Demographic Modeling Introduction

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Demographic models and population viability analysis (PVA)

Calculating a population growth rate

We will use census data from a multi-year bottlenose dolphin census to calculate annual growth rates. NOTE: These data are made up for the purposes of this demonstration, and do not represent real population sizes for this species.

#bottlenose_census <- read.csv(file = "https://raw.githubusercontent.com/mlammens/DemModel-ShortCourse/bottlenose_census <- read.csv(file = "~/Google Drive/Professional/Demographic-Modeling-Short-Course/bot

Let's have a quick look at these data

head(bottlenose_census)

```
## year pop_size
## 1 1990 1000
## 2 1991 996
## 3 1992 1185
## 4 1993 1015
## 5 1994 923
## 6 1995 1120
```

tail(bottlenose_census)

summary(bottlenose_census)

```
##
         year
                      pop_size
##
   Min.
           :1990
                   Min. : 923
   1st Qu.:1996
                   1st Qu.:1130
  Median :2002
                   Median:1468
           :2002
##
  Mean
                          :1740
                   Mean
##
   3rd Qu.:2009
                   3rd Qu.:1950
##
   Max.
           :2015
                          :3615
                   Max.
```

Since we have 26 years worth of data, we should be able to calculate 25 R values. Recall the formula for R is:

$$R = \frac{N(t+1)}{N(t)}$$

```
bottlenose_R <- bottlenose_census$pop_size[2:length(bottlenose_census$pop_size)] /
bottlenose_census$pop_size[1:(length(bottlenose_census$pop_size) - 1)]</pre>
```

Let's look at those values by printing them out to the screen and looking at a few summaries.

bottlenose_R

```
## [1] 0.9960000 1.1897590 0.8565401 0.9093596 1.2134345 1.1017857 0.8176661 
## [8] 1.0931615 1.0525839 1.0680448 1.2862903 0.9028213 1.0993056 0.9456728 
## [15] 1.1349365 1.1571513 0.6846389 1.3268945 1.0643897 1.2288269 1.1712329 
## [22] 1.0500731 1.2464323 1.0094946 0.9106501
```

```
min(bottlenose_R)

## [1] 0.6846389

max(bottlenose_R)

## [1] 1.326895

Geometric mean of R

prod(bottlenose_R)^(1/length(bottlenose_R))

## [1] 1.048814

Standard deviation of R

sd(bottlenose_R)

## [1] 0.1565993

Standard error of the mean

sd(bottlenose_R)/sqrt(length(bottlenose_R))
```

Simulating population growth

OK. Let's now use this information to simulate the population change for bottlenose dolphins, assuming the past population growth rates are a reasonable representation of future population conditions.

We will use two different methods to do this.

Bootstrap method

[1] 0.03131986

First, let's use a bootstrapping method, where we restrict our selves to using only observed R values.

Assume that we want to forecast our population for 25 years. Then we need to draw 25 R values from our current set. As this is bootstrapping for forecasting, we want to sample **with** replacement.

```
bottlenose_R_proj <- sample(bottlenose_R, size = 25, replace = TRUE)</pre>
```

Let's look at our new values.

```
bottlenose_R_proj
```

```
## [1] 1.2134345 0.6846389 0.6846389 1.0500731 1.3268945 1.2862903 1.0680448
## [8] 1.1897590 0.9456728 1.0993056 1.1712329 1.2862903 1.1571513 1.0094946
## [15] 1.1571513 0.6846389 1.0094946 1.0993056 1.1712329 0.8176661 1.0094946
## [22] 1.0500731 1.0643897 0.6846389 1.2862903
```

```
min(bottlenose_R_proj)

## [1] 0.6846389

max(bottlenose_R_proj)

## [1] 1.326895

Geometric mean

prod(bottlenose_R_proj)^(1/length(bottlenose_R_proj))
```

```
## [1] 1.027755
```

In order to use these values to forecast the population, we're going to need to use a for loop.

```
# Set our initial value
n_init <- 1000

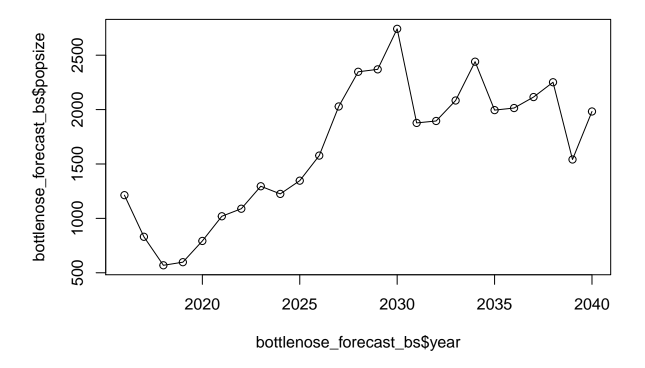
# Create a vector to store new values in
n <- vector(length = length(bottlenose_R_proj))

n_old <- n_init
for (t in 1:length(bottlenose_R_proj)){
    n_new <- n_old * bottlenose_R_proj[t]
    n[t] <- n_new
    n_old <- n_new
}</pre>
```

Make a data.frame with these new values

Plot our values through time

```
plot(x = bottlenose_forecast_bs$year, y = bottlenose_forecast_bs$popsize, type = "o")
```



Calculating extinction risk and other measures

The work above gave us one possible trajectory for this population, but in order to really say something about what we might expect in the future, we need to run this simulation 100s or 1000s of times, and use those values to make probabilistic statements.

Let's run 1000 simulations.

First generate 1000 * 25 R values.

```
bottlenose_R_proj_multi <- sample(bottlenose_R, size = (1000*25), replace = TRUE)
```

Make these into a 2D matrix

```
bottlenose_R_proj_multi <- matrix(bottlenose_R_proj_multi, nrow = 25, byrow = FALSE)
```

In order to run these simulations 1000 times, we're going to use a for loop inside another for loop.

Let's start by making an empty matrix to store all of our simulation results. This matrix needs to have 25 rows and 1000 columns.

```
bottlenose_forecast_bs_multi <- matrix(rep(NA, 1000*25), nrow = 25 )</pre>
```

Why did I use NAs here?

Let's make that double for loop.

```
# Set our initial value
n_init <- 1000
n_sims <- ncol(bottlenose_forecast_bs_multi)</pre>
```

```
for( sim in 1:n_sims){

# Create a vector to store new values in

n <- vector(length = nrow(bottlenose_R_proj_multi))

n_old <- n_init
for (t in 1:nrow(bottlenose_R_proj_multi)){
    n_new <- n_old * bottlenose_R_proj_multi[t, sim]
    n[t] <- n_new
    n_old <- n_new
}

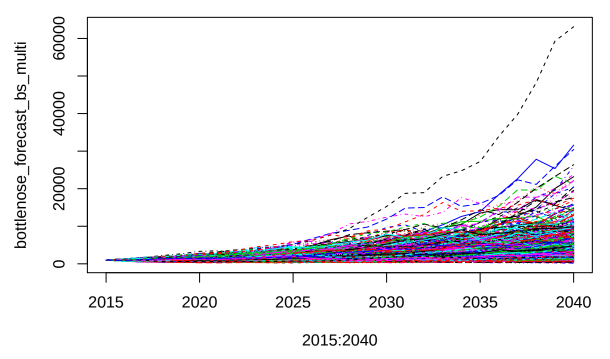
bottlenose_forecast_bs_multi[ , sim] <- n
}</pre>
```

Add the initial value as a new **first** row.

```
bottlenose_forecast_bs_multi <- rbind(rep(n_init, n_sims), bottlenose_forecast_bs_multi)</pre>
```

Let's plot all of the simulations.

```
matplot(x = 2015:2040, y = bottlenose_forecast_bs_multi, type = "1")
```



We can determine the number of simulations where the population goes extinct by looking for how many populations are 0 individuals after 25 years.

```
sum(bottlenose_forecast_bs_multi[26, ] == 0)
```

[1] 0

How about less than some threshold value? Let's say 500.

```
sum(bottlenose_forecast_bs_multi[26, ] < 700)</pre>
```

```
## [1] 26
```

We can use these values to calculate the probability of this event happening. In this case, the probability of decline to below 700 individuals is:

```
sum(bottlenose_forecast_bs_multi[26, ] < 700) / n_sims</pre>
```

```
## [1] 0.026
```

Another useful statistic is the **Expected Minimum Abundance**. Calculating this value will give us a chance to become familiar with a few more R functions.

In order to calculate the EMA, we first need to calculate the minimum value for each simulation.

```
bottlenose_forecast_min <- apply(X = bottlenose_forecast_bs_multi, MARGIN = 2, FUN = min)</pre>
```

Next we can calculate the average minimum value.

```
mean(bottlenose_forecast_min)
```

```
## [1] 859.9625
```

Challange

- 1. What happens if the population size is much smaller? Say 100 individuals?
- 2. What happens if boat tours lead to a 0.025 reduction in population growth rate?

Using statistical distribution methods

Instead of bootstrapping, we could parameterize a statistical distribution with the data we've collected.

```
bottlenose_R_proj <- rlnorm(25, meanlog = log(prod(bottlenose_R)^(1/length(bottlenose_R))), sdlog = sd(
```

Repeat steps as above

In order to use these values to forecast the population, we're going to need to use a for loop.

```
# Set our initial value
n_init <- 1000

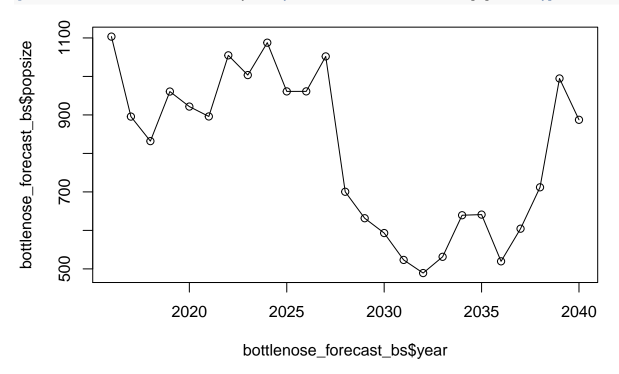
# Create a vector to store new values in
n <- vector(length = length(bottlenose_R_proj))

n_old <- n_init
for (t in 1:length(bottlenose_R_proj)){
    n_new <- n_old * bottlenose_R_proj[t]
    n[t] <- n_new
    n_old <- n_new
}</pre>
```

Make a data.frame with these new values

Plot our values through time

```
plot(x = bottlenose_forecast_bs$year, y = bottlenose_forecast_bs$popsize, type = "o")
```



Caveats regarding PVA

- Be careful
- Only as good as data
- "All models are wrong, but some are useful" George Box

Matrix projection models

Let's parameterize a projection matrix.

```
bottlenose_mat <- matrix(c(0.3, 1, 0.2, 0.8), nrow = 2, byrow = TRUE) eigen(bottlenose_mat)
```

```
## $values
## [1] 1.06234754 0.03765246
##
## $vectors
## [,1] [,2]
## [1,] -0.7952619 -0.9672672
## [2,] -0.6062660 0.2537602
```

Set an initial population size - but now it must be separated by age/stage.

```
n_init_stage <- c(500,500)</pre>
```

Get the abundance for each stage at the next time-step.

```
bottlenose_mat %*% n_init_stage
```

```
## [,1]
## [1,] 650
## [2,] 500
```

Challenge

Write a for loop to forecast the population size changes of the bottlenose dolphin population 25 years into the future.

Considering other sources of variability

Environmental

Similar to how we did it above, but sample for each parameter, rather than R.

Demographic

Nature happens in whole numbers.

If we have estimates of variability for reproduction, we can use a Poisson distribution to determine the number of births.

We can use binomial distribution to determine the number of deaths.