## Introduction to the surface package

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This tutorial walks through the steps of a SURFACE analysis using a simple demonstration data set. The main use of SURFACE is to construct a macroevolutionary adaptive landscape for a clade, given only a phylogenetic tree and measurements of one or more continuous traits for each member species. This is done by using stepwise AIC algorithms to fit a series of 'Hansen' models using ouch functions, with two distinct phases. In the *forward* phase, new selective regimes are added to the model, and in the *backward* phase multiple regimes may be 'collapsed' into convergent regimes discovered independently by different lineages. The final model provides an estimate of the macroevolutionary adaptive landscape, and can be compared to simulated data to evaluate whether the clade contains exceptional phenotypic convergence.

The surface package can be downloaded from CRAN (http://cran.r-project.org/web/packages/surface/). Also required are the packages ape, ouch, geiger and igraph and their dependencies. First, load the surface package along with its dependencies, then import the data object surfaceDemo contained in the package. This object consists of a 25-taxon tree in phylo format, and a simulated data set that includes measurements of three continuous traits for each species.

- > library(surface)
- > data(surfaceDemo)
- > tree<-surfaceDemo\$tree
- > dat<-surfaceDemo\$sim\$dat

At this point, the tree and dat objects could be provided as input to the wrapper function runSurface, which carries out both phases of the analysis in a single step. Instead, let's look at the two steps separately to make the sequence of events clear. The function nameNodes ensures that each node in the tree has a unique label, for easier conversion between formats later (if nodes are already labeled, as is the case for the demo tree, the function returns the tree unchanged). convertTreeData converts the tree and data objects into a format ready for analysis with the ouch function hansen.

- > tree<-nameNodes(tree)
- > olist<-convertTreeData(tree,dat)
- > otree<-olist[[1]]; odata<-olist[[2]]</pre>

The data set is now ready for analysis with SURFACE. We use the function surfaceForward to add one selective regimes at a time to the model, by repeatedly calling the function addRegime. By default, the starting model is a single-regime OU model, although one can optionally start an analysis with a model containing some regimes by specifying a starting\_list.

The default arguments should be suitable for many analyses, but one can specify options such as a different aic\_threshold for accepting model improvements, a limit to the max\_steps of the algorithm, whether to sample\_shifts within sample\_threshold AIC units of the best model rather than choosing the best model at each step, and whether to exclude a fraction of the worst candidate models from the previous step (see documentation for surfaceForward for more). By default progress is not printed out, but one can choose to print (fairly extensive) output to the console with verbose=TRUE, to view a visual representation of the change in AIC at each step in the graphics device with plotaic=TRUE, or to save the output to a file filename at each step to guard against crashes or exceeding run time limits using save\_steps=TRUE. This step should take a little under a minute, but can be quite long for larger data sets.

```
> fwd<-surfaceForward(otree, odata, aic_threshold = 0, exclude = 0,
+ verbose = FALSE, plotaic = FALSE)
> k<-length(fwd)</pre>
```

The object returned by surfaceForward, which has been named fwd, is a list of length k, each element of which is another list containing the output of one call to addRegime. The object fwd is unwieldy to view all at once, but surfaceSummary can be used to retrieve information about the forward phase, such as the sequence of AIC values:

```
> fsum<-surfaceSummary(fwd)</pre>
```

> names(fsum)

```
[1] "n_steps" "lnls" "n_regimes_seq" "aics"
```

[5] "shifts" "n\_regimes" "alpha" "phylhalflife"

[9] "sigma\_squared" "theta"

> fsum\$aics

surfaceSummary also returns the parameter estimates and regime placements from fwd[[k]], the model returned from the forward phase of SURFACE, which is also the starting model for the backward phase. surfaceBackward repeatedly calls the function collapseRegimes, identifying cases where the same (or very similar) regimes are found independently on different branches of the tree, and where the model simplification obtained by collapsing them into single regimes results in a further improvement in the AIC. By default multiple regimes can be collapsed at each step if they are mutually compatible (see documentation), but will be limited to one collapse per step if the option only\_best=TRUE is specified. Other defaults and options are similar to the forward phase. This step will run in a few seconds, but the time taken increases quickly with the number of taxa and the number of regimes found in the forward phase.

```
> bwd<-surfaceBackward(otree, odata, starting_model = fwd[[k]], aic_threshold = 0,
```

- + only\_best = FALSE, verbose = FALSE, plotaic = FALSE)
- > bsum<-surfaceSummary(bwd)</pre>
- > kk<-length(bwd)

surfaceBackward returns another list of lists, and again the final element (bwd[[kk]]) is generally the element of interest. surfaceSummary again summarizes the steps and final parameter values, and also displays measures of the extent of convergence in the data set (deltak and c).

> bsum\$alpha

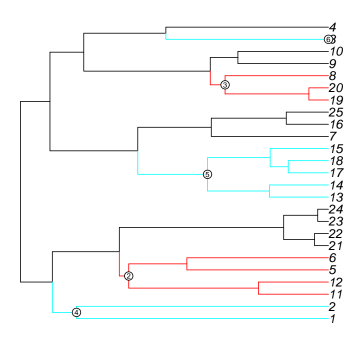
> bsum\$sigma\_squared

> bsum\$theta

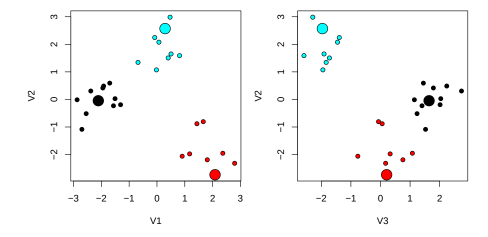
> bsum\$n\_regimes

The fitted SURFACE model can be visualized either as paintings of regimes on the tree, or as color-coded points in trait space. Colors can be generated automatically, and by default convcol = TRUE, meaning convergent regimes are colorful and non-convergent regimes are grey or black.

> surfaceTreePlot(tree, bwd[[kk]], labelshifts = T)



- > par(mfrow=c(1,2), mai=c(0.8,0.8,0.2,0.2))
- > surfaceTraitPlot(dat, bwd[[kk]], whattraits = c(1,2))
- > surfaceTraitPlot(dat, bwd[[kk]], whattraits = c(3,2))



Another way to visualize the results of the SURFACE analysis is with the sequence of AICc values from phase of the analysis. It may be interesting to compare this to the AICc of simpler models. The Brownian motion (BM) and single-regime OU (OU1) models can be fitted using starting-Model, which calculates AICc by adding log-likelihoods across traits so they are comparable to the SURFACE AICc values.

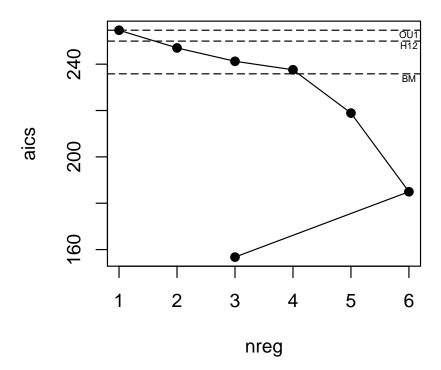
- > bm<-startingModel(otree,odata,brownian=TRUE)
- > ou1<-startingModel(otree,odata)

startingModel can also be used to fit a Hansen model that includes specific shift placements. While the purpose of SURFACE is to avoid such a priori information, it may be of interest to compare the model produced by SURFACE to other models, or to use SURFACE to add shifts while forcing certain other shifts to be present. To impose some regime shifts, one can build the desired model using startingModel, and supply this object to the starting\_list argument in surfaceForward. Here, we will simply fit a hypothetical alternative model 'H12', which we will pretend is based on a priori information about shifts into two habitats (H1 and H2).

> H12<-startingModel(otree,odata,shifts=c("26"="H1","13"="H1","5"="H2","19"="H2"))

Now, we can view the sequence of AICc values from the SURFACE run, and compare it to the values for the other models.

- > surfaceAICPlot(fwd, bwd)
- > abline(h=bm[[1]]\$aic,lty="longdash")
- > abline(h=H12[[1]]\$aic,lty="longdash")
- > text(c(6,6),c(bm[[1]]\$aic, ou1[[1]]\$aic, H12[[1]]\$aic)-2,c("BM","OU1","H12"),cex=0.5)

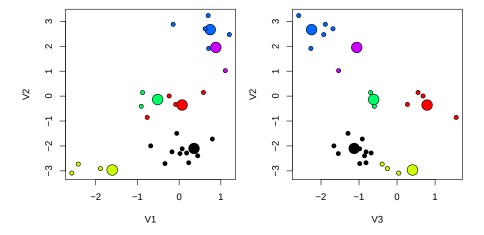


The function propRegMatch can be used to compare the fitted model to the 'true', generating model based on the proportion of pairs of branches (either all branches or only taxa at the tip branches) that are correctly assigned to either the same regime or to different regimes. In this simple case, SURFACE recovers the generating model perfectly, so both values are 1.

- > truefit<-surfaceDemo\$sim\$fit
  > propRegMatch(truefit, bwd[[kk]]\$fit, internal = FALSE)
- [1] 1
- > propRegMatch(truefit, bwd[[kk]]\$fit, internal = TRUE)
- [1] 1

To compare the extent of convergence in a data set to a null expectation, data can be simulated under a model that lacks true convergence: either a simple stochastic model such as Brownian motion, or a Hansen model based on the fitted model from the forward phase; with multiple evolutionary regime shifts to *non-convergent* adaptive peaks. Either can be done using surfaceSimulate: the following code carries out one simulation under the Hansen null model (set.seed is used here only for reproducibility).

- > set.seed(10)
  > newsim<-surfaceSimulate(tree, type="hansen-fit", hansenfit=fwd[[k]]\$fit,
  + shifts=fwd[[k]]\$savedshifts, sample\_optima=TRUE)</pre>
- We can then visualize the simulated trait data; as there is no convergence in the true model, setting convcol = FALSE makes a more colorful figure.
- > par(mfrow=c(1,2),mai=c(0.8,0.8,0.2,0.2))
- > surfaceTraitPlot(newsim\$data, newsim, whattraits = c(1,2), convcol = FALSE)
- > surfaceTraitPlot(newsim\$data, newsim, whattraits = c(3,2), convcol = FALSE)



We can then run SURFACE on the simulated data set, here doing the entire analysis in one step using runSurface (this should take less than a minute). We can then use surfaceSummary to extract the results, and compare the number of regime shifts (k) and the extent of convergence (deltak or c) to what we saw in the 'real' data set. In this case, one instance of convergence is recovered in the simulated data set where two regimes were relatively close to one another in trait space; as we know that the regimes were nonetheless distinct from one another in the generating model, this represents 'incidental' convergence.

- > newout<-runSurface(tree, newsim\$dat)</pre>
- > newsum<-surfaceSummary(newout\$bwd)</pre>
- > newkk<-length(newout\$bwd)</pre>
- > newsum\$n\_regimes

```
        k
        kprime
        deltak
        c
        kprime_conv

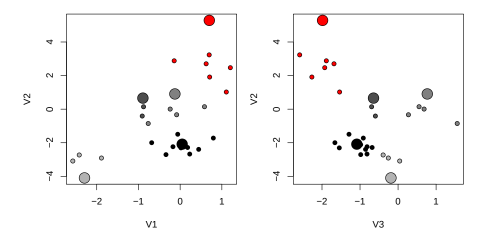
        6
        5
        1
        2
        1

        kprime_nonconv
        4
        4
        4
        4
```

> bsum\$n\_regimes

k	kprime	deltak	С	kprime_conv
6	3	3	5	2
kprime_nonconv				
1				

- > par(mfrow=c(1,2),mai=c(0.8,0.8,0.2,0.2))
- > surfaceTraitPlot(newsim\$data, newout\$bwd[[newkk]], whattraits = c(1,2))
- > surfaceTraitPlot(newsim\$data, newout\$bwd[[newkk]], whattraits = c(3,2))



If we wanted to carry out a hypothesis test of whether the 'real' data set contains more convergence than expected by chance, we could repeat the step of simulating data sets (with different random number seeds) many times, and running SURFACE on them to get a null distribution of deltak or c. We could calculate the statistical significance of such a test as the proportion of null values that meet or exceed the observed value (note that in a small data set like this statistical power is poor, but that statistical properties are much better given larger trees and at least two trait dimensions).

For more information about the method, see the manuscript in *Methods in Ecology and Evolution*, the help files for each function, and the SURFACE homepage (http://www.people.fas.harvard.edu/ingram/web/surface.html).