'SongEvo' package

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Introduction

SongEvo simulates the cultural evolution of quantitative traits of bird song. SongEvo is an individual- (agent-) based model. SongEvo is spatially-explicit and can be parameterized with, and tested against, measured song data. Functions are available for model implementation, sensitivity analyses, parameter optimization, model validation, and hypothesis testing.

Overview of Functions

- 1. SongEvo implements the model
- 2. par.sens allows sensitivity analyses
- 3. par.opt allows parameter optimization
- 4. mod.val allows model validation
- 5. h.test allows hypothesis testing

Getting Started

Load and attach SongEvo package

library(SongEvo)

Functions

 ${\tt SongEvo~implements~the~model~par.sens~allows~sensitivity~analyses~par.opt~allows~parameter~optimization~mod.val~allows~model~validation~h.test~allows~hypothesis~testing}$

Examples

EXAMPLE 1

Load the example data: song.data and global parameters

To explore the SongEvo package, we will use a database of songs from Nuttall's white-crowned sparrow (Zonotrichia leucophrys nuttalli) recorded at three locations in 1969 and 2005.

data("song.data")

Examine global parameters. Global parameters describe our understanding of the system and may be measured or hypothesized. They are called "global" because they are used by many many functions and subroutines within functions. For descriptions of all adjustable parameters, see ?song.data.

```
data("glo.parms")
str(glo.parms)
#> List of 15
#> $ learning.error.d : num 0
#> $ learning.error.sd : num 430
#> $ n.territories : num 40
#> $ mortality.a
                     : num 0.468
#> $ mortality.j
                     : num 0.5
#> $ lifespan
                      : num 2.08
#> $ phys.lim.min : num 1559
#> $ phys.lim.max : num 4364
#> $ male.fledge.n.mean: num 1.35
\# $ male.fledge.n.sd : num 0.5
#> $ disp.age
                 : num 2
#> $ disp.distance.mean: num 110
#> $ disp.distance.sd : num 100
#> $ terr.turnover : num 0.5
#> $ male.fledge.n : num [1:40] 1 1 2 1 0 2 2 2 2 1 ...
```

Share global parameters with the global environment. We make these parameters available in the global environment so that we can access them with minimal code.

```
list2env(glo.parms, globalenv())
#> <environment: R_GlobalEnv>
```

Examine song data

Data include the population name (Bear Valley, PRBO, or Schooner), year of song recording (1969 or 2005), and the frequency bandwidth of the trill.

Simulate bird song evolution with SongEvo()

Define initial individuals

In this example, we use songs from individual birds recorded in one population (PRBO) in the year 1969, which we will call starting.trait.

```
starting.trait <- subset(song.data, Population=="PRBO" & Year==1969)$Trill.FBW
```

We want a starting population of 40 individuals, so we generate additional trait values to complement those from the existing 30 individuals. Then we create a data frame that includes a row for each individual; we add identification numbers, ages, and geographical coordinates for each individual.

```
starting.trait2 <- c(starting.trait, rnorm(n.territories-length(starting.trait),
    mean=mean(starting.trait), sd=sd(starting.trait)))
init.inds <- data.frame(id = seq(1:n.territories), age = 2, trait = starting.trait2)
init.inds$x1 <- round(runif(n.territories, min=-122.481858, max=-122.447270), digits=8)
init.inds$y1 <- round(runif(n.territories, min=37.787768, max=37.805645), digits=8)</pre>
```

Specify and call the SongEvo model

SongEvo() includes several settings, which we specify before running the model. For this example, we run the model for 10 iterations, over 36 years (i.e. 1969–2005). When conducting research with SongEvo(), users will want to increase the number iterations (e.g. to 100 or 1000). Each timestep is one year in this model (i.e. individuals complete all components of the model in 1 year). We specify territory turnover rate here as an example of how to adjust parameter values. We could adjust any other parameter value here also. The learning method specifies that individuals integrate songs heard from adults within the specified integration distance (intigrate.dist, in kilometers). In this example, we do not include a lifespan, so we assign it NA. In this example, we do not model competition for mates, so specify it as FALSE. Last, specify all as TRUE in order to save data for every single simulated individual because we will use those data later for mapping. If we do not need data for each individual, we set all to FALSE because the all.inds data.frame becomes very large!

```
iteration <- 10
years <- 36
timestep <- 1
terr.turnover <- 0.5
learning.method <- "integrate"
integrate.dist <- 0.1
lifespan <- NA
mate.comp <- FALSE
prin <- FALSE
all <- TRUE</pre>
```

Now we call SongEvo with our specifications and save it in an object called SongEvo1.

```
SongEvo1 <- SongEvo(init.inds = init.inds, iteration = iteration, steps = years,
    timestep = timestep, n.territories = n.territories, terr.turnover = terr.turnover,
    learning.method = learning.method, integrate.dist = integrate.dist,
    learning.error.d = learning.error.d, learning.error.sd = learning.error.sd,
    mortality.a = mortality.a, mortality.j = mortality.j, lifespan = lifespan,
    phys.lim.min = phys.lim.min, phys.lim.max = phys.lim.max,
    male.fledge.n.mean = male.fledge.n.mean, male.fledge.n.sd = male.fledge.n.sd,
    male.fledge.n = male.fledge.n, disp.age = disp.age,
    disp.distance.mean = disp.distance.mean, disp.distance.sd = disp.distance.sd,
    mate.comp = mate.comp, prin = prin, all)</pre>
```

Examine results from SongEvo model

The model required the following time to run on your computer:

```
SongEvo1$time
#> user system elapsed
#> 2.082  0.046  2.136
```

Three main objects hold data regarding the SongEvo model. Additional objects are used temporarily within modules of the model.

First, currently alive individuals are stored in a data frame called "inds." Values within "inds" are updated throughout each of the iterations of the model, and "inds" can be viewed after the model is completed.

```
head(SongEvo1$inds, 5)

#> coordinates id age trait x1 y1

#> 1447 (-122.456, 37.79347) 1447 10 4340.245 -122.4560 37.79347

#> 1475 (-122.4806, 37.8062) 1475 10 4364.000 -122.4806 37.80620

#> 1668 (-122.4569, 37.79691) 1668 6 2362.918 -122.4569 37.79691
```

```
#> 1681 (-122.4587, 37.79652) 1681
                                      6 3593.968 -122.4587 37.79652
#> 1728 (-122.4572, 37.79628) 1728
                                      5 3548.710 -122.4572 37.79628
#>
        male.fledglings territory father
                                                 x0
                                                           y0
                                     1050 -122.4561 37.79483
#> 1447
                      0
                                 0
#> 1475
                      0
                                 0
                                     1409 -122.4810 37.80569
                      0
                                 0
#> 1668
                                     1533 -122.4569 37.79621
#> 1681
                      0
                                 0
                                     1600 -122.4581 37.79600
#> 1728
                                     1600 -122.4581 37.79600
```

Second, an array (i.e. a multi-dimensional table) entitled "summary.results" includes population summary values for each time step (dimension 1) in each iteration (dimension 2) of the model. Population summary values are contained in five additional dimensions: population size for each time step of each iteration ("sample.n"), the population mean and variance of the song feature studied ("trait.pop.mean" and "trait.pop.variance"), with associated lower ("lci") and upper ("uci") confidence intervals.

```
dimnames(SongEvo1$summary.results)
#> $iteration
   [1] "iteration 1"
                      "iteration 2"
                                      "iteration 3"
                                                     "iteration 4"
    [5] "iteration 5"
                       "iteration 6"
                                      "iteration 7"
                                                     "iteration 8"
    [9] "iteration 9"
                       "iteration 10"
#>
#> $step
  [1] "1" "2" "3" "4" "5" "6"
                                      "7"
                                           "8"
                                                "9" "10" "11" "12" "13" "14"
#> [15] "15" "16" "17" "18" "19" "20" "21" "22" "23" "24" "25" "26" "27" "28"
#> [29] "29" "30" "31" "32" "33" "34" "35" "36"
#>
#> $feature
#> [1] "sample.n"
                            "trait.pop.mean"
                                                 "trait.pop.variance"
#> [4] "lci"
                            "uci"
```

Third, individual values may optionally be concatenated and saved to one data frame entitled "all.inds." all.inds can become quite large, and is therefore only recommended if additional data analyses are desired.

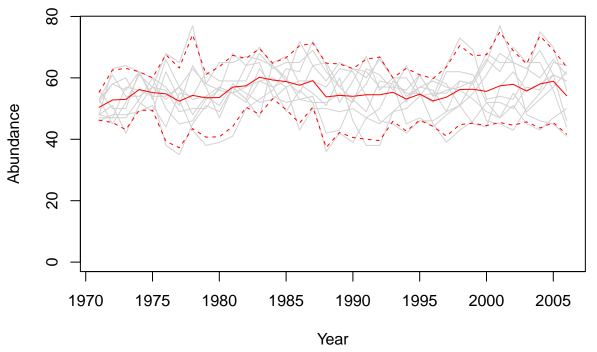
```
head(SongEvo1$all.inds, 5)
#>
                     coordinates id age trait
#> I1.T1.1 (-122.4799, 37.78858)
                                 1 2 4004.8 -122.4799 37.78858
#> I1.T1.2 (-122.4712, 37.79482)
                                  2
                                     2 3765.0 -122.4712 37.79482
                                    2 3237.4 -122.4739 37.79595
#> I1.T1.3 (-122.4739, 37.79595)
                                 3
#> I1.T1.4 (-122.4518, 37.8041) 4
                                      2 3621.1 -122.4518 37.80410
#> I1.T1.5 (-122.4606, 37.79133) 5 2 3285.4 -122.4606 37.79133
#>
           male.fledglings territory father x0 y0 timestep iteration
                                          0 0 0
                                                                    1
#> I1.T1.1
                                   1
                                                         1
                         1
#> I1.T1.2
                         1
                                   1
                                          0
                                             0 0
                                                          1
                                                                    1
                                          0
#> I1.T1.3
                         2
                                   1
                                             0
                                               0
                                                                    1
                                                          1
#> I1.T1.4
                         1
                                   1
                                          0
                                                0
                                                          1
                                                                    1
#> I1.T1.5
```

Simulated population size

We see that the simulated population size remains relatively stable over the course of 36 years. This code uses the summary results array.

```
for(p in 1:iteration){
    lines(SongEvo1$summary.results[p, , "sample.n"], col="light gray")
    }
n.mean <- apply(SongEvo1$summary.results[, , "sample.n"], 2, mean, na.rm=TRUE)
lines(n.mean, col="red")

#Plot 95% quantiles
quant.means <- apply (SongEvo1$summary.results[, , "sample.n"], MARGIN=2, quantile,
    probs=c(0.975, 0.025), R=600, na.rm=TRUE)
lines(quant.means[1,], col="red", lty=2)
lines(quant.means[2,], col="red", lty=2)</pre>
```



Load Hmisc package for plotting functions.

```
library("Hmisc")
```

Simulated trait values

We see that the mean trait values per iteration varied widely, though mean trait values over all iterations remained relatively stable. This code uses the summary results array.

```
#Plot 95% quantiles
quant.means <- apply (SongEvo1$summary.results[, , "trait.pop.mean"], MARGIN=2, quantile,
    probs=c(0.95, 0.05), R=600, na.rm=TRUE)
lines(quant.means[1,], col="blue", lty=2)
lines(quant.means[2,], col="blue", lty=2)
#plot mean and CI for historic songs.
#plot original song values
library("boot")
#> Attaching package: 'boot'
#> The following object is masked from 'package:survival':
#>
       aml
#> The following object is masked from 'package:lattice':
#>
       melanoma
sample.mean <- function(d, x) {</pre>
    mean(d[x])
}
boot_hist <- boot(starting.trait, statistic=sample.mean, R=100)#, strata=mn.res$iteration)
ci.hist <- boot.ci(boot_hist, conf=0.95, type="basic")</pre>
low <- ci.hist$basic[4]</pre>
high <- ci.hist$basic[5]</pre>
points(0, mean(starting.trait), pch=20, cex=0.6, col="black")
errbar(x=0, y=mean(starting.trait), high, low, add=TRUE)
 #text and arrows
text(x=5, y=2720, labels="Historical songs", pos=1)
arrows(x0=5, y0=2750, x1=0.4, y1=mean(starting.trait), length=0.1)
      4000
      3500
Bandwidth (Hz)
                Historical songs
      2500
             1970
                                                                                 2005
                      1975
                                1980
                                          1985
                                                    1990
                                                             1995
                                                                       2000
```

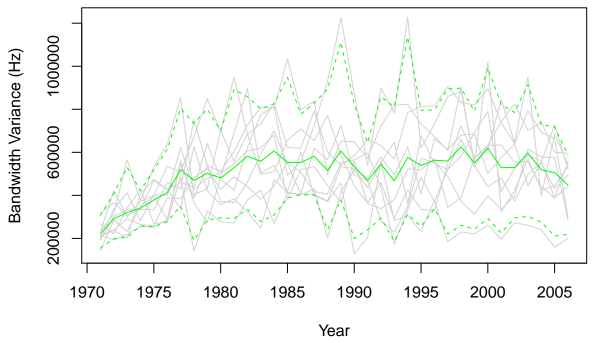
Trait variance

Year

We see that variance for each iteration per year increased in the first few years and then stabilized. This code uses the summary results array.

```
#plot variance for each iteration per year
plot(SongEvo1$summary.results[1, , "trait.pop.variance"], xlab="Year",
    ylab="Bandwidth Variance (Hz)", type="n", xaxt="n",
    ylim=c(min(SongEvo1$summary.results[, , "trait.pop.variance"], na.rm=TRUE),
    max(SongEvo1$summary.results[, , "trait.pop.variance"], na.rm=TRUE)))
axis(side=1, at=seq(0, 40, by=5), labels=seq(1970, 2010, by=5))
    for(p in 1:iteration){
        lines(SongEvo1$summary.results[p, , "trait.pop.variance"], col="light gray")
        }
n.mean <- apply(SongEvo1$summary.results[, , "trait.pop.variance"], 2, mean, na.rm=TRUE)
lines(n.mean, col="green")

#Plot 95% quantiles
quant.means <- apply (SongEvo1$summary.results[, , "trait.pop.variance"], MARGIN=2, quantile,
        probs=c(0.975, 0.025), R=600, na.rm=TRUE)
lines(quant.means[1,], col="green", lty=2)
lines(quant.means[2,], col="green", lty=2)</pre>
```



Maps

The simulation results include geographical coordinates and are in a standard spatial data format, thus allowing calculation of a wide variety of spatial statistics.

Load packages for making maps.

```
library("sp")
library("reshape2")
library("lattice")
```

Convert data frame from long to wide format. This is necessary for making a multi-panel plot.

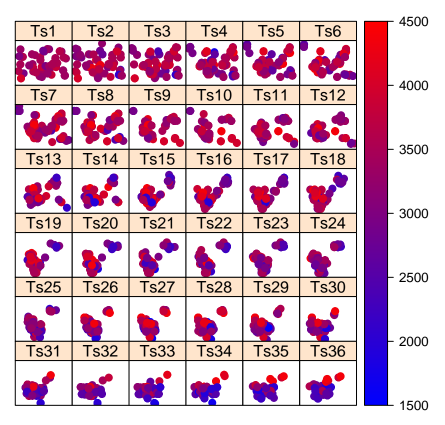
```
all.inds1 <- subset(SongEvo1$all.inds, iteration==1)
w <- dcast(as.data.frame(all.inds1), id ~ timestep, value.var="trait", fill=0)
all.inds1w <- merge(all.inds1, w, by="id")
names(all.inds1w) <- c(names(all.inds1), paste("Ts", seq(1:years), sep=""))</pre>
```

Create a function to generate a continuous color palette—we will use the palette in the next call to make color ramp to represent the trait value.

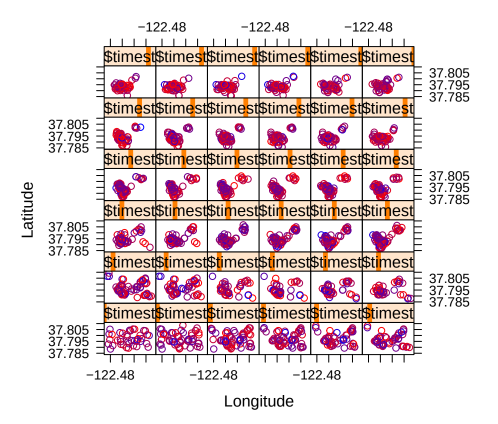
```
rbPal <- colorRampPalette(c('blue', 'red')) #Create a function to generate a continuous color palette
```

Plot maps, including a separate panel for each timestep (each of 36 years). Our example shows that individuals move across the landscape and that regional dialects evolve and move. The x-axis is longitude, the y-axis is latitude, and the color ramp indicates trill bandwidth in Hz.

```
spplot(all.inds1w[,-c(1:ncol(all.inds1))], as.table=TRUE,
    cuts=c(0, seq(from=1500, to=4500, by=10)), ylab="",
    col.regions=c("transparent", rbPal(1000)),
    #cuts specifies that the first level (e.g. <1500) is transparent.
colorkey=list(
   right=list(
          fun=draw.colorkey,
          args=list(
                key=list(
                at=seq(1500, 4500, 10),
                col=rbPal(1000),
                labels=list(
                at=c(1500, 2000, 2500, 3000, 3500, 4000, 4500),
                labels=c("1500", "2000", "2500", "3000", "3500", "4000", "4500")
                )
                )
                )
            )
    )
```



In addition, you can plot simpler multi-panel maps that do not take advantage of the spatial data class.



Test model sensitivity with par.sens()

This function allows testing the sensitivity of SongEvo to different parameter values.

Specify and call par.sens()

Here we test the sensitivity of the Acquire a Territory submodel to variation in territory turnover rates, ranging from 0.8–1.2 times the published rate (40–60% of territories turned over). The call for the par.sens function has a format similar to SongEvo. The user specifies the parameter to test and the range of values for that parameter. The function currently allows examination of only one parameter at a time and requires at least two iterations.

```
parm <- "terr.turnover"
par.range = seq(from=0.4, to=0.6, by=0.05)
sens.results <- NULL</pre>
```

Now we call the par.sens function with our specifications.

Examine par.sens results

Examine results objects, which include two arrays:

The first array, sens.results, contains the SongEvo model results for each parameter. It has the following dimensions:

```
dimnames(par.sens1$sens.results)
#> [[1]]
#> [1] "iteration 1" "iteration 2" "iteration 3" "iteration 4"
#> [5] "iteration 5" "iteration 6" "iteration 7" "iteration 8"
#> [9] "iteration 9" "iteration 10"
#> [[2]]
#> [1] "1" "2" "3" "4" "5" "6" "7" "8" "9" "10" "11" "12" "13" "14"
#> [15] "15" "16" "17" "18" "19" "20" "21" "22" "23" "24" "25" "26" "27" "28"
#> [29] "29" "30" "31" "32" "33" "34" "35" "36"
#>
#> [[31]
#> [1] "sample.n"
                          "trait.pop.mean"
                                              "trait.pop.variance"
#> [4] "lci"
                           "uci"
#>
#> [[4]]
#> [1] "par.val 0.4" "par.val 0.45" "par.val 0.5" "par.val 0.55"
#> [5] "par.val 0.6"
```

The second array, sens.results.diff contains the quantile range of trait values across iterations within a parameter value. It has the following dimensions:

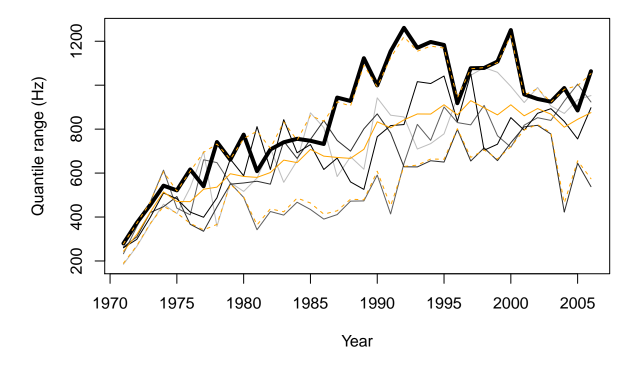
```
dimnames(par.sens1$sens.results.diff)
#> [[1]]
#> [1] "par.val 0.4" "par.val 0.45" "par.val 0.5" "par.val 0.55"
#> [5] "par.val 0.6"
#>
#> [[2]]
#> [1] "Quantile diff 1" "Quantile diff 2" "Quantile diff 3"
#> [4] "Quantile diff 4" "Quantile diff 5" "Quantile diff 6"
#> [7] "Quantile diff 7" "Quantile diff 8" "Quantile diff 9"
#> [10] "Quantile diff 10" "Quantile diff 11" "Quantile diff 12"
#> [13] "Quantile diff 13" "Quantile diff 14" "Quantile diff 15"
#> [16] "Quantile diff 16" "Quantile diff 17" "Quantile diff 18"
#> [19] "Quantile diff 19" "Quantile diff 20" "Quantile diff 21"
#> [22] "Quantile diff 22" "Quantile diff 23" "Quantile diff 24"
#> [25] "Quantile diff 28" "Quantile diff 29" "Quantile diff 27"
#> [28] "Quantile diff 28" "Quantile diff 29" "Quantile diff 30"
```

```
#> [31] "Quantile diff 31" "Quantile diff 32" "Quantile diff 33"
#> [34] "Quantile diff 34" "Quantile diff 35" "Quantile diff 36"
```

To assess sensitivity of SongEvo to a range of parameter values, plot the range in trait quantiles per year by the parameter value. We see that territory turnover values of 0.4–0.6 provided means and quantile ranges of trill bandwidths that are similar to those obtained with the published estimate of 0.5, indicating that the Acquire a Territory submodel is robust to realistic variation in those parameter values.

In the figure, solid gray and black lines show the quantile range of song frequency per year over all iterations as parameterized with the published territory turnover rate (0.5; thick black line) and a range of values from 0.4 to 0.6 (in steps of 0.05, light to dark gray). Orange lines show the mean and 2.5th and 97.5th quantiles of all quantile ranges.

```
#plot of range in trait quantiles by year by parameter value
plot(1:years, par.sens1$sens.results.diff[1,], ylim=c(min(par.sens1$sens.results.diff,
    na.rm=TRUE), max(par.sens1$sens.results.diff, na.rm=TRUE)), type="1",
   ylab="Quantile range (Hz)", xlab="Year", col="transparent", xaxt="n")
axis(side=1, at=seq(0, 35, by=5), labels=seq(1970, 2005, by=5))
  #Make a continuous color ramp from gray to black
grbkPal <- colorRampPalette(c('gray', 'black'))</pre>
  #Plot a line for each parameter value
for(i in 1:length(par.range)){
lines(1:years, par.sens1$sens.results.diff[i,], type="1",
    col=grbkPal(length(par.range))[i])
}
  #Plot values from published parameter values
lines(1:years, par.sens1$sens.results.diff[2,], type="1", col="black", lwd=4)
  #Calculate and plot mean and quantiles
quant.mean <- apply(par.sens1$sens.results.diff, 2, mean, na.rm=TRUE)
lines(quant.mean, col="orange")
#Plot 95% quantiles (which are similar to credible intervals)
  #95% quantiles of population means (narrower)
quant.means <- apply (par.sens1$sens.results.diff, MARGIN=2, quantile,
    probs=c(0.975, 0.025), R=600, na.rm=TRUE)
lines(quant.means[1,], col="orange", lty=2)
lines(quant.means[2,], col="orange", lty=2)
```



Optimize parameter values with par.opt()

This function follows par.sens to help users optimize values for imperfectly known parameters for SongEvo. The goals are to maximize accuracy and precision of model prediction. Accuracy is quantified by three different approaches: i) the mean of absolute residuals of the predicted population mean values in relation to target data (e.g. observed or hypothetical values (smaller absolute residuals indicate a more accurate model)), ii) the difference between the bootstrapped mean of predicted population means and the mean of the target data, and iii) the proportion of simulated population trait means that fall within (i.e. are "contained by") the confidence intervals of the target data (a higher proportion indicates greater accuracy). Precision is measured with the residuals of the predicted population variance to the variance of target data (smaller residuals indicate a more precise model).

Prepare current song values

```
target.data <- subset(song.data, Population=="PRBO" & Year==2005)$Trill.FBW
```

Specify and call par.opt()

Users specify the timestep ("ts") at which to compare simulated trait values to target trait data ("target.data") and save the results in an object (called par.opt1 here).

Examine results objects (residuals and target match).

```
par.opt1$Residuals
#> , , Residuals of mean
#>

#>

Iteration 1 Iteration 2 Iteration 3 Iteration 4 Iteration 5
#> par.val 0.4 383.6657 47.60517 47.43456 324.85793 734.2031
```

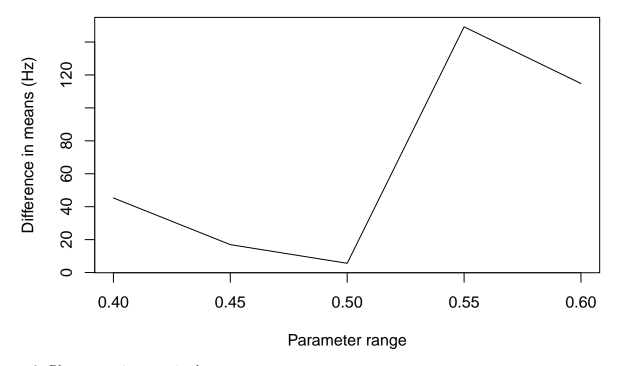
```
#> par.val 0.45
                  419.2283
                             406.21328 136.03468
                                                    232.35778
                                                                 334.1901
#> par.val 0.5
                             548.67152 233.99923
                  123.0665
                                                    271.87325
                                                                 142.8514
#> par.val 0.55
                  143.4617
                             225.08001
                                         28.93880
                                                     65.91219
                                                                 544.6026
#> par.val 0.6
                  116.8529
                              28.26789 117.15954
                                                    695.56947
                                                                 168.6774
               Iteration 6 Iteration 7 Iteration 8 Iteration 9 Iteration 10
#>
#> par.val 0.4
                  504.3934
                              219.9672
                                         64.62456
                                                     254.0140
                                                                 90.14688
#> par.val 0.45
                  503.6697
                             448.5809
                                         674.34969
                                                     504.3921
                                                                 301.89841
                                                                 255.21635
#> par.val 0.5
                  152.7058 254.2320
                                        716.48343
                                                     209.9715
#> par.val 0.55
                  254.9533
                             201.7253
                                         122.62141
                                                     193.9677
                                                                 172.28947
                                         599.29129
                                                     307.9787
#> par.val 0.6
                  182.9460
                              236.0133
                                                                  61.08874
#>
#> , , Residuals of variance
#>
#>
               Iteration 1 Iteration 2 Iteration 3 Iteration 4 Iteration 5
                             535479.5
                                         171520.5
                                                     578709.1
#> par.val 0.4
                  175754.1
                                                                 244465.4
#> par.val 0.45
                  286613.8
                             306324.9
                                         953998.4
                                                     324503.1
                                                                 253660.0
#> par.val 0.5
                  513340.8
                              478060.7
                                         617687.0
                                                     481771.9
                                                                 763181.2
#> par.val 0.55
                  854512.5
                              300257.1
                                         585129.9
                                                     526303.9
                                                                 583339.0
#> par.val 0.6
                  322840.6
                              898194.7
                                                     286737.2 1089690.1
                                         223242.3
#>
               Iteration 6 Iteration 7 Iteration 8 Iteration 9 Iteration 10
#> par.val 0.4
                  256209.3
                             384433.7
                                         542116.7
                                                     384384.8
                                                                  408530.5
#> par.val 0.45
                  257798.1
                             578514.4
                                         209509.2
                                                     722086.2
                                                                  762550.0
#> par.val 0.5
                                                                  478619.3
                  930202.0 647553.9
                                         110073.1
                                                     279279.7
#> par.val 0.55
                  416482.2
                              888578.1
                                         457855.7
                                                     296017.6
                                                                  717650.0
#> par.val 0.6
                              335369.9
                                         181271.5
                  554666.7
                                                     396157.8
                                                                  604308.1
par.opt1$Target.match
#>
             Difference in means Proportion contained
#> par.val 0.4
                        45.302898
                                                   0.4
#> par.val 0.45
                         16.959666
                                                   0.0
#> par.val 0.5
                                                   0.0
                         5.632687
#> par.val 0.55
                        149.222433
                                                   0.2
#> par.val 0.6
                        114.811394
                                                   0.2
```

Plot results of par.opt()

Accuracy

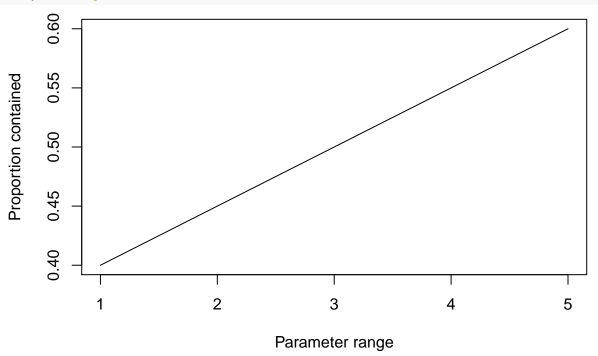
1. Difference in means.

```
plot(par.range, par.opt1$Target.match[,1], type="l", xlab="Parameter range",
    ylab="Difference in means (Hz)")
```



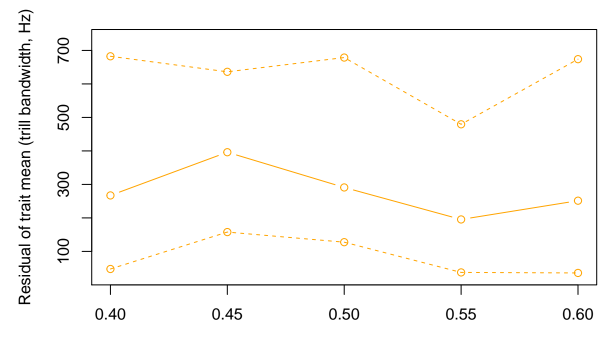
2. Plot proportion contained.

```
plot(par.range, par.opt1$Prop.contained, type="1", xlab="Parameter range",
    ylab="Proportion contained")
```



3. Calculate and plot mean and quantiles of residuals of mean trait values.

```
xlab="Parameter value (territory turnover rate)",
   ylab="Residual of trait mean (trill bandwidth, Hz)")
points(par.range, res.mean.quants[1,], col="orange")
points(par.range, res.mean.quants[2,], col="orange")
lines(par.range, res.mean.quants[1,], col="orange", lty=2)
lines(par.range, res.mean.quants[2,], col="orange", lty=2)
```



Precision

Parameter value (territory turnover rate)



Visual inspection of accuracy and precision: plot trait values for range of parameters

```
par(mfcol=c(3,2),
   mar=c(2.1, 2.1, 0.1, 0.1),
    cex=0.8)
for(i in 1:length(par.range)){
plot(par.sens1$sens.results[ , , "trait.pop.mean", ], xlab="Year", ylab="Bandwidth (Hz)",
    xaxt="n", type="n", xlim=c(-0.5, years),
   ylim=c(min(par.sens1$sens.results[ , , "trait.pop.mean", ], na.rm=TRUE),
   max(par.sens1$sens.results[ , , "trait.pop.mean", ], na.rm=TRUE)))
   for(p in 1:iteration){
        lines(par.sens1$sens.results[p, , "trait.pop.mean", i], col="light gray")
freq.mean <- apply(par.sens1$sens.results[, , "trait.pop.mean", i], 2, mean, na.rm=TRUE)</pre>
lines(freq.mean, col="blue")
axis(side=1, at=seq(0, 35, by=5), labels=seq(1970, 2005, by=5))
#Plot 95% quantiles
quant.means <- apply (par.sens1$sens.results[, , "trait.pop.mean", i], MARGIN=2, quantile,
   probs=c(0.95, 0.05), R=600, na.rm=TRUE)
lines(quant.means[1,], col="blue", lty=2)
lines(quant.means[2,], col="blue", lty=2)
#plot mean and CI for historic songs.
#plot original song values
library("boot")
sample.mean <- function(d, x) {</pre>
   mean(d[x])
boot_hist <- boot(starting.trait, statistic=sample.mean, R=100)#, strata=mn.res$iteration)
```

```
ci.hist <- boot.ci(boot_hist, conf=0.95, type="basic")</pre>
low <- ci.hist$basic[4]</pre>
high <- ci.hist$basic[5]</pre>
points(0, mean(starting.trait), pch=20, cex=0.6, col="black")
library("Hmisc")
errbar(x=0, y=mean(starting.trait), high, low, add=TRUE)
 #plot current song values
library("boot")
sample.mean <- function(d, x) {</pre>
    mean(d[x])
}
boot_curr <- boot(target.data, statistic=sample.mean, R=100)#, strata=mm.res$iteration)
ci.curr <- boot.ci(boot_curr, conf=0.95, type="basic")</pre>
low <- ci.curr$basic[4]</pre>
high <- ci.curr$basic[5]</pre>
points(years, mean(target.data), pch=20, cex=0.6, col="black")
library("Hmisc")
errbar(x=years, y=mean(target.data), high, low, add=TRUE)
  #plot panel title
text(x=3, y=max(par.sens1$sens.results[ , , "trait.pop.mean", ], na.rm=TRUE)-100,
    labels=paste("Par = ", par.range[i], sep=""))
}
     Par = 0.4
                                                      Par = 0.55
3500
                                                 3500
2500
                                                 2500
    1970
               1980
                          1990
                                      2000
                                                      1970
                                                                 1980
                                                                            1990
                                                                                       2000
     Par = 0.45
                                                       Par = 0.6
                                                 3500
3500
2500
                                                 2500
    1970
               1980
                          1990
                                      2000
                                                      1970
                                                                 1980
                                                                            1990
                                                                                       2000
      Par = 0.5
3500
2500
    1970
               1980
                          1990
                                      2000
```

Model validation with mod.val()

This function allows users to assess the validity of the specified model by testing model performance with a population different from the population used to build the model. The user first runs SongEvo with initial trait values from the validation population. mod.val() uses the summary.results array from SongEvo, along with target values from a specified timestep, to calculate the same three measures of accuracy and one measure of precision that are calculated in par.opt.

We parameterized SongEvo with initial song data from Schooner Bay, CA in 1969, and then compared simulated data to target (i.e. observed) data in 2005.

Prepare initial song data for Schooner Bay.

Specify and call SongEvo() with validation data

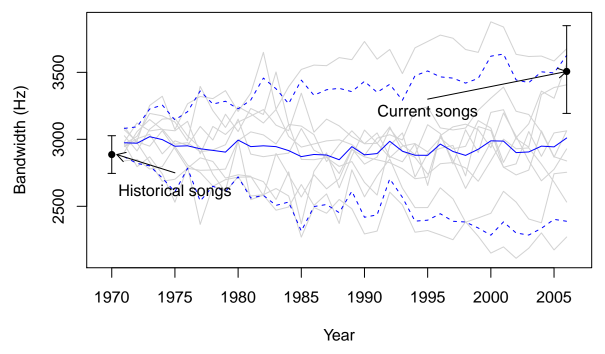
```
iteration <- 10
years <- 36
timestep <- 1
terr.turnover <- 0.5

SongEvo2 <- SongEvo(init.inds = init.inds, iteration = iteration, steps = years,
    timestep = timestep, n.territories = n.territories, terr.turnover = terr.turnover,
    learning.method = learning.method, integrate.dist = integrate.dist,
    learning.error.d = learning.error.d, learning.error.sd = learning.error.sd,
    mortality.a = mortality.a, mortality.j = mortality.j, lifespan = lifespan,
    phys.lim.min = phys.lim.min, phys.lim.max = phys.lim.max,
    male.fledge.n.mean = male.fledge.n.mean, male.fledge.n.sd = male.fledge.n.sd,
    male.fledge.n = male.fledge.n, disp.age = disp.age,
    disp.distance.mean = disp.distance.mean, disp.distance.sd = disp.distance.sd,
    mate.comp = mate.comp, prin = prin, all)</pre>
```

Specify and call mod.val

Plot results from mod.val()

```
#Plot 95% quantiles
quant.means <- apply (SongEvo2\summary.results[, , "trait.pop.mean"], MARGIN=2, quantile,
    probs=c(0.95, 0.05), R=600, na.rm=TRUE)
lines(quant.means[1,], col="blue", lty=2)
lines(quant.means[2,], col="blue", lty=2)
#plot mean and CI for historic songs.
#plot original song values
library("boot")
sample.mean <- function(d, x) {</pre>
    mean(d[x])
boot_hist <- boot(starting.trait, statistic=sample.mean, R=100)</pre>
ci.hist <- boot.ci(boot_hist, conf=0.95, type="basic")</pre>
low <- ci.hist$basic[4]</pre>
high <- ci.hist$basic[5]</pre>
points(0, mean(starting.trait), pch=20, cex=0.6, col="black")
library("Hmisc")
errbar(x=0, y=mean(starting.trait), high, low, add=TRUE)
#text and arrows
text(x=5, y=2720, labels="Historical songs", pos=1)
arrows(x0=5, y0=2750, x1=0.4, y1=mean(starting.trait), length=0.1)
#plot current song values
library("boot")
sample.mean <- function(d, x) {</pre>
    mean(d[x])
boot_curr <- boot(target.data, statistic=sample.mean, R=100)</pre>
ci.curr <- boot.ci(boot_curr, conf=0.95, type="basic")</pre>
low <- ci.curr$basic[4]</pre>
high <- ci.curr$basic[5]</pre>
points(years, mean(target.data), pch=20, cex=0.6, col="black")
library("Hmisc")
errbar(x=years, y=mean(target.data), high, low, add=TRUE)
#text and arrows
text(x=25, y=3100, labels="Current songs", pos=3)
arrows(x0=25, y0=3300, x1=36, y1=mean(target.data), length=0.1)
```



The model did reasonably well predicting trait evolution in the validation population, suggesting that it is valid for our purposes: the mean bandwidth was abs(mean(target.data)-freq.mean)Hz from the observed values, ~21% of predicted population means fell within the 95% confidence intervals of the observed data, and residuals of means (~545 Hz) and variances (~415181 Hz) were similar to those produced by the training data set.

Hypothesis testing with h.test()

This function allows hypothesis testing with SongEvo. To test if measured songs from two time points evolved through mechanisms described in the model (e.g. drift or selection), users initialize the model with historical data, parameterize the model based on their understanding of the mechanisms, and test if subsequently observed or predicted data match the simulated data. The output data list includes two measures of accuracy: the proportion of observed points that fall within the confidence intervals of the simulated data and the residuals between simulated and observed population trait means. Precision is measured as the residuals between simulated and observed population trait variances. We tested the hypothesis that songs of Z. l. nuttalli in Bear Valley, CA evolved through cultural drift from 1969 to 2005.

Prepare initial song data for Bear Valley.

Specify and call SongEvo() with test data

```
SongEvo3 <- SongEvo(init.inds = init.inds, iteration = iteration, steps = years,
    timestep = timestep, n.territories = n.territories, terr.turnover = terr.turnover,
    learning.method = learning.method, integrate.dist = integrate.dist,
    learning.error.d = learning.error.d, learning.error.sd = learning.error.sd,</pre>
```

```
mortality.a = mortality.a, mortality.j = mortality.j, lifespan = lifespan,
phys.lim.min = phys.lim.min, phys.lim.max = phys.lim.max,
male.fledge.n.mean = male.fledge.n.mean, male.fledge.n.sd = male.fledge.n.sd,
male.fledge.n = male.fledge.n, disp.age = disp.age,
disp.distance.mean = disp.distance.mean, disp.distance.sd = disp.distance.sd,
mate.comp = mate.comp, prin = prin, all)
```

Specify and call h.test()

The output data list includes two measures of accuracy: the proportion of observed points that fall within the confidence intervals of the simulated data and the residuals between simulated and observed population trait means. Precision is measured as the residuals between simulated and observed population trait variances.

Eighty percent of the observed data fell within the central 95% of the simulated values, providing support for the hypothesis that cultural drift as described in this model is sufficient to describe the evolution of trill frequency bandwidth in this population.

```
h.test1
#> $Residuals
               Residuals of mean Residuals of variance
#>
#> Iteration 1
                       291.5200
                                              340238.2
#> Iteration 2
                        192.8924
                                              582643.8
#> Iteration 3
                       961.2465
                                              152831.5
#> Iteration 4
                        159.1739
                                              480857.1
                        572.3262
#> Iteration 5
                                              445170.6
#> Iteration 6
                       221.5838
                                              443112.3
#> Iteration 7
                       138.7057
                                              300428.8
#> Iteration 8
                                              813656.6
                        429.0511
#> Iteration 9
                                              224839.5
                        775.2170
#> Iteration 10
                        661.0336
                                              138152.7
#> $Prop.contained
#> [1] 0.8
```

We can plot simulated data in relation to measured data.

```
lines(quant.means[2,], col="blue", lty=2)
 #plot original song values
library("boot")
sample.mean <- function(d, x) {</pre>
    mean(d[x])
boot hist <- boot(starting.trait, statistic=sample.mean, R=100)#, strata=mn.res$iteration)
ci.hist <- boot.ci(boot_hist, conf=0.95, type="basic")</pre>
low <- ci.hist$basic[4]</pre>
high <- ci.hist$basic[5]</pre>
points(0, mean(starting.trait), pch=20, cex=0.6, col="black")
library("Hmisc")
errbar(x=0, y=mean(starting.trait), high, low, add=TRUE)
#plot current song values
points(rep(ts, length(target.data)), target.data)
library("boot")
sample.mean <- function(d, x) {</pre>
    mean(d[x])
boot_curr <- boot(target.data, statistic=sample.mean, R=100)#, strata=mm.res$iteration)
ci.curr <- boot.ci(boot_curr, conf=0.95, type="basic")</pre>
low <- ci.curr$basic[4]</pre>
high <- ci.curr$basic[5]</pre>
points(years, mean(target.data), pch=20, cex=0.6, col="black")
library("Hmisc")
errbar(x=years, y=mean(target.data), high, low, add=TRUE)
#text and arrows
text(x=11, y=2850, labels="Historical songs", pos=1)
arrows(x0=5, y0=2750, x1=0.4, y1=mean(starting.trait), length=0.1)
text(x=25, y=2900, labels="Current songs", pos=1)
arrows(x0=25, y0=2920, x1=years, y1=mean(target.data), length=0.1)
```

