar(a = 0.2))
m1 <- a4aM(level = mod02)
m1

## a4aM object:
## shape: ~1
## level: ~a
## trend: ~1</pre>
```

More interesting natural mortality shapes can be set up using biological knowledge. The following example uses an exponential decay over ages (implying that resulting FLQuant generated by the m() method will be age based). We also use Jensen's second estimator (Kenshington, 2013) as a scaling level model, which is based on the von Bertalanffy K parameter, M = 1.5K.

```
shape2 <- FLModelSim(model = ~exp(-age - 0.5))
level2 <- FLModelSim(model = ~1.5 * k, params = FLPar(k = 0.4))
m2 <- a4aM(shape = shape2, level = level2)
m2

## a4aM object:
## shape: ~exp(-age - 0.5)
## level: ~1.5 * k
## trend: ~1</pre>
```

Note that the shape model has age as a parameter of the model but is not set using the params argument.

The shape model does not have to be age-based. For example, here we set up a shape model using Gislason's second estimator (Kenshington, 2013): $M_l = K(\frac{L_i n f}{l})^{1.5}$. We use the default level and trend models.

Check a length-base shape model.

```
shape_len <- FLModelSim(model = ~K * (linf/len)^1.5, params = FLPar(linf = 60,
        K = 0.4))
m_len <- a4aM(shape = shape_len)</pre>
```

As an alternative, an external factor may impact the natural mortality. This can be added through the trend model. Suppose M can be modelled with a dependency on the NAO index, due to some mechanism that results in having lower M when NAO is negative and higher when it's positive. The impact is represented by the NAO value on the quarter before spawning, which occurs in the second quarter.

We use this to make a complicated natural mortality model with an age based shape model, a level model based on K and a trend model driven by NAO, where M increases 50% if NAO is positive on the first quarter.

```
# Get NAO
nao.orig <- read.table("http://www.cdc.noaa.gov/data/correlation/nao.data",</pre>
    skip = 1, nrow = 62, na.strings = "-99.90")
dnms <- list(quant = "nao", year = 1948:2009, unit = "unique", season = 1:12,
    area = "unique")
# Build an FLQuant from the NAO data
nao.flq <- FLQuant(unlist(nao.orig[, -1]), dimnames = dnms, units = "nao")</pre>
# Build covar by calculating mean over the first 3 months
nao <- seasonMeans(nao.flq[, , , 1:3])</pre>
# Turn into Boolean
nao \leftarrow (nao > 0)
# Constructor
trend3 <- FLModelSim(model = ~1 + b * nao, params = FLPar(b = 0.5))</pre>
shape3 <- FLModelSim(model = ~exp(-age - 0.5))</pre>
level3 <- FLModelSim(model = ~1.5 * k, params = FLPar(k = 0.4))</pre>
m3 <- a4aM(shape = shape3, level = level3, trend = trend3)
m3
## a4aM object:
     shape: ~exp(-age - 0.5)
##
    level: ~1.5 * k
## trend: ~1 + b * nao
```

5.2 Adding multivariate normal parameter uncertainty

Uncertainty on natural mortality is added through uncertainty on the parameters. In the case of the a4aM class it makes use of the class FLModelSim method mvrnorm(), which is a wrapper for the method mvrnorm() distributed by the package MASS.

```
shape4 <- FLModelSim(model = ~exp(-age - 0.5))</pre>
level4 <- FLModelSim(model = ^{k}0.66 * t0.57, params = FLPar(k = 0.4, t = 10),
    vcov = array(c(0.002, 0.01, 0.01, 1), dim = c(2, 2)))
trend4 <- FLModelSim(model = ~1 + b * nao, params = FLPar(b = 0.5), vcov = matrix(0.02))
m4 <- a4aM(shape = shape4, level = level4, trend = trend4)
m4 <- mvrnorm(100, m4)
m4
## a4aM object:
##
     shape: ^{\sim}exp(-age - 0.5)
##
     level: ^{^{\sim}}k^{^{\sim}}0.66 * t^{^{\sim}}0.57
##
     trend: ^{\sim}1 + b * nao
# Look at the level model (for example)
m4@level
## An object of class "FLModelSim"
## Slot "model":
## ~k^0.66 * t^0.57
##
## Slot "params":
## An object of class "FLPar"
## iters: 100
##
## params
##
## 0.40071(0.0439) 10.01613(0.9546)
## units: NA
##
## Slot "vcov":
##
         [,1] [,2]
## [1,] 0.002 0.01
## [2,] 0.010 1.00
##
## Slot "distr":
## [1] "norm"
# Note the variance in the parameters. The trend model also has uncertainty
params(trend(m4))
## An object of class "FLPar"
## iters: 100
##
## params
## 0.50226(0.131)
## units: NA
# However, the shape model has no parameters and no uncertainty
params(shape(m4))
```

```
## An object of class "FLPar"
## param
##
## NA
## units: NA
```

In this particular case, the **shape** model will not be randomized because it doesn't have a variance covariance matrix. Also note that because there is only one parameter in the **trend** model, the randomization will use a univariate normal distribution.

The same model could be achieved using mnrnorm() on each model component:

```
m4 <- a4aM(shape = shape4, level = mvrnorm(100, level4), trend = mvrnorm(100, trend4))</pre>
```

5.3 Adding parameter uncertainty with copulas

We can also use copulas to add parameter uncertainty to the natural mortality model, similar to the way we use them for the growth model in Section 4.3. As stated above these processes make use of the methods implemented for the FLModelSim class.

In the following example we'll use again Gislason's second estimator, $M_l = K(\frac{L_i n f}{l})^{1.5}$ and a triangle copula to model parameter uncertainty in natural mortality.

```
linf <- 60
k < -0.4
# vcov matrix (make up some values)
mm <- matrix(NA, ncol = 2, nrow = 2)
# 10% cv
diag(mm) \leftarrow c((linf * 0.1)^2, (k * 0.1)^2)
# 0.2 correlation
mm[upper.tri(mm)] <- mm[lower.tri(mm)] <- c(0.05)</pre>
# a good way to check is using cov2cor
cov2cor(mm)
          [,1]
                  [,2]
## [1,] 1.0000 0.2083
## [2,] 0.2083 1.0000
# create object
mgis2 <- FLModelSim(model = ~k * (linf/len)^1.5, params = FLPar(linf = linf,</pre>
    k = k), vcov = mm)
# set the lower, upper and (optionally) centre of the parameters (without
# the centre, the triangle is symmetrical)
pars <- list(list(a = 55, b = 65), list(a = 0.3, b = 0.6, c = 0.35))
mgis2 <- mvrtriangle(1000, mgis2, paramMargins = pars)</pre>
mgis2
## An object of class "FLModelSim"
## Slot "model":
## ~k * (linf/len)~1.5
##
## Slot "params":
## An object of class "FLPar"
## iters: 1000
##
```

```
## params
##
               linf
## 59.87739(2.2710)
                     0.40742(0.0762)
##
  units: NA
##
##
  Slot "vcov":
##
         [,1]
                [,2]
   [1,] 36.00 0.0500
##
   [2,]
        0.05 0.0016
## Slot "distr":
## [1] "un t copula family triangle"
```

The resulting parameter estimates and marginal distributions can be seen in Figure 10 and 11

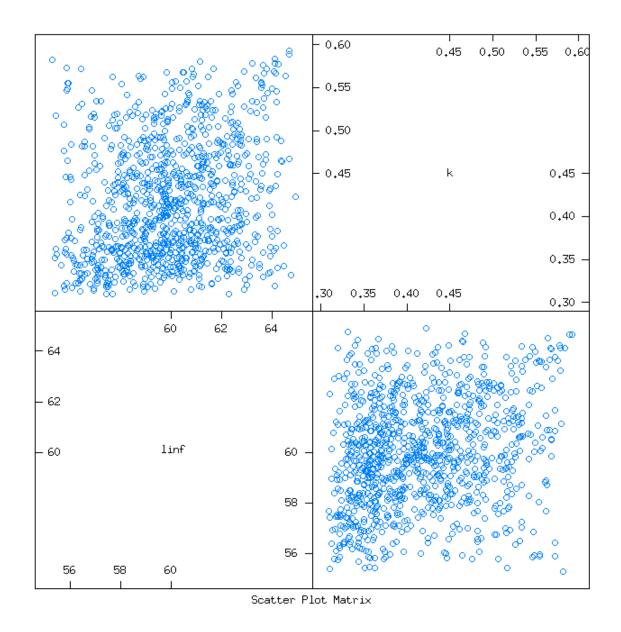
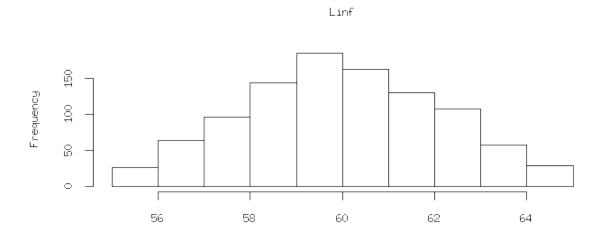


Figure 10: Parameter estimates for Gislason's second natural mortality model from using a triangle distribution.



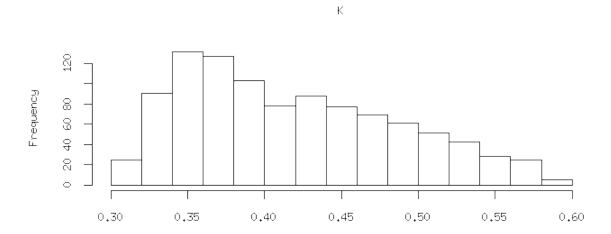


Figure 11: Marginal distributions of the parameters for Gislason's second natural mortality model using a triangle distribution.

We now have a new model that can be used for the shape model. Use the constructor or the set method to add the new model. Note that we have a quite complex method now for M. A length based shape model from Gislason's work, Jensen's third based temperature level and a time trend depending on NAO. All of the component models have uncertainty in their parameters.

```
m5 <- a4aM(shape = mgis2, level = level4, trend = trend4)
# or
m5 <- m4
shape(m5) <- mgis2</pre>
```

5.4 The "m" method

Now that we have set up the natural mortality model and added parameter uncertainty, we are ready to generate the FLQuant of natural mortality. For that we need the m() method.

The m method is the workhorse method for computing natural mortality. The method returns an FLQuant that can be inserted in an FLStock for usage by the assessment method. Note that if the models use age and/or year as terms, the method expects these to be included in the call. If they're not, the method will use the range slot to work out the ages and/or years that should be predicted.

The size of the FLQuant object is determined by the min, max, minyear and maxyear elements of the range slot of the a4aM object. By default the values of these elements are set to 0. Giving an FLQuant with length 1 in the quant and year dimension. The range slot can be set by hand, or by using the rngquant() and rngyear() methods.

The name of the first dimension of the output FLQuant (e.g. 'age' or 'len') is determined by the parameters of the shape model. If it is not clear what the name should be then the name is set to 'quant'.

Using the m method to make an FLQuant of constant natural mortality:

```
# Start with the simplest model
m1
## a4aM object:
     shape: ~1
##
##
     level: ~a
     trend: ~1
# Check the range
range(m1)
##
                   max plusgroup
                                    minyear
                                              maxyear
                                                        minmbar
                                                                   maxmbar
         min
##
           0
                     0
                                        0
                                                    0
                               0
                                                              0
                                                                         0
# Simple - no ages or years
m(m1)
## An object of class "FLQuant"
## , , unit = unique, season = all, area = unique
##
##
        year
## quant 0
       0 0.2
##
##
## units: NA
# Set the quant range
rngquant(m1) <- c(0, 7) # set the quant range</pre>
range(m1)
##
         min
                   max plusgroup
                                    minyear
                                              maxyear
                                                        minmbar
                                                                   maxmbar
##
           0
                                        0
                                                    0
                     7
                                                                         0
                              0
                                                          0
m(m1)
## An object of class "FLQuant"
## , , unit = unique, season = all, area = unique
##
##
        year
## quant 0
##
       0 0.2
##
       1 0.2
```

```
##
      2 0.2
       3 0.2
##
       4 0.2
##
##
       5 0.2
##
       6 0.2
##
       7 0.2
##
## units: NA
# Set the year range too
rngyear(m1) \leftarrow c(2000, 2010) # set the year range
range(m1)
##
                  max plusgroup
                                  minyear
                                            maxyear
                                                     minmbar
        min
                                                               maxmbar
##
                                     2000
                                               2010
                                                           0
m(m1)
## An object of class "FLQuant"
## , , unit = unique, season = all, area = unique
##
##
       year
## quant 2000 2001 2002 2003 2004 2005 2006 2007 2008 2009 2010
             0.2 0.2 0.2 0.2 0.2 0.2 0.2
                                                0.2
                                                    0.2 0.2
##
       0 0.2
##
       1 0.2
             0.2
                  0.2
                      0.2
                            0.2
                                0.2
                                      0.2
                                           0.2
                                                0.2
                                                    0.2
       2 0.2
                  0.2
                      0.2
                            0.2
                                 0.2
                                      0.2
                                           0.2
                                                0.2
##
             0.2
                                                     0.2
##
       3 0.2
             0.2 0.2 0.2
                            0.2
                                0.2
                                      0.2
                                           0.2
                                                0.2
                                                    0.2
##
       4 0.2 0.2 0.2 0.2
                            0.2
                                0.2 0.2
                                           0.2
                                                0.2
                                                    0.2 0.2
##
       5 0.2 0.2 0.2 0.2
                           0.2 0.2 0.2
                                           0.2
                                                0.2
                                                    0.2 0.2
       6 0.2 0.2 0.2 0.2 0.2 0.2 0.2 0.2
##
                                                0.2
                                                    0.2 0.2
##
       7 0.2 0.2 0.2 0.2 0.2 0.2 0.2 0.2
                                                0.2 0.2 0.2
##
## units: NA
# Note the name of the first dimension is 'quant'
```

The next example has an age based shape. As the shape model has 'age' as a variable which is not included in the *FLPar* slot it is used as the name of the first dimension. Note that in this case mbar becames relevant. It's the range of quants (in this case, ages) that is used to compute the mean level. This mean level will match the value given by the level model.

The mbar range can be changed with the rngmbar() method.

Using the m method to make an FLQuant with age varying natural mortality:

```
# Remind ourselves of the model
m2

## a4aM object:
## shape: ~exp(-age - 0.5)
## level: ~1.5 * k
## trend: ~1

# Simple with no ages or years - note that the first dimension is 'age'
m(m2)
```

```
## An object of class "FLQuant"
## , , unit = unique, season = all, area = unique
##
##
     year
## age 0
   0 0.6
##
##
## units: NA
# With ages
rngquant(m2) \leftarrow c(0, 7)
m(m2)
## An object of class "FLQuant"
## , , unit = unique, season = all, area = unique
##
##
     year
## age 0
## 0 0.6000000
## 1 0.22072766
## 2 0.08120117
##
    3 0.02987224
##
    4 0.01098938
## 5 0.00404277
## 6 0.00148725
## 7 0.00054713
##
## units: NA
# With ages and years
rngyear(m2) <- c(2000, 2003)
m(m2)
## An object of class "FLQuant"
## , , unit = unique, season = all, area = unique
##
##
     year
## age 2000
                  2001
                             2002
                                        2003
## 0 0.60000000 0.60000000 0.60000000 0.60000000
## 1 0.22072766 0.22072766 0.22072766
     2 0.08120117 0.08120117 0.08120117 0.08120117
##
    3 0.02987224 0.02987224 0.02987224 0.02987224
##
     4 0.01098938 0.01098938 0.01098938 0.01098938
     5\ 0.00404277\ 0.00404277\ 0.00404277\ 0.00404277
##
     6\;\; 0.00148725\;\; 0.00148725\;\; 0.00148725\;\; 0.00148725
##
     7 0.00054713 0.00054713 0.00054713 0.00054713
##
##
## units: NA
# Note that the level value is:
predict(level(m2))
##
      iter
##
         1
##
   1 0.6
```

```
# Is the same as
m(m2)["0"]
## An object of class "FLQuant"
## , , unit = unique, season = all, area = unique
##
##
     year
## age 2000 2001 2002 2003
## 0 0.6 0.6 0.6 0.6
##
## units: NA
# This is because the mbar range is currently set to '0' and '0'
range(m2)
##
                  max plusgroup
                                  minyear
                                             maxyear
                                                       minmbar
                                                                 maxmbar
         min
##
           0
                              0
                                      2000
                                                2003
# The mean natural mortality value over this range is given by the level
# model We can change the mbar range
rngmbar(m2) \leftarrow c(0, 5)
range(m2)
##
                  max plusgroup
                                  minyear
                                             maxyear
                                                       minmbar
                                                                 maxmbar
                    7
                                      2000
                                                2003
                         0
# This rescales the natural mortality at age:
m(m2)
## An object of class "FLQuant"
## , , unit = unique, season = all, area = unique
##
##
     year
                2001
## age 2000
                          2002
                                    2003
## 0 2.2812888 2.2812888 2.2812888 2.2812888
## 1 0.8392392 0.8392392 0.8392392 0.8392392
    2 0.3087389 0.3087389 0.3087389 0.3087389
##
    3 0.1135787 0.1135787 0.1135787 0.1135787
##
    4 0.0417833 0.0417833 0.0417833 0.0417833
    5 0.0153712 0.0153712 0.0153712 0.0153712
##
    6 0.0056547 0.0056547 0.0056547 0.0056547
    7 0.0020803 0.0020803 0.0020803 0.0020803
##
##
## units: NA
# Check that the mortality over the mean range is the same as the level
# model
quantMeans(m(m2)[as.character(0:5)])
## An object of class "FLQuant"
## , , unit = unique, season = all, area = unique
##
##
       year
## age 2000 2001 2002 2003
   all 0.6 0.6 0.6 0.6
##
## units: NA
```

The next example uses a time trend for the trend model. We use the m3 model we made earlier. The trend model for this model has a covariate, 'nao'. This needs to be passed in to the m() method. The year range of the 'nao' covariate should match that of the range slot.

Using the m method to make an FLQuant with a time trend:

```
# Simple, pass in a single nao value (only one year)
m(m3, nao = 1)
## An object of class "FLQuant"
## , , unit = unique, season = all, area = unique
##
##
      year
## age 0
##
    0 0.9
##
## units: NA
# Set some ages
rngquant(m3) \leftarrow c(0, 7)
m(m3, nao = 0)
## An object of class "FLQuant"
## , , unit = unique, season = all, area = unique
##
##
     year
## age 0
    0 0.60000000
##
##
   1 0.22072766
##
    2 0.08120117
##
    3 0.02987224
    4 0.01098938
##
##
    5 0.00404277
##
    6 0.00148725
##
    7 0.00054713
##
## units: NA
# With ages and years - passing in the NAO data as numeric (1,0,1,0)
rngyear(m3) <- c(2000, 2003)
m(m3, nao = as.numeric(nao[, as.character(2000:2003)]))
## An object of class "FLQuant"
## , , unit = unique, season = all, area = unique
##
##
     year
## age 2000
                  2001
                             2002
                                         2003
    0 0.90000000 0.60000000 0.90000000 0.60000000
    1 0.33109150 0.22072766 0.33109150 0.22072766
##
    2 0.12180175 0.08120117 0.12180175 0.08120117
##
    3 0.04480836 0.02987224 0.04480836 0.02987224
    4 0.01648407 0.01098938 0.01648407 0.01098938
##
    5 0.00606415 0.00404277 0.00606415 0.00404277
##
##
    6 0.00223088 0.00148725 0.00223088 0.00148725
    7 0.00082069 0.00054713 0.00082069 0.00054713
##
## units: NA
```

The final example show how m() can be used to make an FLQuant with uncertainty (see Figure 12). We use the m4 from earlier with uncertainty on the level and trend parameters.

Using the m method to make an FLQuant with uncertainty:

```
# Simple - no time trend but with iterations
m(m4, nao = 1)
## An object of class "FLQuant"
## iters: 100
##
## , , unit = unique, season = all, area = unique
##
##
    year
## age 0
   0 3.0431(0.416)
##
##
## units: NA
dim(m(m4, nao = 1))
##
          year unit season area
                                      iter
      age
##
                                        100
       1
             1
                     1
                           1
                                  1
# With ages
rngquant(m4) \leftarrow c(0, 7)
m(m4, nao = 0)
## An object of class "FLQuant"
## iters: 100
##
## , , unit = unique, season = all, area = unique
##
     year
##
## age 0
    0 2.0260853(0.183187)
##
## 1 0.7453551(0.067391)
## 2 0.2742008(0.024792)
## 3 0.1008728(0.009120)
    4 0.0371090(0.003355)
##
   5 0.0136517(0.001234)
    6 0.0050222(0.000454)
##
##
    7 0.0018476(0.000167)
##
## units: NA
dim(m(m4, nao = 0))
##
           year unit season
                               area
                                      iter
                    1
                         1
                                        100
# With ages and years
rngyear(m4) <- c(2000, 2003)</pre>
m(m4, nao = as.numeric(nao[, as.character(2000:2003)]))
```

```
## An object of class "FLQuant"
## iters: 100
##
## , , unit = unique, season = all, area = unique
##
##
     year
## age 2000
                                              2002
                          2001
##
   0 3.0430775(0.415739) 2.0260853(0.183187) 3.0430775(0.415739)
    1 1.1194856(0.152942) 0.7453551(0.067391) 1.1194856(0.152942)
##
    2 0.4118358(0.056264) 0.2742008(0.024792) 0.4118358(0.056264)
##
    3 0.1515059(0.020698) 0.1008728(0.009120) 0.1515059(0.020698)
##
    4 0.0557359(0.007615) 0.0371090(0.003355) 0.0557359(0.007615)
    5 0.0205041(0.002801) 0.0136517(0.001234) 0.0205041(0.002801)
    6 0.0075430(0.001031) 0.0050222(0.000454) 0.0075430(0.001031)
##
##
    7 0.0027749(0.000379) 0.0018476(0.000167) 0.0027749(0.000379)
##
     year
## age 2003
## 0 2.0260853(0.183187)
## 1 0.7453551(0.067391)
## 2 0.2742008(0.024792)
## 3 0.1008728(0.009120)
## 4 0.0371090(0.003355)
##
    5 0.0136517(0.001234)
##
    6 0.0050222(0.000454)
##
    7 0.0018476(0.000167)
##
## units: NA
dim(m(m4, nao = as.numeric(nao[, as.character(2000:2003)])))
                  unit season area iter
##
     age
           year
              4 1 1
```

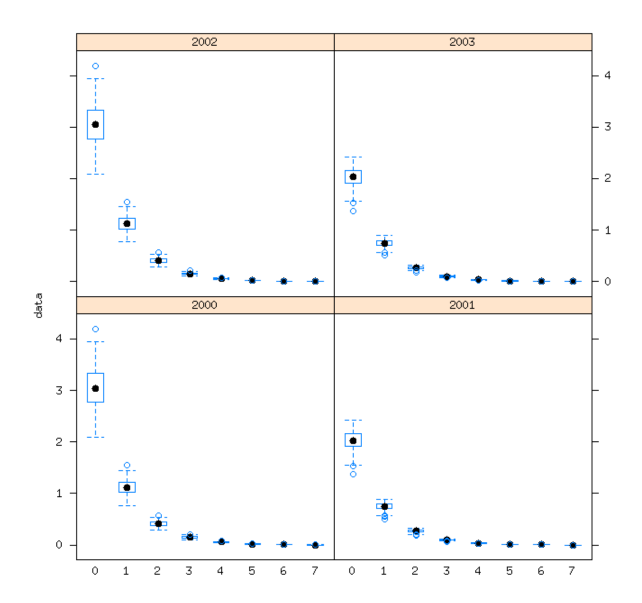


Figure 12: Natural mortality with age and year trend.

6 Running assessments

In the a4a assessment model, the model structure is defined by submodels, which are the different parts of a statistical catch at age model that requires structural assumptions.

There are effectively 5 submodels in operation: the model for F-at-age, a model for the initial age structure, a model for recruitment, a (list) of model(s) for abundance indices catchability-at-age, and a list of models for the observation variance of catch-at-age and abundance indices. In practice, we fix the variance models and the initial age structure models, but in theory these can be changed.

The submodels form is given using linear models, which opens the possibility of using the linear modelling tools available in R: see for example gam formulas, or factorial design formulas using lm. In R's linear modelling language, a constant model is coded as ~ 1 , while a slope over age would simply be $\sim age$. For example we can write a traditional year/age separable F model like $\sim factor(age) + factor(year)$.

There are two basic types of assessments available from using a4a, the management procedure fit and the full assessment fit. The management procedure fit does not compute estimates of covariances and is therefore quicker to execute, while the full assessment fit returns parameter estimates and their covariances at the expense of longer fitting time.

6.1 Stock assessment model details

The statistical catch at age model is based on the well known Baranov catch equation

$$e^{\mathrm{E}[\log C]} = \frac{\mathbf{F}}{\mathbf{F} + M} \left(1 - e^{-\mathbf{F} - M} \right) \mathbf{R} e^{-\sum \mathbf{F} + M}$$

and the survival equation

$$e^{\mathrm{E}[\log I]} = \mathbf{Q} \mathbf{R} e^{-\sum \mathbf{F} + M}$$

where

$$\operatorname{Var}\left[\log C_{ay}\right] = \sigma_{\mathbf{ay}}^{\mathbf{2}} \qquad \operatorname{Var}\left[\log I_{ays}\right] = \tau_{\mathbf{ays}}^{\mathbf{2}}$$

The quantities $\log F$, $\log Q$, $\log R$, $\log observation variances$ and $\log initial age structure$ (in red in the equations above), need to be given form, which is done using linear models. Recruitment is a special case. It's modelled as a fixed variance random effect, using the hard coded models Ricker, Beverton Holt, smooth hockeystick or geometric mean, which can use linear models for their parameters $\log a$ or $\log b$, where relevant. In alternative the $\log R$ submodel can use a linear model like the other submodels

The 'language' of linear models has been developing within the statistical community for many years, and constitutes an elegant way of defining models without going through the complexity of mathematical representations. This approach makes it also easier to communicate among scientists

- 1965 J. A. Nelder, notation for randomized block design
- 1973 Wilkinson and Rodgers, symbolic description for factorial designs
- 1990 Hastie and Tibshirani, introduced notation for smoothers
- 1991 Chambers and Hastie, further developed for use in S

6.2 Quick and dirty

The default settings of the stock assessment model work reasonably well. It's an area of research that will improve with time. Note that because the survey index for plaice has missing values we get a warning saying that we assume these values are missing at random, and not because the observations were zero.

```
data(ple4)
data(ple4.indices)
fit <- sca(ple4, ple4.indices)</pre>
## Note: The following observations are treated as being missing at random:
##
         fleet year age
##
       BTS-Isis 1997 1
##
       BTS-Isis 1997 2
## BTS-Tridens 1997 1
## BTS-Tridens 1997 2
           SNS 1997 1
##
           SNS 1997 2
##
##
           SNS 2003 1
##
           SNS 2003 2
##
           SNS 2003 3
        Predictions will be made for missing observations.
##
```

The residuals() method will compute standardized residuals which can be plotted using a set of packed methods.

```
res <- residuals(fit, ple4, ple4.indices)</pre>
```

Figure 13 shows a scatterplot of residuals by age and survey, with a smoother to guide (or mis-guide ...) your visual analysis.

```
plot(res, main = "Residuals")
```

log residuals of catch and abundance indices

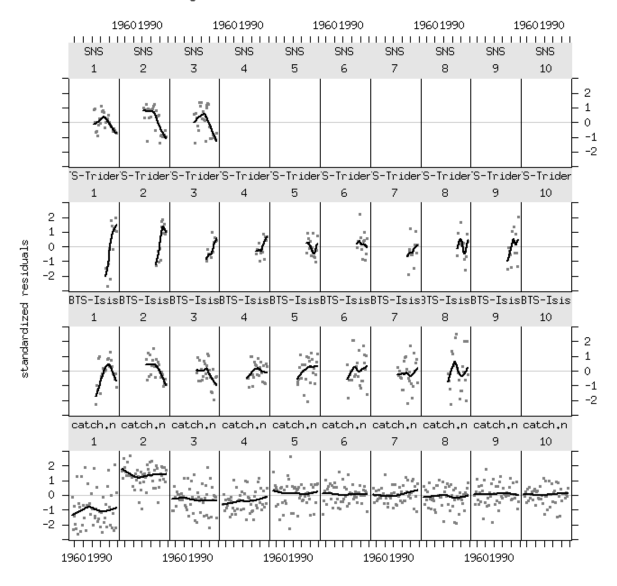


Figure 13: Standardized residuals

The common bubble plot by year and age for each survey are shown in Figure 14.

bubbles(res)

log residuals of catch and abundance indices

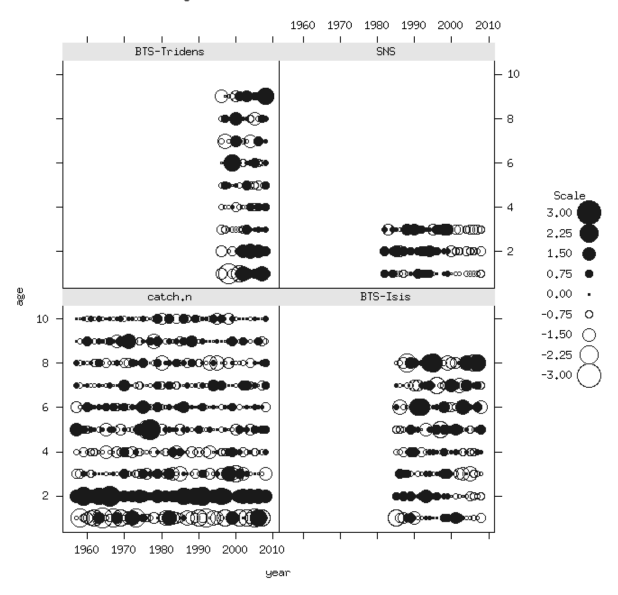


Figure 14: Bubbles plot of standardized residuals.

Finally, Figure ?? shows a quantile-quantile plot to assess how well do the residuals match the normal distribution.

qqmath(res)



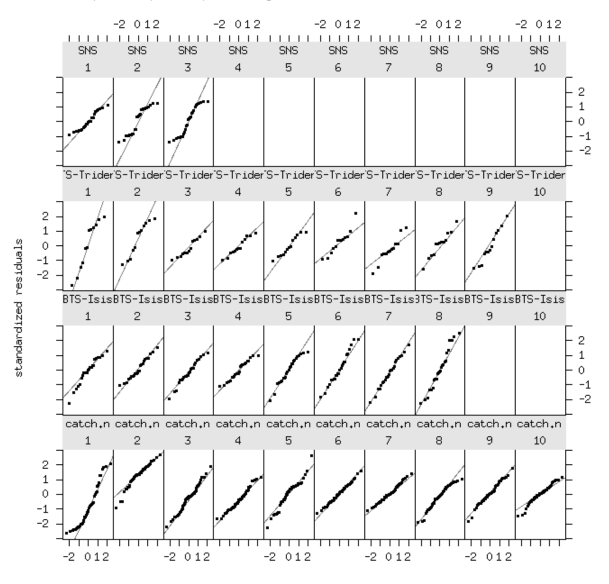


Figure 15: Quantile-quantile plot of standardized residuals.

To inspect the summaries (Fbar, SSB, catch and recruitment) of the fit the user may add the fit to the original stock object, using the method + and plot the result (Figure 16).

```
stk <- ple4 + fit
plot(stk)</pre>
```



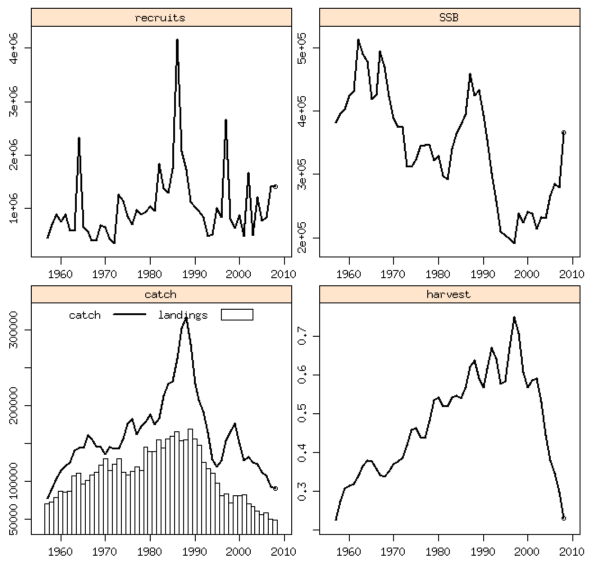


Figure 16: Stock summary

In more detail, one can plot a 3D representation of fishing mortality (Figure 17),

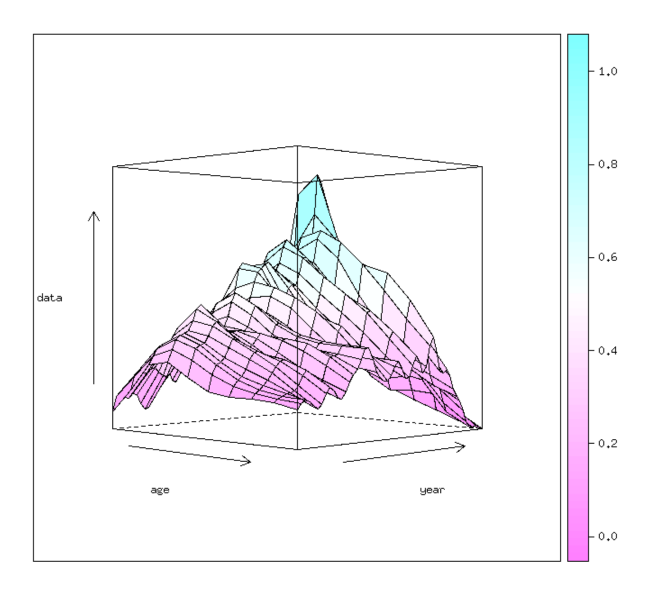


Figure 17: Fishing mortality

population abundance (Figure 18),

```
wireframe(data ~ age + year, data = as.data.frame(stock.n(stk)), drape = TRUE,
    screen = list(x = -90, y = -45))
```

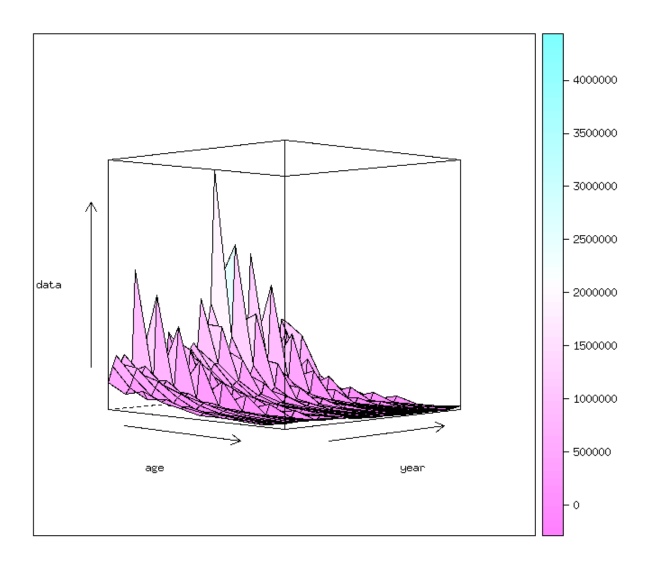


Figure 18: Population abundance

or catch-at-age (Figure 19).

```
wireframe(data ~ age + year, data = as.data.frame(catch.n(stk)), drape = TRUE)
```

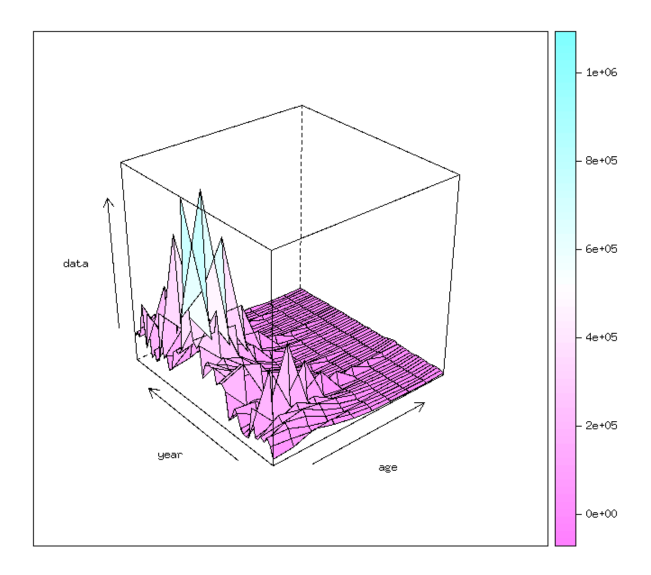


Figure 19: Catches

6.3 Data structures

As mentioned above, the output of the stock assessment method may be simpler, without all the information about the parameters of the model like the variance-covariance matrix, or may be more complex carrying all the information about the parameters. In the first case the class of the output object is a4aFit while in the second case is a4aFitSA.

This section will describe the data structures of these classes and the classes that compose them.

Starting with the basic model output class, a4aFit, the slots of this class are shown on the code below and Figure 20.

```
showClass("a4aFit")

## Class "a4aFit" [package "FLa4a"]
##
```

```
## Slots:
##
## Name:
               call
                        clock
                                fitSumm
                                           stock.n
                                                     harvest
                                                                catch.n
## Class:
               call
                                           FLQuant
                                                     FLQuant
                                                                FLQuant
                      numeric
                                   array
##
## Name:
              index
                         name
                                    desc
                                             range
## Class: FLQuants character character
                                           numeric
##
## Extends: "FLComp"
##
## Known Subclasses: "a4aFitSA"
```



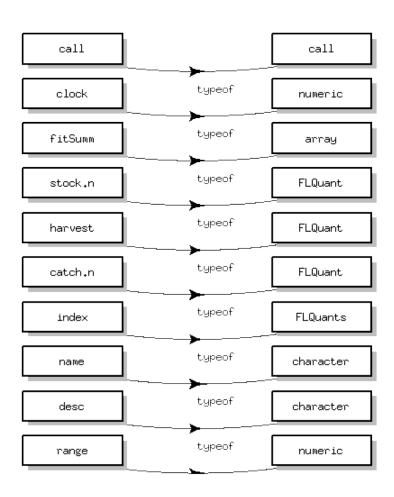


Figure 20: The a4aFit class

Fitted values are stored in the stock.n, harvest, catch.n and index slots. It also contains information carried over from the stock object used to fit the model, like the the name of the stock in name, any description provided in desc and the age and year range and mean F range in range. There is also a wall clock that has a breakdown of the time taken o run the model in clock.

The full assessment fit returns an object of class a4aFitSA, the slots of this class are shown on the code below and Figure 21.

```
showClass("a4aFitSA")
## Class "a4aFitSA" [package "FLa4a"]
##
## Slots:
##
## Name:
                                clock fitSumm
            pars
                       call
                                                 stock.n
                                                          harvest
## Class: SCAPars
                       call
                                                 FLQuant
                                                          FLQuant
                              numeric
                                        array
##
## Name: catch.n
                      index
                                 name
                                          desc
                                                   range
## Class: FLQuant FLQuants character character
                                                 numeric
##
## Extends:
## Class "a4aFit", directly
## Class "FLComp", by class "a4aFit", distance 2 \,
```

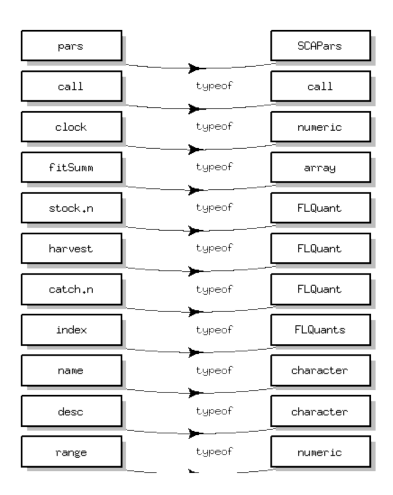


Figure 21: The a4aFitSA class

The additional slots in the assessment output are the fitSumm and pars slots, which are containers for model summaries and the model parameters. The pars slot is a class of type SCAPars (Figure 22) which is itself composed of sub-classes, designed to contain the information necessary to simulate from the model.

```
showClass("SCAPars")

## Class "SCAPars" [package "FLa4a"]

##

## Slots:

##

## Name: stkmodel qmodel vmodel

## Class: a4aStkParams submodels submodels

showClass("a4aStkParams")

## Class "a4aStkParams" [package "FLa4a"]
```

```
##
## Slots:
##
                                 srMod params vcov centering ormula FLPar array numeric
## Name: fMod n1Mod
## Class: formula formula
                              formula
##
                      m
## Name:
          distr
                                                      desc
                                units
                                          name
                                                               range
## Class: character FLQuant character character character
                                                             numeric
## Extends: "FLComp"
showClass("submodel")
## Class "submodel" [package "FLa4a"]
##
## Slots:
##
                     params vcov centering distr name FLPar array numeric character character
## Name: Mod
## Class: formula
##
## Name:
              desc
                      range
## Class: character
                     numeric
##
## Extends: "FLComp"
```

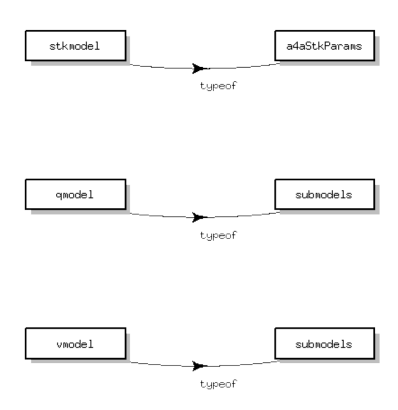


Figure 22: The SCAPars class

The SCAPars is built using objects of class a4aStkParams (Figure 23) and submodel (Figure 24). These classes have the following slots.

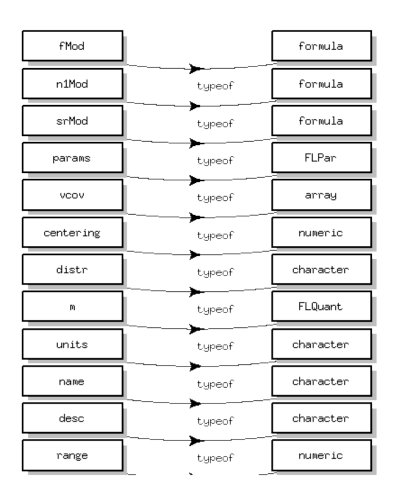


Figure 23: The a4aStkParams class

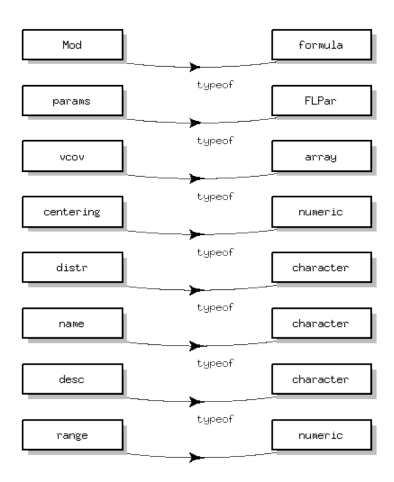


Figure 24: The submodel class

For example, all the parameters required so simulate a time-series of mean F trends is contained in the stkmodel slot, which is a class of type a4aStkParams. This class contains the relevant submodels (see later), their parameters params and the joint covariance matrix vcov for all stock related parameters.

6.4 The sca method - statistical catch-at-age

The sca() method used in the previous section with the default settings, can be parametrized to control other features of the stock assessment framework. The most interesting ones are the submodels for F, Q and R.

An important argument for sca() is the type of fit, which controls if a full assessment will be performed or a management procedure type of assessment. The argument is called fit and can have the values 'assessment' or 'MP', regarding a full assessment or a simpler assessment, repectively. By default sca() uses fit='MP'.

We'll start by looking at the submodel for F, then Q and finally R.

Please note that each of these model forms have not been tuned to the data. The degrees of freedom of each model can be better tuned to the data by using model selection procedures such as AIC or BIC, etc.

6.4.1 Fishing mortality submodel

We will now take a look at some examples for F models and the forms that we can get. We'll fix the Q and R submodels.

Lets start with a separable model in which age and year effects are modelled as dummy variables, using the factor coding provided by R (Figure 25).

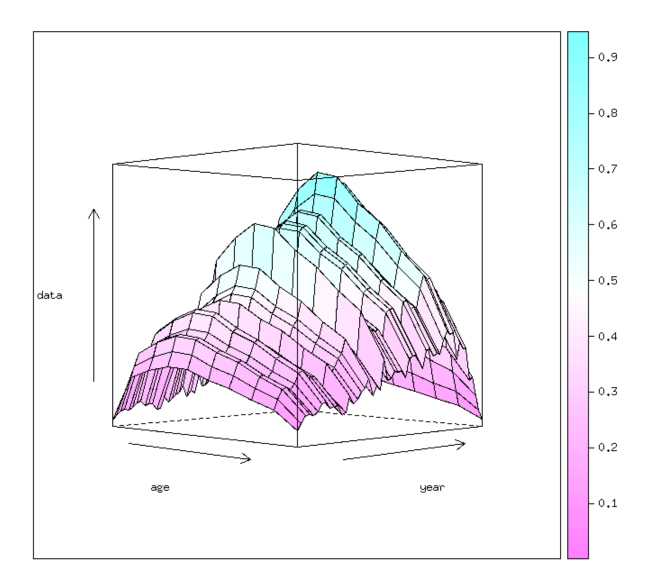


Figure 25: Fishing mortality separable model

Next we may make things a bit more interesting by using an (unpenalised) thin plate spline, where we'll borrow the method s() provided by package mgcv. We're using the North Sea Plaice data again, and since it has 10 ages we will use a simple rule of thumb that the spline should have fewer than $\frac{10}{2} = 5$

degrees of freedom, and so we opt for 4 degrees of freedom. We will also do the same for year and model the change in F through time as a smoother with 20 degrees of freedom. Note that this is still a separable model, it's a smoothed version of the previous model (Figure 26).

```
fmod <- ~s(age, k = 4) + s(year, k = 20)
fit1 <- sca(ple4, ple4.indices[1], fmod, qmod, srmod)</pre>
```

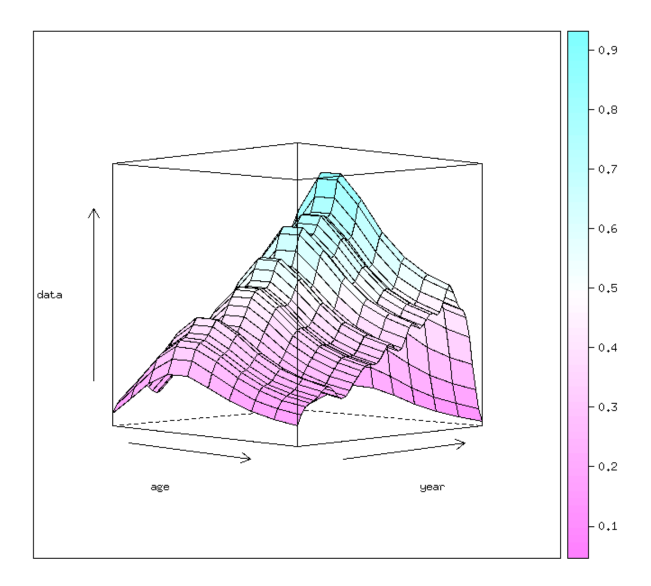
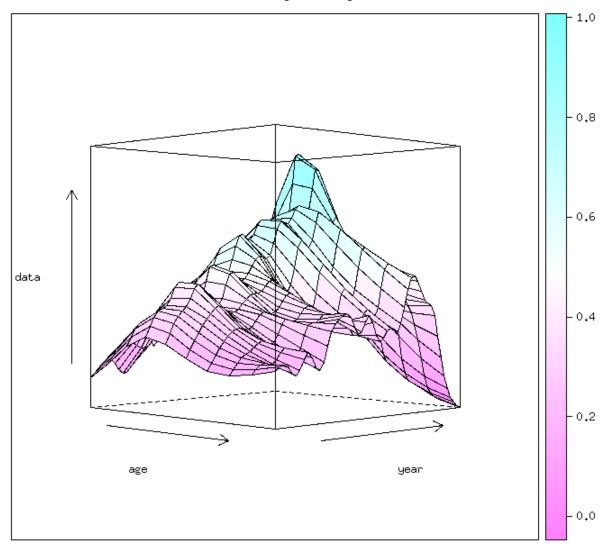


Figure 26: Fishing mortality smoothed separable model

A non-separable model, where we consider age and year to interact can be modeled using a smooth interaction term in the fmodel using a tensor product of cubic splines with the te method, again borrowed from mgcv.

```
fmod <- ~te(age, year, k = c(4, 20))
fit3 <- sca(ple4, ple4.indices[1], fmod, qmod, srmod)</pre>
```

Fishing mortality



In the last examples the Fs are linked across age and time. What if we want to free up a specific age class because in the residuals we see a consistent pattern. This can happen, for example, if the spatial distribution of juveniles is disconnected to the distribution of adults. The fishery focuses on the adult fish, and therefore the the F on young fish is a function of the distribution of the juveniles and could deserve a specific model. This can be achieved by adding a component for the year effect on age 1 (Figure 27).

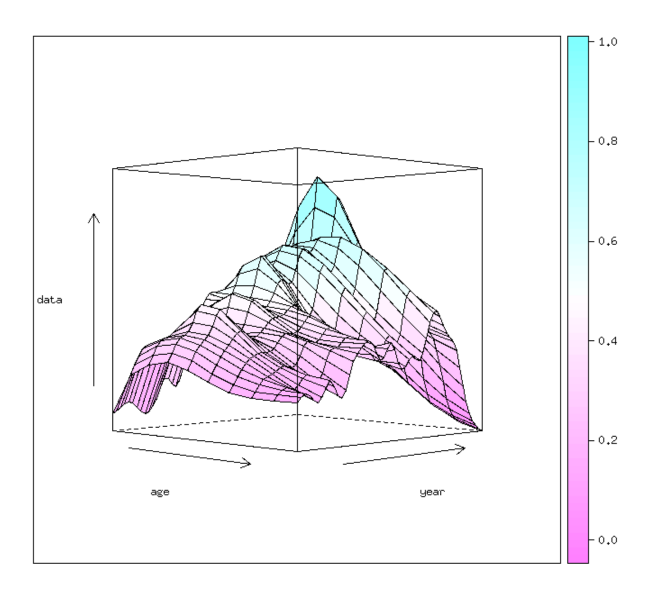


Figure 27: Fishing mortality age-year interaction model with extra age 1 smoother.

6.4.2 Catchability submodel

The catchability submodel is set up the same way as the F submodel and the tools available are the same. The only difference is that the submodel is set up as a list of formulas, where each formula relates with one abundance index.

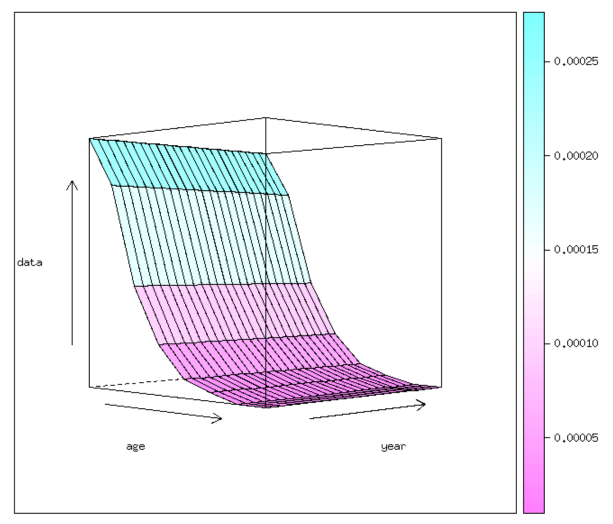
We'll start by fixing the F and R models and compute the fraction of the year the index relates too, which will allows us to compute catchability at age and year.

```
sfrac <- mean(range(ple4.indices[[1]])[c("startf", "endf")])
fmod <- ~factor(age) + factor(year)
srmod <- ~factor(year)</pre>
```

A first model is simply a dummy effect on age, which means that a coefficient will be estimated for each age. Note that this kind of model considers that levels of the factor are independent.

```
qmod <- list(~factor(age))
fit <- sca(ple4, ple4.indices[1], fmod, qmod, srmod)

# compute N for the fraction of the year the survey is carried out
Z <- (m(ple4) + harvest(fit)) * sfrac
lst <- dimnames(fit@index[[1]])
lst$x <- stock.n(fit) * exp(-Z)
stkn <- do.call("trim", lst)</pre>
```

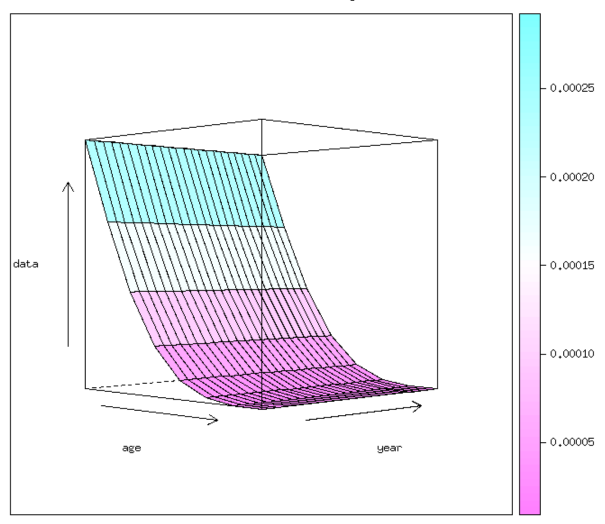


If one considers catchability at a specific age to be dependent on catchability on the other ages, similar to a selectivity modelling approach, one option is to use a smoother at age, and let the data 'speak' regarding the shape.

```
qmod <- list(~s(age, k = 4))
fit1 <- sca(ple4, ple4.indices[1], fmod, qmod, srmod)

# compute N for the fraction of the year the survey is carried out
Z <- (m(ple4) + harvest(fit1)) * sfrac
lst <- dimnames(fit1@index[[1]])</pre>
```

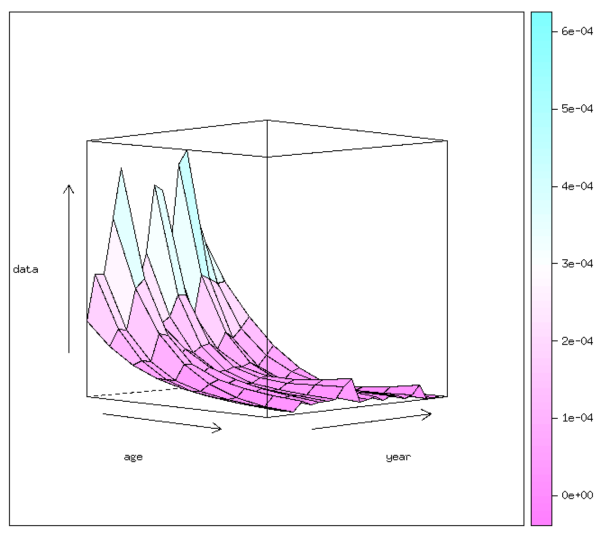
```
lst$x <- stock.n(fit1) * exp(-Z)
stkn <- do.call("trim", lst)</pre>
```



As in the case of F, one may consider catchability to be a process that evolves with age and year, including an interaction between the two effects. Such model can be modelled using the tensor product of cubic splines, the same way we did for the F model.

```
qmod <- list(~te(age, year, k = c(3, 40)))
fit2 <- sca(ple4, ple4.indices[1], fmod, qmod, srmod)

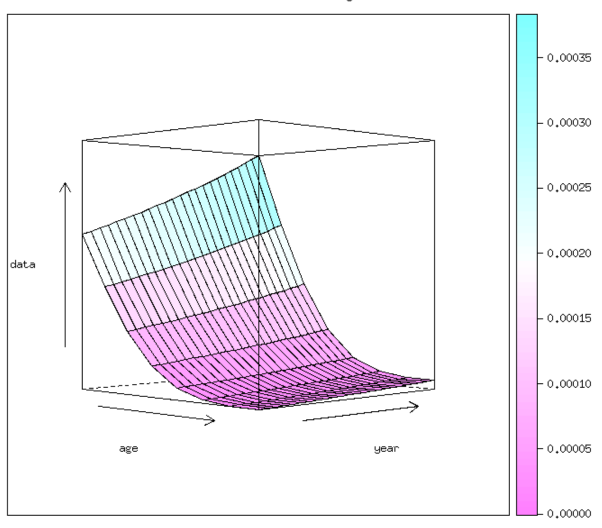
# compute N for the fraction of the year the survey is carried out
Z <- (m(ple4) + harvest(fit2)) * sfrac
lst <- dimnames(fit2@index[[1]])
lst$x <- stock.n(fit2) * exp(-Z)
stkn <- do.call("trim", lst)</pre>
```



Finally, one may want to investigate a trend in catchability with time, very common in indices built from CPUE data. In the example given here we'll use a linear trend in time, set up by a simple linear model.

```
qmod <- list(~s(age, k = 4) + year)
fit3 <- sca(ple4, ple4.indices[1], fmod, qmod, srmod)

# compute N for the fraction of the year the survey is carried out
Z <- (m(ple4) + harvest(fit3)) * sfrac
lst <- dimnames(fit3@index[[1]])
lst$x <- stock.n(fit3) * exp(-Z)
stkn <- do.call("trim", lst)</pre>
```



6.4.3 Catchability submodel for age aggregated indices

The previous section was focused on age disaggregated indices, but age aggregated indices (biomass, DEPM, etc) may also be used to tune the total biomass of the population. In these cases the *FLIndex* has to be named "bio" in the FLIndices object to be passed to the fitting method. At the moment only one "bio" index is allowed. Note that in this case the qmodel should be set without age factors. It could have a "year" component though.

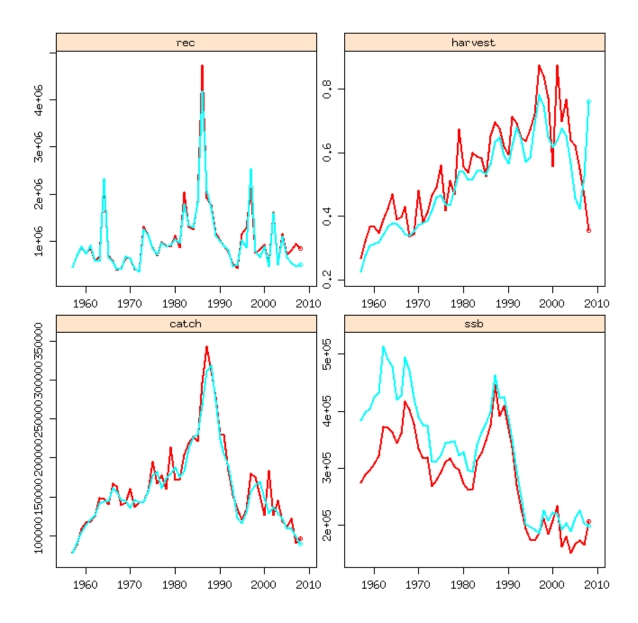
```
# creating an index
bioidx <- FLIndex(FLQuant(NA, dimnames = list(age = "all", year = range(ple4)["minyear"]:range(ple4)["mindex(bioidx) <- stock(ple4) * 0.001
index(bioidx) <- index(bioidx) * exp(rnorm(index(bioidx), sd = 0.1))
range(bioidx)[c("startf", "endf")] <- c(0, 0)

# fitting the model
fit <- sca(ple4, FLIndices(bio = bioidx), qmodel = list(~1))

# how is it fiting catchability ? not too well ...
index(fit)[[1]]/(quantSums(stock.n(fit) * stock.wt(ple4)))</pre>
```

```
## An object of class "FLQuant"
  , , unit = unique, season = all, area = unique
##
##
##
       year
## age
       1957
                  1958
                             1959
                                       1960
                                                  1961
                                                            1962
##
    all 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05
##
       year
## age
        1963
                  1964
                             1965
                                       1966
                                                  1967
    all 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05
##
       year
                  1970
                                       1972
## age
       1969
                            1971
                                                  1973
                                                            1974
    all 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05
##
       year
## age
       1975
                  1976
                             1977
                                       1978
                                                  1979
    all 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05
##
##
       year
## age
                                       1984
       1981
                  1982
                            1983
                                                  1985
    all 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05
##
##
       year
## age
       1987
                  1988
                             1989
                                       1990
                                                  1991
                                                            1992
##
    all 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05
##
       year
                  1994
                             1995
                                       1996
                                                  1997
## age
        1993
    all 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05
       year
##
## age
                  2000
                             2001
                                       2002
       1999
                                                  2003
                                                            2004
    all 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05 4.7796e-05
       year
## age
        2005
                  2006
                             2007
                                       2008
##
    all 4.7796e-05 4.7796e-05 4.7796e-05
##
## units: kg
```

Residuals are not so good ... one shouldn't expect much from a model with only one biomass index anyway. Note that the results are not so bad because most information is coming from the catch at age information.

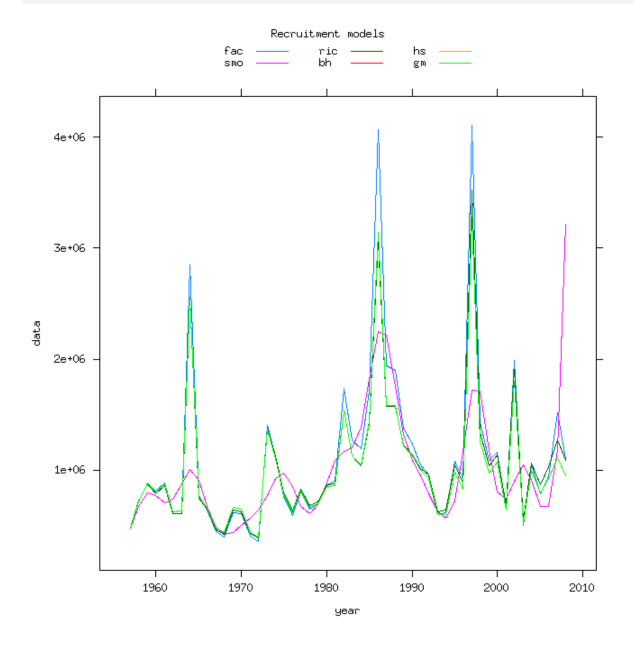


6.4.4 Stock-recruitment submodel

The S/R submodel is a special case, in he sense that it can be set up with the same linear tools as the F and Q models, but it can also use some hard coded models. The example shows how to set up a simple dummy model with factor(), a smooth model with s(), a Ricker model (ricker()), a Beverton and Holt model (bevholt()), a hockey stick model (hockey()), and a geometric mean model (geomean). As mentioned before, the 'structural' models have a fixed variance, which must be set by defining the coefficient of variation. We now fix the F and Q submodels before fiddling around with the S/R model.

```
fmod <- ~s(age, k = 4) + s(year, k = 20)
qmod <- list(~s(age, k = 4))</pre>
```

```
srmod <- ~factor(year)
fit <- sca(ple4, ple4.indices[1], fmod, qmod, srmod)
srmod <- ~s(year, k = 20)
fit1 <- sca(ple4, ple4.indices[1], fmod, qmod, srmod)
srmod <- ~ricker(CV = 0.05)
fit2 <- sca(ple4, ple4.indices[1], fmod, qmod, srmod)</pre>
```



6.5 The a4aSCA method - advanced features

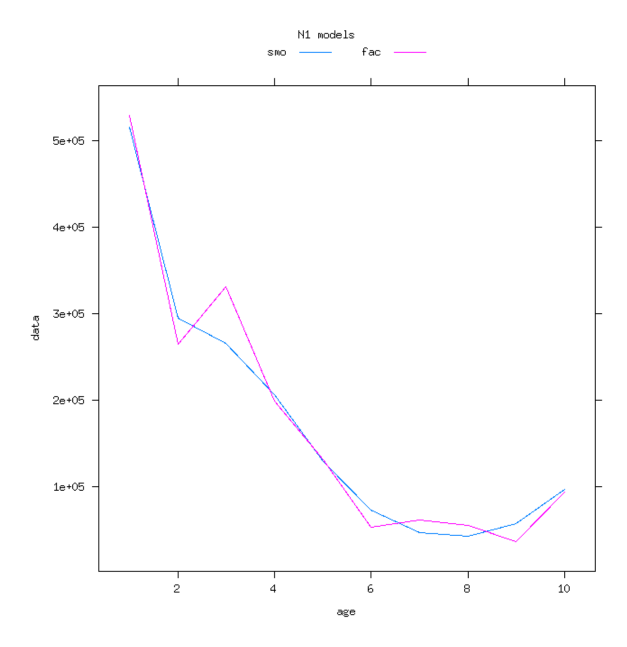
A more advanced method for stock assessment can be used through the a4aSCA() method. This method gives access to the submodels for N1, σ_{ay}^2 and I_{ays} as well as arguments to get the ADMB files, and etc. Check the manual pages with ?a4aSCA for more information. This method has 'assessment' as default fit, which means that the hessian is going to be computed and all the information about the parameters will be returned by default. Note that the default models of each submodel can be accessed with

```
\# fmodel <- \tilde{} s(age, k=4) + s(year, k = 20) qmodel <- list( \tilde{} s(age, k=4) +
# year) srmodel <- ~s(year, k=20)</pre>
fit <- a4aSCA(ple4, ple4.indices[1])</pre>
submodels(fit)
     fmodel: ~s(age, k = 3) + factor(year)
## srmodel: ~factor(year)
## n1model: ~factor(age)
##
     qmodel:
##
       BTS-Isis: ~1
##
     vmodel:
##
       catch:
                  s(age, k = 3)
       BTS-Isis: ~1
```

6.5.1 N1 model

The submodel for the stock number at age in the first year of the time series is set up with the usual linear tools, but bare in mind that the year effect does not make sense here.

```
n1mod <- ~s(age, k = 4)
fit1 <- a4aSCA(ple4, ple4.indices[1], n1model = n1mod)
flqs <- FLQuants(smo = stock.n(fit1)[, 1], fac = stock.n(fit)[, 1])</pre>
```

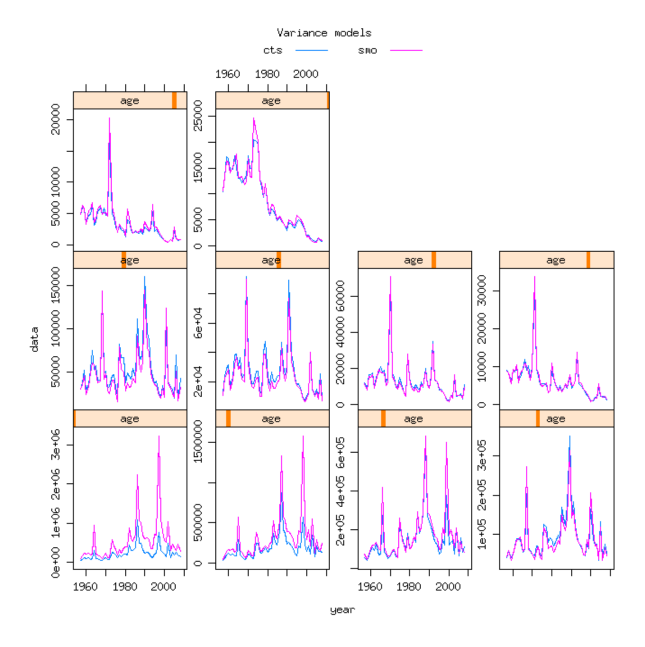


6.5.2 Variance model

The variance model allows the user to set up the shape of the observation variances σ_{ay}^2 and I_{ays} . This is an important subject related with fisheries data used for input to stock assessment models. It's quite common to have more precision on the most represented ages and less precision on the less frequent ages. Due to the fact that the last do not show so often on the auction markets, on the fishing operations or on survey samples.

By default the model assumes constant variance over time and ages (1 model) but it can use other models specified by the user. As with the other submodels, R linear model capabilities are used.

```
vmod <- list(~1, ~1)
fit1 <- a4aSCA(ple4, ple4.indices[1], vmodel = vmod)
vmod <- list(~s(age, k = 4), ~1)
fit2 <- a4aSCA(ple4, ple4.indices[1], vmodel = vmod)
flqs <- FLQuants(cts = catch.n(fit1), smo = catch.n(fit2))</pre>
```



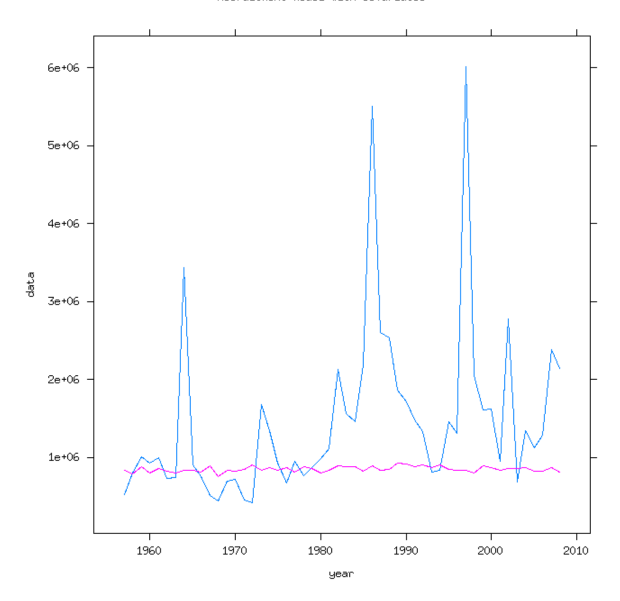
6.5.3 Working with covariates

In linear model one can use covariates to explain part of the variance observed on the data that the 'core' model does not explain. The same can be done in the a4a framework. The example below uses the North Atlantic Oscillation (NAO) index to model recruitment.

First by simply assuming that the index drives recruitment.

```
srmod <- ~nao
fit2 <- a4aSCA(ple4, ple4.indices[1], qmodel = list(~s(age, k = 4)), srmodel = srmod)
flqs <- FLQuants(fac = stock.n(fit)[1], cvar = stock.n(fit2)[1])</pre>
```

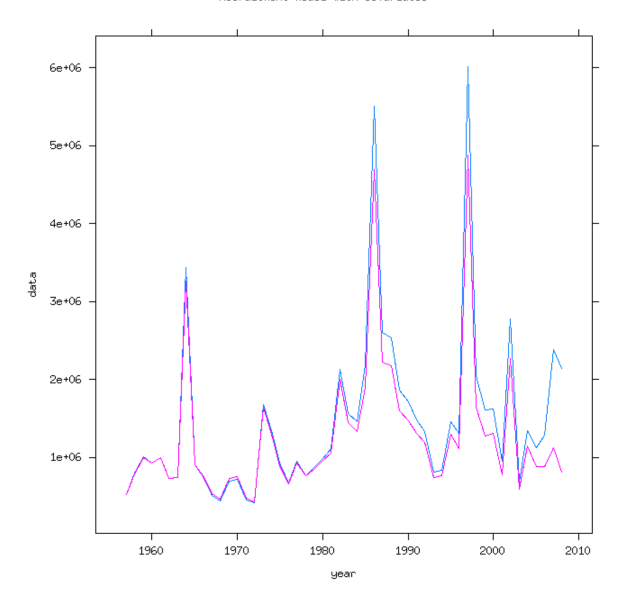
Recruitment model with covariates



In a second model we're using the NAO index not to model recruitment directly but to model one of the parameters of the $\rm S/R$ function.

```
srmod <- ~ricker(a = ~nao, CV = 0.1)
fit3 <- a4aSCA(ple4, ple4.indices[1], qmodel = list(~s(age, k = 4)), srmodel = srmod)
flqs <- FLQuants(fac = stock.n(fit)[1], cvar = stock.n(fit3)[1])</pre>
```





Note that covariates can be added to any submodel using the linear model capabilities of R.

6.5.4 Assessing ADMB files

The framework gives access to the files produced to run the ADMB fitting routine through the argument wkdir. When set up all the ADMB files will be left in the directory. Note that the ADMB tpl file is distributed with the FLa4a. One can get it from your R library, under the folder myRlib/FLa4a/admb/.

```
fit1 <- a4aSCA(ple4, ple4.indices, wkdir = "mytest")
## Model and results are stored in working directory [mytest]</pre>
```

6.6 Predict and simulate

Predicting and simulating are important methods for statistical models. R uses the methods predict() and simulate(), which were implemented in FLa4a in the same fashion.

```
# fmodel <- ~ s(age, k=4) + s(year, k = 20) qmodel <- list( ~ s(age, k=4) +
# year) srmodel <- ~ s(year, k=20)
fit <- a4aSCA(ple4, ple4.indices[1])</pre>
```

6.6.1 Predict

Predict simply computes the quantities of interest using the estimated coefficients.

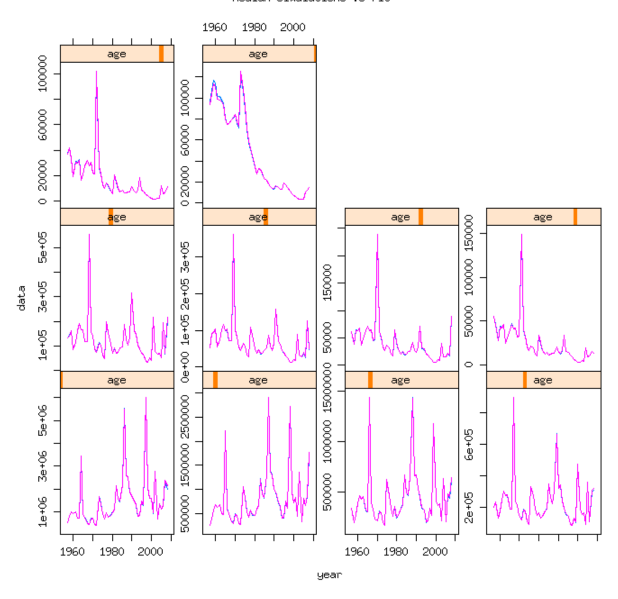
```
fit.pred <- predict(fit)
lapply(fit.pred, names)

## $stkmodel
## [1] "harvest" "rec" "ny1"
##
## $qmodel
## [1] "BTS-Isis"
##
## $vmodel
## [1] "catch" "BTS-Isis"</pre>
```

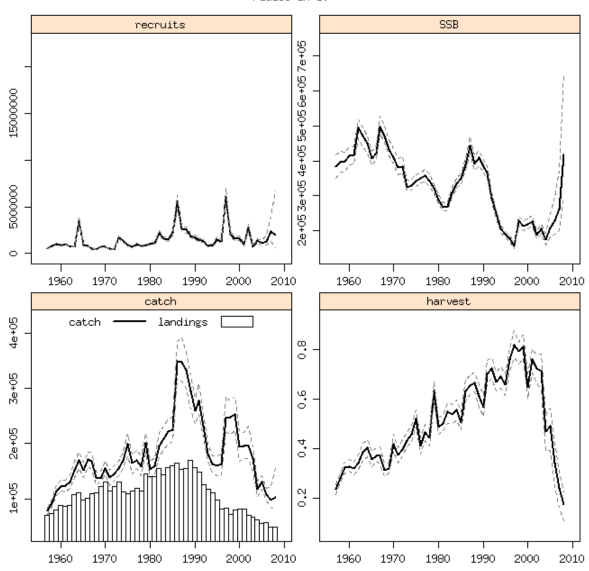
6.6.2 simulate

Simulate uses the variance-covariance matrix computed from the Hessian returned by ADMB and the fitted parameters, to parametrize a multivariate normal distribution. The simulations are carried out using the method mvrnorm() provided by the R package MASS.

```
fits <- simulate(fit, 100)
flqs <- FLQuants(sim = iterMedians(stock.n(fits)), det = stock.n(fit))</pre>
```







6.7 Geeky stuff

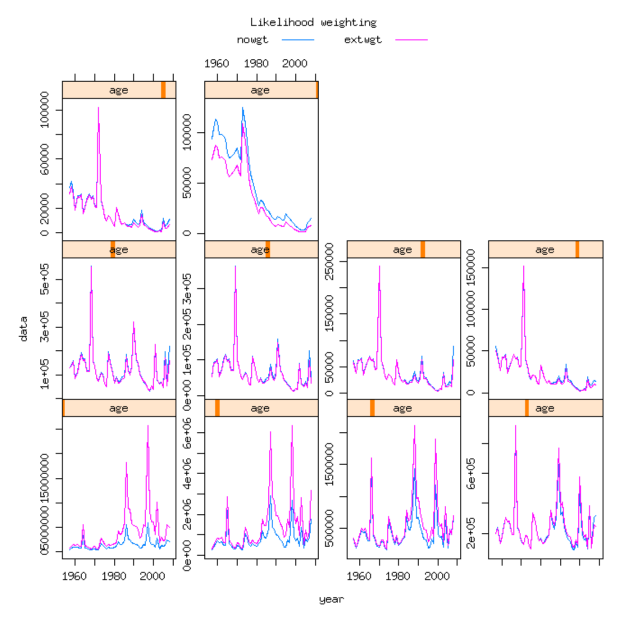
A lot more can be done with the a4a framework. The next sections will describe methods that are more technical. What we'd categorize as 'matters for geeks', in the sense that these methods usually will require the users to 'dive' into R a bit more.

```
# fmodel <- ~ s(age, k=4) + s(year, k = 20) qmodel <- list( ~ s(age, k=4) +
# year) srmodel <- ~ s(year, k=20) vmodel <- list(~1, ~1)
fit <- a4aSCA(ple4, ple4.indices[1])</pre>
```

6.7.1 External weighting of likelihood components

By default the likelihood components are weighted using inverse variance. However the user may change this weights by setting the variance of the input parameters, which is done by adding a variance matrix to the catch.n and index.n slots of the stock and index objects. These variances will be used to penalize the data during the likelihood computation. The values should be given as coefficient of variation in the log scale, so that variance is $\log (CV^2 + 1)$.

```
stk <- ple4
idx <- ple4.indices[1]
# variance of observed catches
varslt <- catch.n(stk)
varslt[] <- 0.4
catch.n(stk) <- FLQuantDistr(catch.n(stk), varslt)
# variance of observed indices
varslt <- index(idx[[1]])
varslt[] <- 0.1
index.var(idx[[1]]) <- varslt
# run
fit1 <- a4aSCA(stk, idx)
flqs <- FLQuants(nowgt = stock.n(fit), extwgt = stock.n(fit1))</pre>
```



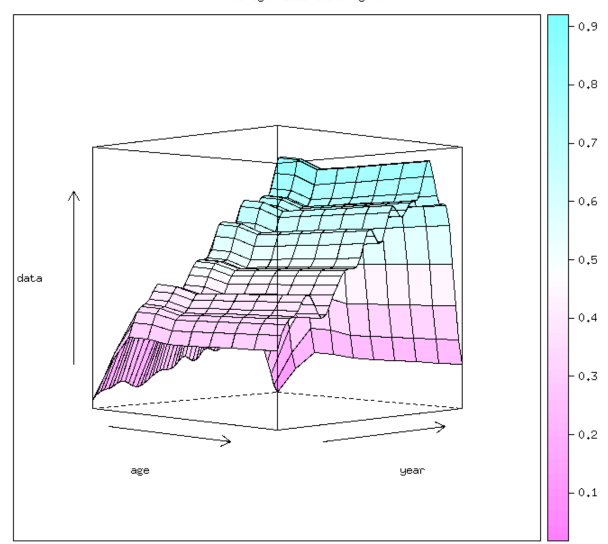
6.7.2 More models

There's a set of methods that allow the user to have more flexibility on applying the models referred before. For example to break the time series in two periods, using the method breakpts(), or fixing some parts of the selection pattern by setting F to be the same for a group of ages, using replace().

The example below replaces all ages above 5 by age 5, which means that a single coefficient is going to be estimated for age 5-10.

```
fmod <- \tilde{s}(replace(age, age > 5, 5), k = 4) + s(year, k = 20) fit <- sca(ple4, ple4.indices, fmod)
```

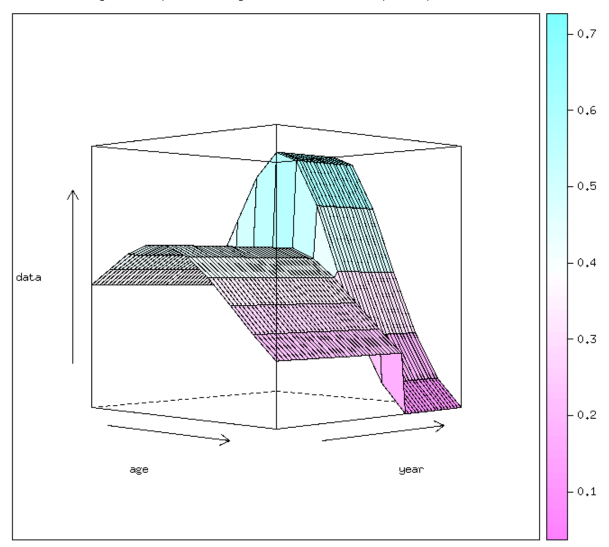




While in the next case we'll use the breakpts() to split the time series in 1990, although keeping the same shape in both periods, a thin plate spline with 3 knots.

```
fmod <- ~s(age, k = 3, by = breakpts(year, 1990))
fit <- sca(ple4, ple4.indices, fmod)</pre>
```

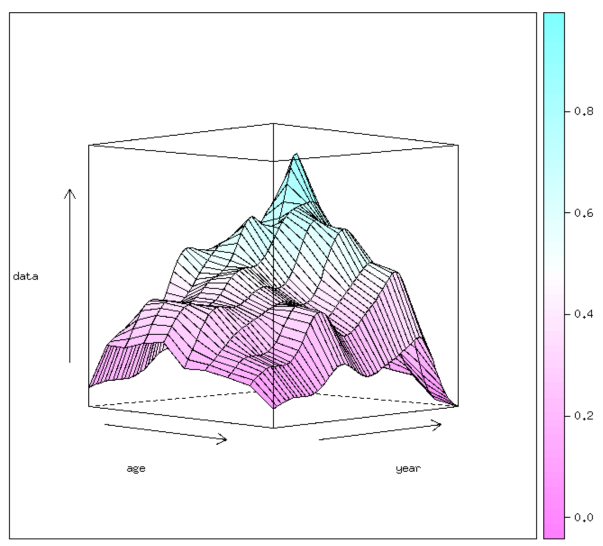
F-at-age in two periods using in both cases a thin plate spline with 3 knots



More complicated models can be built with these tools. For example a model where the age effect is modelled as a smoother (the same thin plate spline) throughout years but independent from each other.

```
fmod <- ~factor(age) + s(year, k = 10, by = breakpts(age, c(2:8)))
fit <- sca(ple4, ple4.indices, fmod)</pre>
```

F-at-age as thin plate spline with 3 knots for each age



6.7.3 Propagate M uncertainty

In this section we give the example of how M uncertainty set up using the m() method and the class $a \nmid a M$, is propagated through the stock assessment. We'll start by fitting the default model to the data.

```
data(ple4)
data(ple4.indices)
fit <- sca(ple4, ple4.indices)</pre>
```

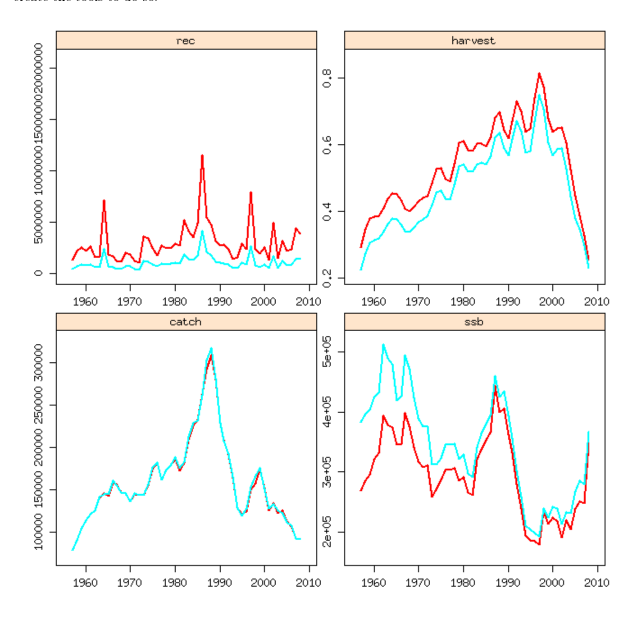
Now, using the a4a methods we'll model natural mortality using a negative exponential model by age, Jensen's estimator for the level and a constant trend with time with some noise.

```
m4 <- a4aM(shape = shape, level = level, trend = trend)
m4 <- mvrnorm(nits, m4)
range(m4)[] <- range(ple4)[]
range(m4)[c("minmbar", "maxmbar")] <- c(1, 1)
flq <- m(m4)[]
quant(flq) <- "age"
stk <- propagate(ple4, nits)
m(stk) <- flq</pre>
```

Fit the same model to the new stock information, which includes the uncertainty on M.

```
fit1 <- sca(stk, ple4.indices)</pre>
```

And compare the two results. It's quite easy to run these kind of tests and a large part of our effort is to create the tools to do so.

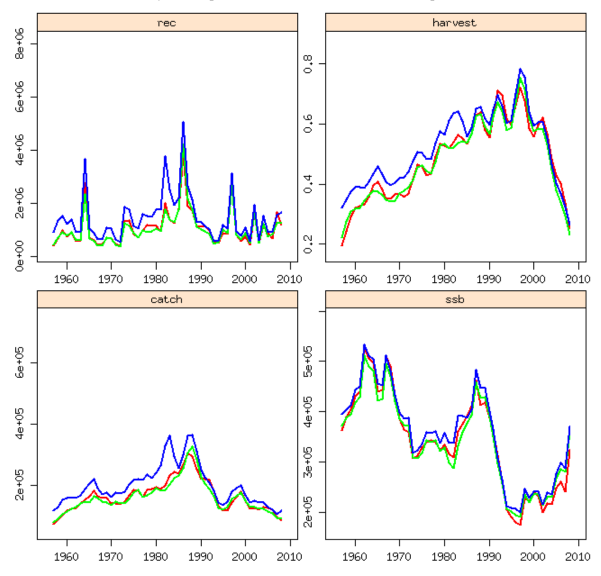


6.7.4 WCSAM exercise - replicating itself

The World Conference on Stock Assessment Methods (WCSAM) promoted a workshop where a large simulation study was used to test the performance of distinct stock assessment models. The first criteria used was that the models should be able to reproduce itself. The process involved fitting the model, simulate observation error using the same model, and refit the model to each iteration. The final results should be similar to the fitted results before adding observation error (see Deroba, et.al, 2014 for details). The following analysis runs this analysis.

```
data(ple4)
data(ple4.indices)
fit <- sca(ple4, ple4.indices, fit = "assessment")</pre>
nits <- 25
stk <- ple4 + fit
fits <- simulate(fit, nits)</pre>
stks <- ple4 + fits
idxs <- ple4.indices[1]</pre>
index(idxs[[1]]) <- index(fits)[[1]]</pre>
library(parallel)
options(mc.cores = 4)
lst <- lapply(split(1:nits, 1:nits), function(x) {</pre>
    out <- try(sca(iter(stks, x), FLIndices(iter(idxs[[1]], x))))</pre>
    if (is(out, "try-error"))
         NULL else out
})
stks2 <- stks
for (i in 1:nits) {
    iter(catch.n(stks2), i) <- catch.n(lst[[i]])</pre>
    iter(stock.n(stks2), i) <- stock.n(lst[[i]])</pre>
    iter(harvest(stks2), i) <- harvest(lst[[i]])</pre>
}
catch(stks2) <- computeCatch(stks2)</pre>
stock(stks2) <- computeStock(stks2)</pre>
```





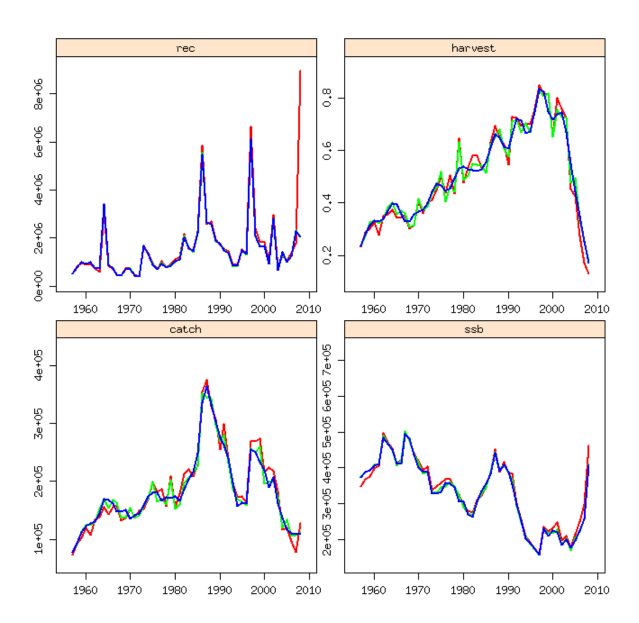
6.7.5 Paralell computing

This is an example of how to use the parallel package to run assessments. In this example each iteration is a dataset, including surveys, and we'll run one assessment for each iteration. Afterwards the data is pulled back together in a FLStock object and plotted. Only 20 iterations are ran to avoid taking too long. Also note that we're using 4 cores. This parameter depends on the computer being used. These days almost all computers have 2 cores.

Finally, compare this code with the one for replicating WCSAM and note that it's exactly the same, except that we're using mclapply() from package paralell instead of lapply().

```
data(ple4)
data(ple4.indices)
nits <- 25
fit <- a4aSCA(ple4, ple4.indices[1])
stk <- ple4 + fit
fits <- simulate(fit, nits)
stks <- ple4 + fits
idxs <- ple4.indices[1]</pre>
```

```
index(idxs[[1]]) <- index(fits)[[1]]</pre>
library(parallel)
options(mc.cores = 4)
lst <- mclapply(split(1:nits, 1:nits), function(x) {</pre>
    out <- try(sca(iter(stks, x), FLIndices(iter(idxs[[1]], x))))</pre>
    if (is(out, "try-error"))
         NULL else out
})
stks2 <- stks
for (i in 1:nits) {
    iter(catch.n(stks2), i) <- catch.n(lst[[i]])</pre>
    iter(stock.n(stks2), i) <- stock.n(lst[[i]])</pre>
    iter(harvest(stks2), i) <- harvest(lst[[i]])</pre>
}
catch(stks2) <- computeCatch(stks2)</pre>
stock(stks2) <- computeStock(stks2)</pre>
stks3 <- FLStocks(orig = stk, sim = stks, fitsim = stks2)</pre>
```

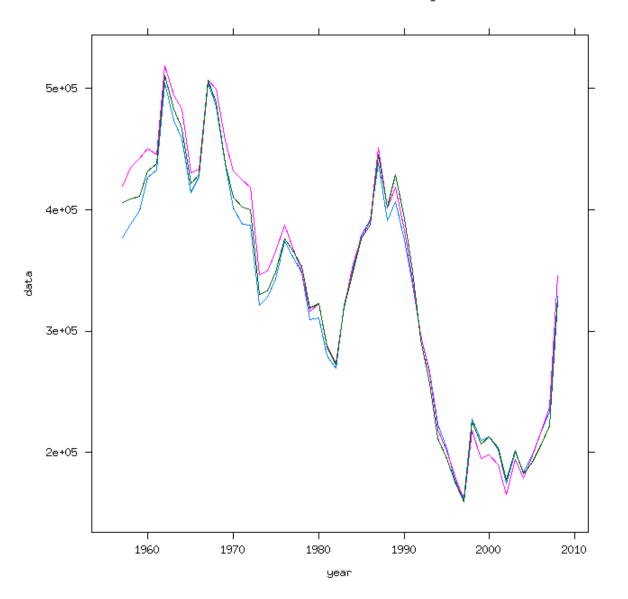


6.8 Model averaging

To merge results from several fits, using distinct models or datasets, we follow the paper Colin et al. The method ma() is the wrapper to the distinct methods, although for now only the AIC averaging is implemented.

```
data(ple4)
data(ple4.indices)
f1 <- sca(ple4, ple4.indices, fmodel = ~factor(age) + s(year, k = 20), qmodel = list(~s(age, k = 4), ~s(age, k = 3)), fit = "assessment")
f2 <- sca(ple4, ple4.indices, fmodel = ~factor(age) + s(year, k = 20), qmodel = list(~s(age, k = 4) + year, ~s(age, k = 4), ~s(age, k = 3)), fit = "assessment")
stock.sim <- ma(a4aFitSAs(list(f1 = f1, f2 = f2)), ple4, AIC, nsim = 100)
stks <- FLStocks(f1 = ple4 + f1, f2 = ple4 + f2, ma = stock.sim)</pre>
```

SSB of the two models and their average



Stock summaries of the two models and their average

