Do salamanders switch prey?

Michael Hutson

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1 Introduction

I am looking for a signal of prey switching by larval marbled salamanders (Ambystoma opacum), because such behavior could contribute to stabilizing their prey assemblage.

I use Chesson's (1983) measure of prey preference, comparing the proportion of the diet comprised by a given prey type against the proportion that the prey type makes up of all available prey in the environment. If a prey type becomes more frequent both in the environment and in the diet, this would not imply switching, and Chesson's preference value would similarly remain unchanged. However, if a prey type becomes more common in the environment and *even more so* in the diet, this would constitute a change in predator behavior in response to prey frequency. I apply this measure to a set of feeding trials where I controlled the available food types, and measured how many of each type were eaten.

In the data I have shared, the first few variables are probably peripheral for our purposes. The first important one is "weight", which is the measure for each salamander.

For the remainder:

"x.start" are the starting counts

"x.end" are the leftover counts

"x.eat" are the deduced number eaten

"x.a" are the preference values given by Chesson's measure

"x.e" are the preference values rescaled so that they range from -1 to +1

(-1 = never eaten regardless of frequency, 0 = equivalent proportion in diet and in environment, and +1 = always eaten regardless of frequency)

> data<-read.csv("data_short.csv",header=TRUE)</pre>

I first tried stepwise model simplification, testing the response with MANOVA:

2 Multivariate Multiple Regression

> # original stepwise model simplification approach
> with(data,{

```
trial.mlm1<-lm(cbind(clado.e,cope.e,ostr.e)~
                   weight*clado.start*cope.start)
    # remove weight:clado.start:cope.start
   trial.mlm2<-lm(cbind(clado.e,cope.e,ostr.e)~
                   weight*clado.start+weight*cope.start+clado.start:cope.start)
   # remove weight:clado.start
   trial.mlm3<-lm(cbind(clado.e,cope.e,ostr.e)~
                   weight+clado.start+cope.start+weight:cope.start
                  +clado.start:cope.start)
   # remove clado.start:cope.start
   trial.mlm4<-lm(cbind(clado.e,cope.e,ostr.e)~
                    weight+clado.start+cope.start+weight:cope.start)
   # trial.mlm4 is selected with this approach
     car::Manova(trial.mlm4, type="II")
+ })
Type II MANOVA Tests: Pillai test statistic
                 Df test stat approx F num Df den Df Pr(>F)
weight
                 1 0.65272 8.1446 3 13 0.0026256 **
                1 0.37524 2.6027 3 13 0.0964491 .
1 0.86810 28.5209 3 13 5.469e-06 ***
clado.start
cope.start
weight:cope.start 1 0.72039 11.1646 3 13 0.0006683 ***
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
  However, to be consistent with using AIC for examining single-taxon re-
```

sponses, I switched to AIC for the multiple multivariate models. I compare the full set of combinations of variables, including two models (6 & 7) that have preference independent of prey frequency.

```
> with(data, {
    trial.mlm1<-lm(cbind(clado.e,cope.e,ostr.e)~
                    weight*clado.start*cope.start)
    trial.mlm2<-lm(cbind(clado.e,cope.e,ostr.e)~
                    weight*clado.start+weight*cope.start+clado.start:cope.start)
   trial.mlm3<-lm(cbind(clado.e,cope.e,ostr.e)~
                    weight+clado.start+cope.start+weight:cope.start
                    +clado.start:cope.start)
   trial.mlm4<-lm(cbind(clado.e,cope.e,ostr.e)~
                     weight+clado.start+cope.start+weight:cope.start)
   trial.mlm5<-lm(cbind(clado.e,cope.e,ostr.e)~1)</pre>
   trial.mlm6<-lm(cbind(clado.e,cope.e,ostr.e)~weight)</pre>
    trial.mlm7<-lm(cbind(clado.e,cope.e,ostr.e)~weight+cope.start)</pre>
```

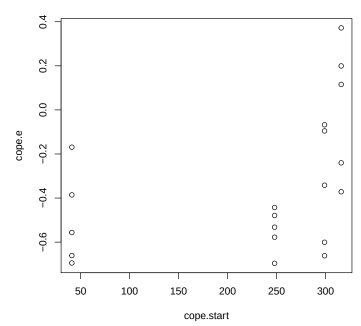
```
trial.mlm8<-lm(cbind(clado.e,cope.e,ostr.e)~weight+clado.start)
   trial.mlm9<-lm(cbind(clado.e,cope.e,ostr.e)~weight+ostr.start)</pre>
   trial.mlm10<-lm(cbind(clado.e,cope.e,ostr.e)~weight+cope.start+clado.start)
   trial.mlm14<-lm(cbind(clado.e,cope.e,ostr.e)~weight*cope.start)
   trial.mlm15<-lm(cbind(clado.e,cope.e,ostr.e)~weight*clado.start)
   trial.mlm16<-lm(cbind(clado.e,cope.e,ostr.e)~weight*ostr.start)</pre>
   trial.mlm17<-lm(cbind(clado.e,cope.e,ostr.e)~weight*clado.start+weight*cope.start)
   models<-list(trial.mlm1, trial.mlm2, trial.mlm3, trial.mlm4, trial.mlm5,
               trial.mlm6,trial.mlm7,trial.mlm8,trial.mlm9,trial.mlm10,
               trial.mlm14,trial.mlm15,
               trial.mlm16,trial.mlm17)
    (unlist(lapply(models,extractAIC)))
+ })
      48.00000 -115.09538
[1]
                             47.00000 -116.84824
                                                   46.00000 -104.78334
[7]
      45.00000 -92.17266
                             41.00000 -43.22355
                                                   42.00000 -49.85570
[13]
      43.00000 -85.39276
                             43.00000 -61.18062
                                                   43.00000 -82.99767
Г197
      44.00000 -90.57832
                             44.00000 -86.42151
                                                   44.00000 -60.37936
[25]
      44.00000 -84.10073
                             46.00000 -97.83989
```

AIC supports models that include prey frequency over models that do not. Therefore the salamanders are sensitive to prey frequency. However, this doesn't tell us how salamanders respond to any single prey taxon, which is why I am also trying to look at the single taxon responses.

3 Single taxon responses

Here I will focus only on copepods, since these are exemplary of the problems I have been facing.

When I looked at the response of cope.e to cope.start, there was clearly an un-



even increase:

My first instinct was to add a quadratic term.

```
> with(data,{
    cope.lm1<-lm(cope.e~weight*cope.start+I(cope.start^2))</pre>
    cope.lm2<-lm(cope.e~weight+cope.start+I(cope.start^2))</pre>
    cope.lm3<-lm(cope.e~cope.start+I(cope.start^2))</pre>
    cope.lm4<-lm(cope.e~weight*cope.start)</pre>
    cope.lm5<-lm(cope.e~weight+cope.start)</pre>
    cope.lm6<-lm(cope.e~cope.start)</pre>
    cope.lm7<-lm(cope.e~weight)</pre>
    cope.lm8<-lm(cope.e~1)</pre>
    cope.models<-list(cope.lm1,cope.lm2,cope.lm3,cope.lm4,cope.lm5,cope.lm6,</pre>
                      cope.lm7,cope.lm8)
    # per Burnham & Anderson, AICc is recommended when n/K<40
    (aicc<-unlist(lapply(cope.models,AICcmodavg::AICc)))</pre>
+ })
[1] 10.412972 6.788754 8.406833 15.661551 12.052644 13.311882 11.004474
[8] 14.153347
```

It looks like the best models are the ones that include cope.start+cope.start².

But I am not sure if the residuals here are showing worrying signs of pattern.

4 Next steps

So the last thing that I can think of trying is fitting an exponential model instead of a quadratic. An exponential curve would increase monotonically, as one might expect, and could fit with the increased slope of the preference curve at high prey frequencies. However, just as with the quadratic, the exponential curve would also start to predict values greater than 1, which is outside the allowable range of preference values. Also, I am not sure if it is possible to compare AIC values for different classes of models such as exponential vs quadratic.