How to use the **ss3sim** package to run simulations in SS3

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First start by installing the latest version of ss3sim and loading the package:

```
> # install.packages("devtools")
> # devtools::install_github("ss3sim", username="seananderson")
> library(ss3sim)
```

You can read the help files and access this vignette again with:

```
> help(package = "ss3sim")
> vignette("ss3sim")
```

Setting up the file structure

This package is set up assuming that you have an established base case operating model and estimation model to work with. Each operating model and estimation model should be in their own folder. The operating should have the files:

- 1. yourmodel.ctl
- 2. yourmodel.dat
- 3. ss3.par
- 4. starter.ss

The estimation model folder should have:

- 1. yourmodel.ctl
- 2. yourmodel.dat

3. starter.ss

In both cases, nothing more and nothing less. The names of the .ctl and .dat files are not important. The package functions will rename them after they are copied to appropriate folders. These files should be derived from the .ss_new files but named as listed above. It's important to use these .ss_new files so they have consistent formatting. Many of the functions in this package depend on that formatting.

For the purposes of the Fish 600 project, we have unique case identifiers which combine to create unique scenarios. The types of cases are: natural mortality (M), fishing mortality (F), data quality (D), and retrospective (R). And the species are cod (cod), sardine-like (sar), and flatfish (fla). It's important to use these three letter abbreviations for the species since the functions assume that the last three letters represent a species (or some other identifier for a different project).

The different version of each case are identified with numbers. So, for example, the base case scenario for cod is identified as M1-F1-D1-R1-cod. We will have a spreadsheet that describes each of these in plain language.

The function copy_ss3models creates a folder structure and copies over the operating and estimation models. The folder structure looks like:

```
M1-F1-D1-R1-cod/1/om
M1-F1-D1-R1-cod/1/em
M1-F1-D1-R1-cod/2/om
M1-F1-D1-R1-cod/2/em
```

If you are using bias correction (bias_correct = TRUE) then there will be some additional folders. In that case the folders will look like:

```
M1-F1-D1-R1-cod/bias/1/om
M1-F1-D1-R1-cod/bias/1/em
M1-F1-D1-R1-cod/bias/2/om
M1-F1-D1-R1-cod/bias/2/em
...
M1-F1-D1-R1-cod/1/om
M1-F1-D1-R1-cod/1/em
M1-F1-D1-R1-cod/2/om
M1-F1-D1-R1-cod/2/om
...
```

Note that the operating and estimating model folders have been renamed om and em within each iteration, the operating and estimation models have been checked to make sure they contain the minimal files (as listed above), the filenames have been made all lowercase, the data file has been renamed data.dat, the control files have been renamed om.ctl or em.ctl, and the starter and control files have been adjusted to reflect these new file names.

The functions in this package assume you've set your working directory in R to be the base folder where you will store the scenario folders.

Creating the input configuration files

You will need to have a folder containing "case" argument definitions. These plain text files are read by get_caseval and turned into argument lists that are passed to run_ss3sim. You can create template input files by running create_argiles. It reads the various functions and parses the arguments and default values into plain text files. The default settings create these files:

```
1. MO-spp.txt
```

- 2. F0-spp.txt
- 3. index0-spp.txt
- 4. agecomp0-spp.txt
- 5. lcomp0-spp.txt
- 6. RO-spp.txt (not implemented yet)

Look in your working directory for the template files. Change the case ID number (defaults to 0) and the species identifier to a three letter identifier. For the Fish 600 project use one of cod, sar, or fla for cod, sardine, or flatfish. An example filename would be M1-sar.txt or lcomp2-fla.txt.

The first column in the text files denotes the argument to be passed to a function. The second argument denotes the value to be passed. You can use any simple R syntax. For example: c(1, 2, 4), or seq(1, 100) or 1:100 or matrix(). Character objects don't need to be quoted, but can be if you'd like. However, be careful not to use the delimiter (set up as a semicolon) anywhere else in the file besides to denote columns. You can add comments after any # symbols. Internally, the functions

evaluate in R any entries that have no character values (e.g. 1:100) or have an alphanumeric character followed by a (. Anything that is character only or has character mixed with numeric but doesn't have the regular expression "[A-Za-z0-9](" gets turned into a character argument.

Putting that all together, here's what an example F1-cod.txt file might look like:

```
years; 1:100
years_alter; NA
fvals; NA
```

Running the models

The run_ss3sim function is a wrapper function. It adjusts the natural mortality, adjusts the fishing mortality, adds recruitment deviations, calls run_ss3model to run the operating model, samples various survey estimates from the operating model, changes the age composition data as specified, changes the length composition data as specified, copies and renames files as necessary, and calls run_ss3model again to run the estimation model.

The run_fish600 function is a high-level wrapper that deals with parsing the case input files and then passes these arguments on to run_ss3sim.

Say you have your input case files setup and you want to run the first 50 iterations of those scenarios. You could run them like this:

```
> # First grab the example package data:
> d <- system.file("extdata", package = "ss3sim")
> f <- pasteO(d, "/run_ss3sim_eg/")
> om_model_dir <- pasteO(f, "cod_om")
> em_model_dir <- pasteO(f, "cod_em")
> case_folder <- pasteO(f, "case-arguments")
> #
> #
> # Without bias correction:
> run_fish600(iterations = 1, scenarios = c("M1-F1-D1-R1-cod"),
+ case_folder = case_folder, om_model_dir = om_model_dir,
+ em_model_dir = em_model_dir)
> #
> #
```

```
> # With bias correction:
> # (Note that bias_nsim should be bigger, say 5, but it is set to 1
> # here so the example runs faster.)
> run_fish600(iterations = 1, scenarios = c("M1-F1-D1-R1-cod"),
+ case_folder = case_folder, om_model_dir = om_model_dir,
+ em_model_dir = em_model_dir, bias_correct = TRUE,
+ bias_nsim = 1)
```

The flat scenario ID structure

There are many advantages to this flat scenario ID fold setup:

- 1. It makes it easier for multiple papers to share scenarios.
- 2. It makes it easier for papers to change which scenarios to compare after.
- 3. It avoids unnecessary nested folder structure.
- 4. It's easier to distribute the model runs across people and computers.
- 5. The functions are more general and applicable to future research.
- 6. Since each folder represents a unique scenario run, it's simple to keep track of progress on model runs in a spreadsheet

We can have a spreadsheet with the following columns:

Scenario ID, Scenario description, Model status

Then, groups can compile a list of scenario IDs they want to extract and compare.

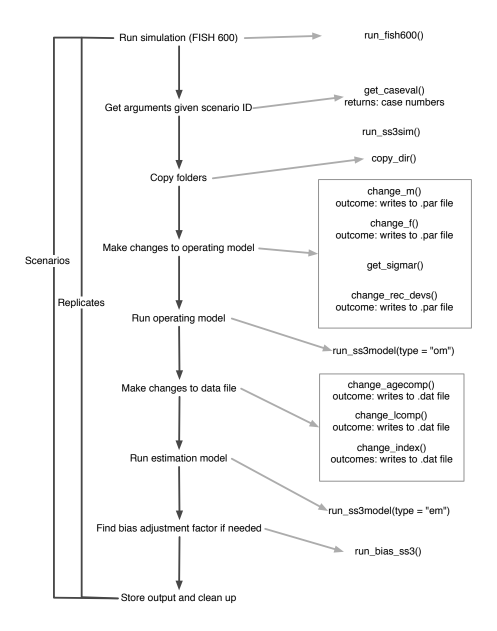


Figure 1: Simulation steps