mvMORPH: an R package for fitting multivariate evolutionary models

To illustrate how **mvMORPH** works, we describe and comment below two simulated examples. These examples are deliberately simple in order to show how the package works; therefore they are not directly related to any real biological context, although we will use names of known environments in order to make the understanding of objectives and results easier.

In the first example, we show how the package works in the classic situation of testing evolution towards two distinct selective regimes (Fig. 1). We simulate two correlated continuous traits evolving according to an Ornstein-Uhlenbeck (OU) process with two optima along the tree. We then fit an OU multivariate model with one (OU1) or two optima (OUM), and we compare the results with those obtained from separate univariate analyses on each simulated trait. This example is intended to show how covariations between traits can blur differences in optima that are not detected using a set of univariate analyses. In this specific case, we further show differences in type I and type II errors under univariate and multivariate settings; we also illustrate how to compare for significant interactions between traits in the path toward the optima.

In the second worked example, we compare the structure of evolutionary covariances in two, ecologically distinct species group using a nested hierarchy of tests thanks to the different rate matrix parameterizations proposed in **mvMORPH**. Such an approach is particularly useful to assess statistically how evolutionary covariations differ between groups of species. Indeed, we can expect various covariance structures depending on the selective pressures acting, e.g., on different ecological groups (e.g., low covariances for phenotypically diversified species when compared to more specialized ones, same directions of trait covariances, common structure...).

Note that the help pages of the **mvMORPH** package (accessible through the question mark argument in the R console; e.g., ?mvBM) also provide simulated examples illustrating how the various functions work.

Example 1: Models with distinct selective regimes

We first load the package and its dependencies, and initialize the random number generator:

```
# Load the package and dependencies (ape, phytools, corpcor, subplex, spam)
library(mvMORPH)
```

```
> Loading required package: phytools
> Loading required package: ape
> Loading required package: maps
> Loading required package: corpcor
> Loading required package: subplex
> Loading required package: spam
> Loading required package: grid
> Spam version 1.0-1 (2014-09-09) is loaded.
> Type 'help( Spam)' or 'demo( spam)' for a short introduction
> and overview of this package.
> Help for individual functions is also obtained by adding the
> suffix '.spam' to the function name, e.g. 'help( chol.spam)'.
>
> Attaching package: 'spam'
```

Then we simulate a random pure-birth tree with 100 taxa using the phytools package:

```
tree<-pbtree(n=100)
```

In this example, we are interested in evaluating the optima of two species groups (one optimum per species group) experiencing different environments. We first map on the tree the location of each regime (Fig. 1), and then simulate traits according to a multivariate OU process with two slightly distinct optima. For the sake of illustration, we hypothesize that the two selective regimes in this case are two well-known environments found in Africa: Savannah and Tropical Evergreen Forest. In real case studies, we may expect, e.g., that morpho-functional traits found in species occurring in each environment display differences in evolutionary rates, or adaptation toward distinct optima due to distinct selective pressures.

We use stochastic character mapping of discrete traits (SIMMAP; Huelsenbeck et al. 2003; Bollback 2006) as proposed by the make.simmap function from **phytools** for mapping the selective regimes:

```
# Simulate two selective regimes
state<-as.vector(c(rep("Forest",60),rep("Savannah",40))); names(state)<-tree$tip.label
# Make the tree with mapped states using SIMMAP
tree<-make.simmap(tree, state, model="ER", nsim=1)</pre>
> make.simmap is sampling character histories conditioned on the transition matrix
> Q =
>
                Forest
                           Savannah
           -0.02289711 0.02289711
> Forest
> Savannah 0.02289711 -0.02289711
> (estimated using likelihood);
> and (mean) root node prior probabilities
> pi =
    Forest Savannah
       0.5
                0.5
> Done.
```

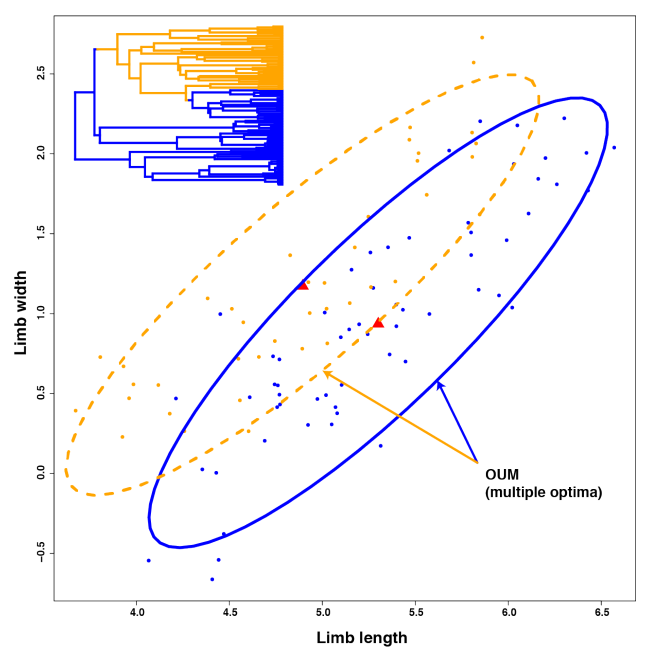
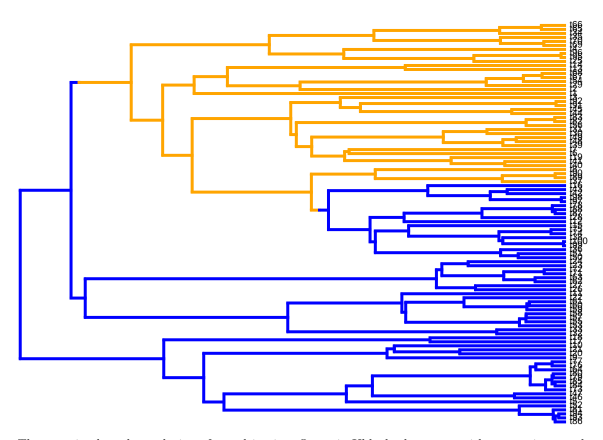


Figure 1. Simulated correlated traits under a multi-optimum OU process (function "mvSIM"). The tree was simulated according to a pure-birth process with 100 taxa and two selective regimes representing hypothesized "Savannah" and "Tropical Evergreen Forest" environments mapped on. The mapping is done using the "make.simmap" function from the phytools package, and is used as an input tree by the mvMORPH functions.

The resulting tree can be plotted with the mapped selective regimes as displayed in Fig. 1 for illustrative purpose:

```
# Plot the phylogeny with the mapped discrete trait
col<-c("blue","orange"); names(col)<-c("Forest","Savannah")
plotSimmap(tree,col, fsize=0.6, node.numbers=FALSE, lwd=3, pts=FALSE)</pre>
```



Then we simulate the evolution of a multivariate Ornstein-Uhlenbeck process with two optima on the tree. In order to illustrate the problem shown in Fig. 1 of the main text, we choose here a configuration where the leading eigenvector is oriented at \sim 45° in the bivariate space of the two evolving traits:

```
# 2 Random traits evolving along the phylogeny as a two-optimum OU
set.seed(101)

alpha<-matrix(c(1.1,-0.9,-0.9,1),2)
sigma<-matrix(c(0.35,0.06,0.06,0.35),2)
theta<-c(5.5,5.1,1.2,1.4)
data<-mvSIM(tree, param=list(sigma=sigma, alpha=alpha, ntraits=2, mu=theta,
names_traits=c("limb.length","limb.width")), model="OUM", nsim=1)</pre>
```

We now fit a single optimum and a multiple optimum Ornstein-Uhlenbeck process to each trait separately (i.e., univariate analysis), and then a single and a multiple OU process to both traits analyzed simultaneously (i.e., multivariate analysis):

```
# Fitting the Ornstein Uhlenbeck on the whole tree
trait1_0U1<- mv0U(tree, data[,1], model="0U1", diagnostic=FALSE, echo=FALSE)
trait2_0U1<- mv0U(tree, data[,2], model="0U1", diagnostic=FALSE, echo=FALSE)
# Fitting the Ornstein Uhlenbeck with multiple optimums
trait1_0UM<- mv0U(tree, data[,1], model="0UM", diagnostic=FALSE, echo=FALSE)
trait2_0UM<- mv0U(tree, data[,2], model="0UM", diagnostic=FALSE, echo=FALSE)
# Compare the AIC values between models fit
AIC(trait1_0UM); AIC(trait1_0U1)</pre>
```

```
> [1] 99.49821
AIC(trait2_OUM); AIC(trait2_OU1)
> [1] 128.1863
> [1] 127.6215
# Now compare with the multivariate fit
OUM<- mvOU(tree, data, model="OUM")</pre>
> successful convergence of the optimizer
> a reliable solution has been reached
> -- Summary results --
> LogLikelihood: -65.75399
> AIC: 151.508
> AICc:
           153.9799
> 10 parameters
> Estimated theta values
> limb.length limb.width
> Forest 5.314069 0.9277244
> Savannah 4.960028 1.2360596
> ML alpha values
      limb.length limb.width
> limb.length 1.787563 -1.511967
> limb.width -1.511967 1.698011
> ML sigma values
  limb.length limb.width
> limb.length 0.32448650 0.03069236
OU1<- mvOU(tree, data, model="OU1")
> successful convergence of the optimizer
> a reliable solution has been reached
> -- Summary results --
> LogLikelihood: -85.98834
> AIC: 187.9767
> AICc:
       189.5591
> 8 parameters
> Estimated theta values
```

limb.length limb.width

```
> OU1 5.178202 1.091494
>
> ML alpha values
> ______
> limb.length limb.width
> limb.length 0.4883319 -0.3071923
> limb.width -0.3071923 0.5873142
>
> ML sigma values
> _____
> limb.length limb.width
> limb.length 0.2538671 0.1145218
> limb.width 0.1145218 0.3924766
```

```
AIC(OUM); AIC(OU1)
```

- > [1] 151.508
- > [1] 187.9767

After model fitting, we extract their related AIC or AICc using the expression "\$AIC" or with the AIC() generic method. Despite of the low number of parameters to estimate in the univariate analysis (3 and 4 for the OU1 and OUM models, respectively), it is difficult to distinguish between the two models and the unique optimum OU1 is slightly preferred. In stark contrast, the multivariate analysis clearly favors the actual simulated (generating) model (two optima) with a delta AIC > 35.

We can further assess the type I and type II error rates through simulations. Note that parametric bootstrapping and simulations could be parallelized using forking with the **parallel** package to speed up the computations:

```
# Simulate 1000 traits under a two optima and unique OU process
library(parallel)
nsim=1000
data1<-mvSIM(tree, param=list(sigma=sigma, alpha=alpha, ntraits=2, mu=theta,
          names_traits=c("head.size","mouth.size")), model="OUM", nsim=nsim)
data2<-mvSIM(tree, param=list(sigma=sigma, alpha=alpha, ntraits=2, mu=theta2,
          names_traits=c("head.size","mouth.size")), model="OU1", nsim=nsim)
# Fit of the models using the parallel package (we will use 2 cores), can take a while...
library(parallel)
nb cores=2L
oum_data1<- mclapply(1:nsim, function(x){</pre>
   mv0U(tree, data1[[x]], model="0UM", method="sparse", diagnostic=F, echo=F)
}, mc.cores = getOption("mc.cores", nb_cores))
ou1_data1<- mclapply(1:nsim, function(x){
    mv0U(tree, data1[[x]], model="0U1", method="sparse", diagnostic=F, echo=F)
}, mc.cores = getOption("mc.cores", nb_cores))
# Now same simulations on the second dataset
oum_data2<- mclapply(1:nsim, function(x){</pre>
```

```
mv0U(tree, data1[[x]], model="0UM", method="sparse", diagnostic=F, echo=F)
}, mc.cores = getOption("mc.cores", nb_cores))
ou1_data2<- mclapply(1:nsim, function(x){
    mvOU(tree, data1[[x]], model="OU1", method="sparse", diagnostic=F, echo=F)
}, mc.cores = getOption("mc.cores", nb_cores))
# Retrieve the results from the simulations
OUM_simul<-sapply(1:nsim, function(x){</pre>
    c(oum_data1[[x]]$AICc,ou1_data1[[x]]$AICc)
})
OU1 simul <- sapply (1:nsim, function(x) {
    c(oum_data2[[x]]$AICc,ou1_data2[[x]]$AICc)
})
# Now compute the type I & II errors
sum(OU1_simul[1,]<OU1_simul[2,])/nsim</pre>
[1] 0.141
sum(OUM_simul[1,]<OUM_simul[2,])/nsim</pre>
```

The simulations in this example shows that both type I and type II errors are quite low (although the type I error is $\sim 14\%$) comparing to the type I and type II errors obtained for the univariate fits (Table 1).

Table 1. Summary of type I and type II errors resulting from the univariate analysis of the simulated dataset.

	Trait 1	Trait 2
Type II	0.538	0.669
Type I	0.221	0.217

Finally, constraining the model fit to an OU process without interactions further shows that the preferred model is indicative of significant interactions between traits in the path toward the optima.

```
> -- Log-likelihood Ratio Test --
> Model OUM symmetric positive versus OUM diagonal
> Number of degrees of freedom : 1
> LRT statistic: 33.44478 p-value: 7.331641e-09 ***
> ---
> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

If we are interested in the evolutionary variance and covariances or correlations, we can use the *stationary* function with the fitted OU object.

```
stationary(OUM)
              limb.length limb.width
> limb.length
                0.4658481 0.4434552
> limb.width
                0.4434552 0.5361603
# we can use the cov2cor function from the R stats package
# to get the evolutionary correlations between traits
cov2cor(stationary(OUM))
              limb.length limb.width
> limb.length
                  1.00000
                             0.88732
> limb.width
                  0.88732
                             1.00000
```

Example 2: Tests for evolutionary correlations between two species groups

In this second simulated example we illustrate how **mvMORPH** can be used for assessing correlated evolution and evolutionary covariance structures in different selective regimes. Constrained models are used to compare different hypotheses of covariations between traits organized hierarchically from a common structure shared between species group, proportional evolutionary covariances, similar variances but different correlations (shared eigenvectors), similar correlations but different variances, and completely unrelated covariance structures.

We first simulate a tree with two mapped regimes, and simulate traits with unrelated covariances in each group (Fig. 2). Unrelated covariance structures can result from different selective pressures within each group, while the other sub-models assume intermediate hypotheses of shared covariance structures such as, e.g., evolution along the same variance directions.

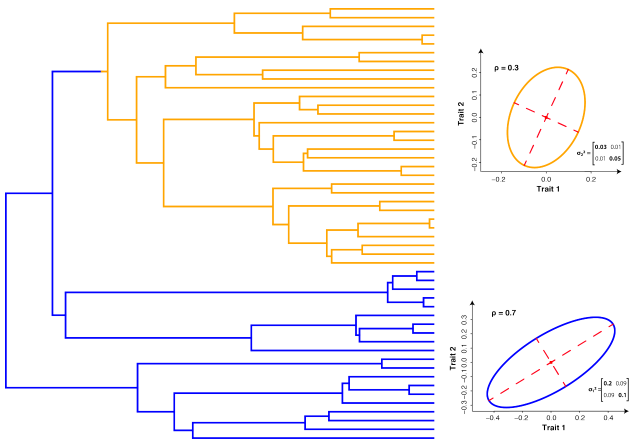


Figure 2. Simulated tree (50 tips) with mapped selective regimes and associated two-trait covariance structures. The first rate matrix associated with the first mapped regime ("blue") show a correlation between traits of 0.7, and the second rate matrix a correlation of 0.3.

Now we fit five nested BM models. We first compare a multiple (unrelated) rate matrix ("BMM") model with an equal rate matrix ("BM1") model. Then we fit a multiple rate matrix for which:

- we constrain the variances to be similar but not the covariances using the "param" list (param=list(constraint="shared")),
- a multiple rate matrix model for which the correlations are identical between groups (param=list(constraint="correlation")),
- and a multiple rate matrix model for which the matrices differs by a proportional scalar (param=list(constraint="proportional")).

AIC and Log-likelihood ratio tests are finally used to compare the four nested models.

```
# Fitting the models
# BM1 - (Equal rate matrix)
model_1<-mvBM(tree, data, model="BM1", diagnostic=FALSE, echo=FALSE)</pre>
# BMM - (Proportional rate matrices)
model_2<-mvBM(tree, data, param=list(constraint="proportional")</pre>
               , diagnostic=FALSE, echo=FALSE)
# BMM - (Shared eigenvectors between rate matrices)
model_3<-mvBM(tree, data, param=list(constraint="shared")</pre>
              , diagnostic=FALSE, echo=FALSE)
# BMM - (Similar correlations between rate matrices)
model_4<-mvBM(tree, data, param=list(constraint="correlation")</pre>
              , diagnostic=FALSE, echo=FALSE)
# BMM - (independent rate matrices)
model_5<-mvBM(tree, data, model="BMM", diagnostic=FALSE, echo=FALSE)
# Compare the models with AIC
AIC(model_1)
```

> [1] 37.4709

```
AIC(model_2)
> [1] 31.20892
AIC(model_3)
> [1] 38.76954
AIC(model_4)
> [1] 29.59168
AIC(model_5)
> [1] 25.99354
# Test significance with LRT
LRT(model_5,model_4)
> -- Log-likelihood Ratio Test --
> Model BMM versus BMM equal correlation
> Number of degrees of freedom : 1
> LRT statistic: 5.598139 p-value: 0.01797956 *
> Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
LRT(model_5,model_3)
> -- Log-likelihood Ratio Test --
> Model BMM versus BMM shared variances
> Number of degrees of freedom : 2
> LRT statistic: 16.776 p-value: 0.0002275816 ***
> Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
LRT(model_5,model_2)
> -- Log-likelihood Ratio Test --
> Model BMM versus BMM proportional
> Number of degrees of freedom : 2
> LRT statistic: 9.21538 p-value: 0.009974832 **
> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
LRT(model_5,model_1)
```

```
> -- Log-likelihood Ratio Test --
> Model BMM versus BM1
> Number of degrees of freedom : 3
> LRT statistic: 17.47736 p-value: 0.0005636605 ***
> ---
> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

As expected, AIC criterion shows that the generating "unrelated matrix model" we simulated is preferred over all other fitted models. LRT tests further show that the difference with the second best-fit models is significant (at a 0.05 level). From the rates matrices (evolutionary covariances), we can compute the evolutionary correlation in both groups using the *cov2cor* function in the R package **stats**.

Note that the alternate generating models proposed in $\mathbf{mvMORPH}$ (e.g., mvOU, mvEB) can also be used to assess whether a Brownian motion process is a good approximation for describing the simulated evolutionary pattern.

References:

Bollback, J.B. (2006). SIMMAP: Stochastic character mapping of discrete traits on phylogenies. *BMC Bioinformatics*, 7, 1-7.

Huelsenbeck, J.P., Nielsen, R. & Bollback, J.B. (2003). Stochastic mapping of morphological characters. *Systematic Biology*, 52, 131–158.