Quantifying metapopulation portfolio effects with the ecofolio package

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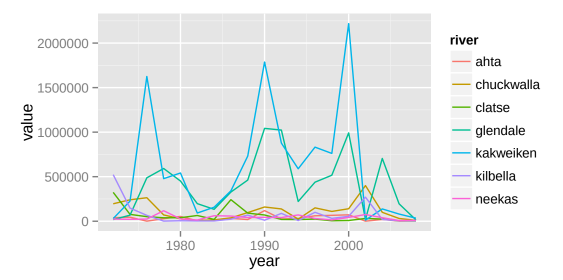
This vignette accompanies the ecofolio R package and the paper *Ecological prophets:* Quantifying metapopulation portfolio effects by Sean C. Anderson, Andrew B. Cooper, and Nicholas K. Dulvy, for the journal *Methods in Ecology and Evolution*. Here, we briefly illustrate the use and interpretation of the functions in the ecofolio package with a sample dataset.

First, let's load the package and load the sample dataset. These data represent pink salmon recruits in the Broughton Archipelago for even years. Each column represents abundance for a different river through time. You can find these data in the supplementary materials of Krkoŝek et al. (2011) and they were originally collected by Fisheries and Oceans Canada.

```
library(ecofolio)
data(pinkbr)
head(pinkbr)
           ahta chuckwalla clatse glendale kakweiken kilbella neekas
     year
## 1 1972 39611
                     195937 324023
                                       18815
                                                  29708
                                                          522499
                                                                   23565
## 2 1974 45248
                     240733
                             76827
                                       67873
                                                 226242
                                                          147115
                                                                   21341
## 3 1976
            812
                     263522
                              51206
                                      487456
                                                1624853
                                                           65881
                                                                   24181
## 4 1978 27983
                      70752
                              38581
                                      591956
                                                 477870
                                                             1415 117885
## 5 1980 54148
                      26999
                             39704
                                                             5400
                                                                  25524
                                      451232
                                                 541478
## 6 1982 4604
                       8156
                             64224
                                      197308
                                                  92077
                                                             3058
                                                                  13447
```

Let's plot the time series:

```
library(reshape)
library(ggplot2)
x_long <- melt(pinkbr, id.vars = "year", variable_name = "river")
ggplot(x_long, aes(year, value, colour = river)) + geom_line()</pre>
```



We can estimate Taylor's power law for this population:

```
fit_taylor(pinkbr[,-1])

## $c

## [1] 0.2806

##

## $z

## [1] 1.978
```

These values reflect the equation:

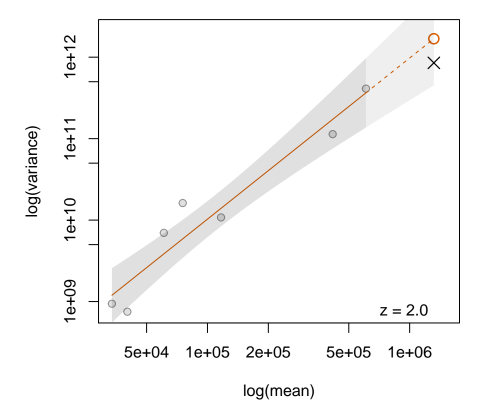
$$\sigma^2 = c\mu^z \tag{1}$$

where σ^2 is the temporal variance and μ is the temporal mean. The parameter c does not affect the portfolio effect. The parameter z is often referred to as Taylor's power law exponent. Specifically, fit_taylor and the function pe_mv fit the mean-variance relationship on a log-log scale:

$$log(\sigma_i^2) = c + z \cdot log(\mu_i) + \epsilon_i \tag{2}$$

where i represents an individual subpopulation, and ϵ_i represents independent and identically distributed residual error with mean zero and an estimated variance.

The package also contains a function plot_mv to visualize the various meanvariance model fits to empirical data. Let's look at the linear model we just fit:



In this plot, we see the $\log(\text{mean})$ and $\log(\text{variance})$ values for each river as grey dots. The orange line shows the mean-variance fit. The \times shows the observed metapopulation $\log(\text{mean})$ and $\log(\text{variance})$ values. The orange-open circle shows the expected $\log(\text{variance})$ value at the observed $\log(\text{mean})$ value for the metapopulation. The ratio of the y-value of the open-orange circle and the \times represents the mean-variance portfolio effect. It's the ratio of the expected to the observed variability at the metapopulation size.

So, z is around 2. Let's look at the default mean-variance portfolio effect, which uses the linear model we just fit:

```
pe_mv(pinkbr[,-1], ci = TRUE)

## $pe
## [1] 1.412
##
## $ci
## [1] 0.727 2.743
```

This tells us that the metapopulation is 1.4 times more stable than if it acted as a homogeneous population. These (frequentist) confidence intervals indicate that under repeated conditions, we would expect 95% of these intervals to contain the true value of the mean-variance portfolio effect given that our model is correct.

We can compare this to the average-CV portfolio effect. In this case, we find that the average-CV portfolio effect is similar for this population. We would expect this given that Taylor's power law z-value is close to 2.

```
pe_avg_cv(pinkbr[,-1], ci = TRUE, boot_reps = 500)
## $pe
## [1] 1.517
##
## $ci
## [1] 1.081 2.179
```

The pe_avg_cv function uses (bias-adjusted) bootstrap confidence intervals (BCa in the boot package). These confidence intervals are obtained by re-sampling the sub-populations with replacement and re-calculating the portfolio effect. If there aren't many subpopulations, these confidence intervals may not be an accurate reflection of the uncertainty in the average-CV portfolio effect.

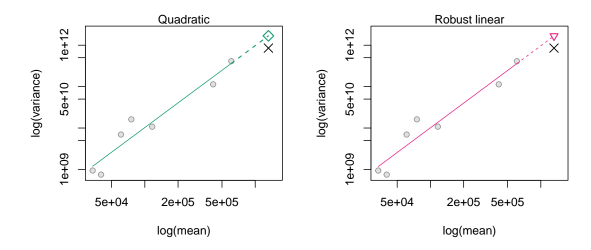
Returning to the mean-variance portfolio effect, we can explore whether different kinds of mean-variance extrapolations would yield different results. Let's fit some different mean-variance extrapolations:

```
pe_mv(pinkbr[,-1], type = "linear_robust")
## [1] 1.406
pe_mv(pinkbr[,-1], type = "quadratic")
```

```
## [1] 1.412
pe_mv(pinkbr[,-1], type = "linear_quad_avg")
## [1] 1.412
```

The quadratic and linear_quad_avg options give us the same value as the linear version because the quadratic model curvature has been bounded to only curve up. In this case, they simplify to the linear model. We could look at these using the plot_mv function:

```
par(mfrow = c(1, 2))
plot_mv(pinkbr[,-1], show = "quadratic", add_z = FALSE)
mtext("Quadratic")
plot_mv(pinkbr[,-1], show = "robust", add_z = FALSE)
mtext("Robust linear")
```



We can also try detrending the time series with linear or loess models to see how that affects the mean-variance portfolio effect. Estimates of variability such as the variance and CV can be biased upward if the time series are non-stationary.

```
pe_mv(pinkbr[,-1], type = "linear")
## [1] 1.412
```

```
pe_mv(pinkbr[,-1], type = "linear_detrended")

## [1] 1.422

pe_mv(pinkbr[,-1], type = "loess_detrended")

## [1] 1.43
```

We've suppressed the confidence intervals here for brevity, but we can see that for this population, the estimates are similar whether or not we detrend the data.

References

Krkoŝek, M., Connors, B.M., Morton, A., Lewis, M.A., Dill, L.M. & Hilborn, R. (2011). Effects of parasites from salmon farms on productivity of wild salmon. Proceedings of the National Academy of Sciences of the United States of America, 108, 14700–14704.