Assessment for All initiative(a4a) Natural Mortality Modelling

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Abstract

The document explains the approach being developed by a4a to integrate uncertainty in natural mortality into stock assessment and advice. It presents a mixture of text and code, where the first explains the concepts behind the methods, while the last shows how these can be run with the software provided.

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1 Background

In the a4a natural mortality is dealt with as an external parameter to the stock assessment model. The rationale to modelling natural mortality 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 natural mortality model and parameters.
- 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.

2 License, documentation and development status

The software is released under the EUPL 1.1.

For more information on the a4a methodologies refer to Jardim, et.al, 2014, Millar, et.al, 2014 and Scott, et.al, 2016.

Documentation can be found at http://flr-project.org/FLa4a. You are welcome to:

- Submit suggestions and bug-reports at: https://github.com/flr/FLa4a/issues
- Send a pull request on: https://github.com/flr/FLa4a/
- Compose a friendly e-mail to the maintainer, see 'packageDescription('FLa4a')'

3 Installing and loading libraries

To run the FLa4a methods the reader will need to install the package and its dependencies and load them. Some datasets are distributed with the package and as such need to be loaded too.

```
# from CRAN
install.packages(c("copula", "triangle", "coda"))
# from FLR
install.packages(c("FLCore", "FLa4a"), repos = "http://flr-project.org/R")
```

```
# libraries
library(FLa4a)
library(XML)
library(reshape2)
library(latticeExtra)
# datasets
data(ple4)
data(ple4.indices)
data(ple4.index)
data(rfLen)
packageVersion("FLCore")
## [1] '2.6.0.20161214'
```

```
packageVersion("FLa4a")
```

```
## [1] '1.0.0'
```

4 How to read this document

The target audience for this document are readers with some experience in R and some background on stock assessment.

The document explains the approach being developed by **a4a** for fish stock assessment and scientific advice. It presents a mixture of text and code, where the first explains the concepts behind the methods, while the last shows how these can be run with the software provided. Moreover, having the code allows the reader to copy/paste and replicate the analysis presented here.

The sections and subsections are as independent as possible, so it can be used as a reference document for the FLa4a.

Finally, this is a live document which will be updated and released often.

5 a4aM - The M class

Natural mortality is implemented in a class named a_4aM . This class is made up of three objects of the class *FLModelSim*. Each object is a model that 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* x *level* x *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
                                                                       numeric
                                                         character
##
## Extends: "FLComp"
```

The a_4aM constructor requires that the models and parameters are provided. The default method will build each of these models as a constant value of 1.

As a simple 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 (meaning that 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 the 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): $M_l = K(\frac{L_{inf}}{l})^{1.5}$. We use the default level and *trend* models.

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

Another option is to model how an external factor may impact the natural mortality. This can be added through the **trend** model. Suppose natural mortality can be modelled with a dependency on the NAO index, due to some mechanism that results in having lower mortality when NAO is negative and higher when it's positive. In this example, 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 mortality increases by 50% if NAO is positive on the first quarter.

```
# Get NAO
nao.orig <- read.table("https://www.esrl.noaa.gov/psd/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 <- (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)
mЗ
## a4aM object:
##
    shape: ~exp(-age - 0.5)
##
    level: ~1.5 * k
##
   trend: ~1 + b * nao
```

6 Adding uncertainty to natural mortality parameters with a multivariate normal distribution

Uncertainty on natural mortality is added through uncertainty on the parameters.

In this section we'll' show how to add multivariate normal uncertainty. We make use of the class *FLModelSim* method mvrnorm(), which is a wrapper for the method mvrnorm() distributed by the package MASS.

We'll create an a4aM object with an exponential shape, a level model based on k and temperature (Jensen's third estimator), and a trend model driven by the NAO (as above). Afterwards a variancecovariance matrix for the level and trend models will be included. Finally, create an object with 100 iterations using the mvrnorm() method.

Create the object:

```
shape4 <- FLModelSim(model = ~exp(-age - 0.5))
level4 <- FLModelSim(model = ~k^0.66 * t^0.57, 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)
# Call mvrnorm()
m4 <- mvrnorm(100, m4)
m4
## a4aM object:
## shape: ~exp(-age - 0.5)
## level: ~k^0.66 * t^0.57
## trend: ~1 + b * nao</pre>
```

Inspect 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
##
                  k
                                    t
## 0.39708(0.0439) 10.14011(0.9030)
## 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:

params(trend(m4))

```
## An object of class "FLPar"
## iters: 100
##
## params
## b
## 0.48259(0.159)
## units: NA
```

Note the shape model has no parameters and no uncertainty:

```
params(shape(m4))
## An object of class "FLPar"
## [1] NA
## units: NA
```

In this particular case, the **shape** model will not be randomized because it doesn't have a variancecovariance 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 by using mnrnorm() on each model component:

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

7 Adding uncertainty to natural mortality parameters with statistical 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 ??. 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, $M_l = K(\frac{L_{inf}}{l})^{1.5}$ and a triangle copula to model parameter uncertainty. The method mvrtriangle() is used to create 1000 iterations.

```
linf <- 60
k <- 0.4
# vcov matrix (make up some values)
mm <- matrix(NA, ncol = 2, nrow = 2)</pre>
```

```
# 10% cv
diag(mm) <- c((linf * 0.1)^2, (k * 0.1)^2)</pre>
# 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,] 1.0000000 0.2083333
## [2,] 0.2083333 1.0000000
# 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))</pre>
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
                                    k
## 59.99983(2.1608) 0.40362(0.0699)
## 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 1 and 2.

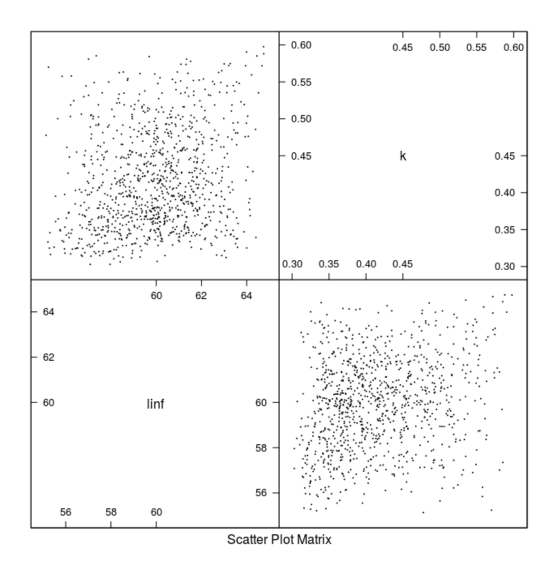


Figure 1: Parameter estimates for Gislason's second natural mortality model from using a triangle distribution.

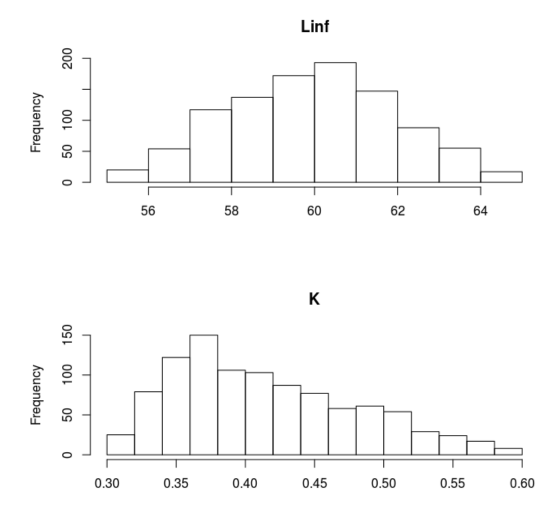


Figure 2: Marginal distributions of the parameters for Gislason's second natural mortality model using a triangle distribution.

We now have a new model that can be used for the **shape** model. You can 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 model based on 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>
```

8 Computing natural mortality time series - the "m" method

Now that we have set up the natural mortality a4aM model and added parameter uncertainty to each component, 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.

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'.

Here we demonstrate m() using the simple a4aM object we created above that has constant natural mortality.

Start with the simplest model:

m1
a4aM object:
shape: ~1
level: ~a
trend: ~1

Check the range:

range(m1)

##	min	max plus	group	minyear	maxyear	minmbar	maxmbar
##	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 and year ranges:

```
rngquant(m1) <- c(0, 7) # set the quant range</pre>
rngyear(m1) <- c(2000, 2010) # set the year range</pre>
range(m1)
##
          min
                     max plusgroup
                                       minyear
                                                  maxyear
                                                             minmbar
                                                                        maxmbar
                                          2000
                                                     2010
##
                       7
            0
                                  0
                                                                    0
                                                                               0
```

Create the object with the M estimates by age and year, note the name of the first dimension is 'quant'.

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 \ 0.2 \ 0.2 \ 0.2 \ 0.2 \ 0.2 \ 0.2 \ 0.2 \ 0.2 \ 0.2 \ 0.2 \ 0.2 \ 0.2
    1 \ 0.2 \ 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 0.2
    ##
##
    ##
    ##
    ##
##
## units: NA
```

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 of the resulting *FLQuant*. Note that in this case the **mbar** values in the range become relevant, once that **mbar** 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. We illustrate this by making an FLQuant with age varying natural mortality.

Check the model and set the ranges:

```
m2
## a4aM object:
## shape: ~exp(-age - 0.5)
  level: ~1.5 * k
##
##
  trend: ~1
rngquant(m2) <- c(0, 7)
rngyear(m2) <- c(2000, 2003)</pre>
range(m2)
##
      min max plusgroup minyear maxyear minmbar maxmbar
##
                 7 0 2000 2003 0
        0
                                                                  0
m(m2)
## An object of class "FLQuant"
## , , unit = unique, season = all, area = unique
##
##
    year
## age 2000
                2001
                          2002
                                    2003
##
    0 0.6000000 0.6000000 0.6000000 0.6000000
##
    1 0.22072766 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 ## 10.6

Which 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" (see above) and the mean natural mortality value over this range is given by the level model.

We can change the **mbar** range:

<pre>rngmbar(m2) <- c(0, 5) range(m2)</pre>									
##	min	max	plusgroup	minyear	maxyear	minmbar	maxmbar		
##	0	7	0	2000	2003	0	5		

Which rescales the the natural mortality at age:

```
m(m2)
## An object of class "FLQuant"
## , , unit = unique, season = all, area = unique
##
##
     year
## age 2000
                 2001
                           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 to the m() method. The year range of the 'nao' covariate should match that of the **range** slot.

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 ages:
rngquant(m3) <- c(0, 7)
m(m3, nao = 0)
## 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 - passing in the NAO data as numeric (1,0,1,0)

```
rngyear(m3) <- c(2000, 2003)</pre>
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 0.9000000 0.6000000 0.9000000 0.6000000
##
  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 3). We use the m4 object from earlier with uncertainty on the **level** and **trend** parameters.

```
rngquant(m4) <- c(0, 7)
rngyear(m4) <- c(2000, 2003)</pre>
flq <- m(m4, nao = as.numeric(nao[, as.character(2000:2003)]))</pre>
flq
## An object of class "FLQuant"
## iters: 100
##
##
  , , unit = unique, season = all, area = unique
##
##
     year
## age 2000
                           2001
                                               2002
##
    0 2.9751858(0.357394) 1.9987979(0.194298) 2.9751858(0.357394)
    1 1.0945097(0.131478) 0.7353167(0.071478) 1.0945097(0.131478)
##
    2 0.4026476(0.048368) 0.2705079(0.026295) 0.4026476(0.048368)
##
    3 0.1481258(0.017794) 0.0995143(0.009674) 0.1481258(0.017794)
##
##
    4 0.0544924(0.006546) 0.0366093(0.003559) 0.0544924(0.006546)
   5 0.0200466(0.002408) 0.0134678(0.001309) 0.0200466(0.002408)
##
##
    6 0.0073747(0.000886) 0.0049545(0.000482) 0.0073747(0.000886)
##
    7 0.0027130(0.000326) 0.0018227(0.000177) 0.0027130(0.000326)
##
    vear
## age 2003
## 0 1.9987979(0.194298)
##
  1 0.7353167(0.071478)
##
  2 0.2705079(0.026295)
##
   3 0.0995143(0.009674)
##
    4 0.0366093(0.003559)
##
    5 0.0134678(0.001309)
##
    6 0.0049545(0.000482)
##
    7 0.0018227(0.000177)
##
## units: NA
dim(flq)
## [1] 8 4 1 1 1 100
```

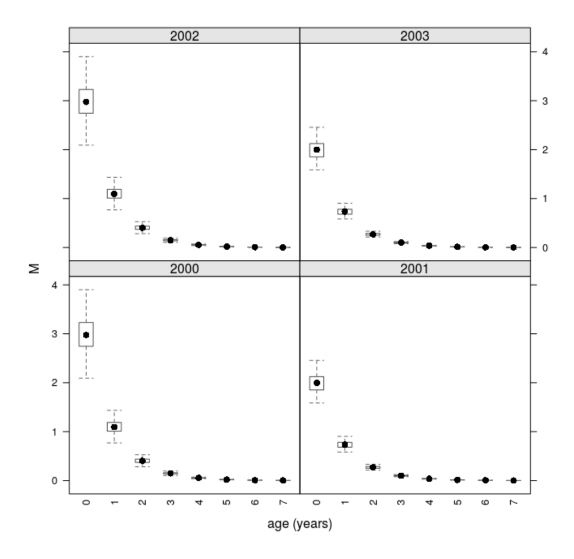


Figure 3: Natural mortality with age and year trend.