Week 13, Lecture 5.3 Exercise

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Exercise 1. Our First ABC - estimating parameters of Gaussian distribution given observed data

A. Normal distribution You took some introductory statistics, so I hope you are familiar with a Gaussian distribution (also referred to as a normal distribution). It's useful to try and attribute data to an underlying probability distribution, so we can understand and make inference about that data. For example, many traits that have a genetic basis follow a normal distribution. Body length in animals is a good example of this. Researchers in the early 1900s were quite interested in better understanding genetic inheritance of traits of interest for animal breeding, which is a key reason why population geneticists (RA Fisher) were the ones who developed classical statistical tests such as Analysis of Variance - the goal was to understand how evolution led to shifts in normally distributed traits. The normal distribution itself was discovered by Carl Friedrich Gauss (a German mathematician and physicist!!) in 1809.

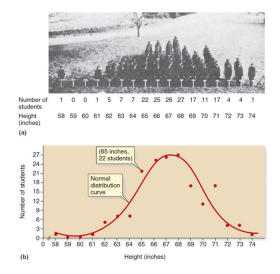


Figure 1: Human height. (a) Height distribution (inches) for 175 students in 1914 attending the Connecticut Agricultural College. (b) Graphical presentation of these student heights showing their close fit to a normal distribution. (a: Reprinted with permission from Albert and Blakeslee: Corn and Man. Journal of Heredity. 1914;5:51. Oxford University Press. b: Reprinted with permission from Brooker RJ: Genetics: Analysis & Principles, 3rd ed. New York: McGrawHill, 2008.)

We can plot data and use our own eyes to consider whether it follows a normal distribution. But I would like you to have an awareness that even claiming / stating "The data x follows a normal distribution" is -you guessed it - formulating a model! So then, given a set of data, we can propose a model, and estimate the parameter values when fitting that data to the model. We will do this with a normal distribution today.

B. The easiest / typical way to estimate parameters of a normal model when fit to observed data The focus of today's exercise will be to use Approximate Bayesian Computation (ABC) for fitting data to models. ABC is most needed with very complicated models where the underlying parameters are often unknown and difficult to estimate. However, its best to show you how ABC works by applying it to a simpler model. So we will learn to estimate the parameters in a normal distribution given a set of data.

The dataset of interest is body weight of a common household insect. Please run the following command to load the dataset:

```
# Dataset with measures of 1000 household insects body mass
# (g)
dat <- read.csv("https://raw.githubusercontent.com/jhpantel/ude-ecomod/main/data/insect.csv",
    header = TRUE, row.names = 1)</pre>
```

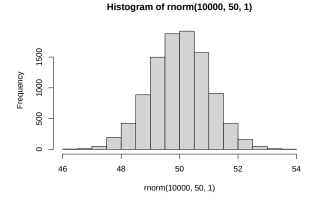
You can visualize and summarize the data - what is the mean (mean) and standard deviation (sd) of the dataset? What is the natural logarithm of the standard deviation of the dataset (log(sd))?

A normal distribution has two parameters to describe its shape - mean (μ) and standard deviation (σ) . The formula to generate a normal curve is:

$$p(x) = \frac{1}{\sigma\sqrt{2\pi}}e^{-\frac{1}{2}(\frac{x-\mu}{\sigma})^2}$$

Where p(x) is the probability of observing a value of x given the model. For example, if data is drawn from a normal distribution centered on a mean mean $(\mu) = 50$ and standard deviation $(\sigma) = 1$, the normal distribution looks like this:

```
hist(rnorm(10000, 50, 1))
```



And the probability of observing a value of 60 given this model and these parameters is quite low:

```
# This calculates p(60) using the formula given above dnorm(60, 50, 1)
```

```
## [1] 7.694599e-23
```

Its of interest to 'fit your data to a normal model'. We can do this very quickly in R - we provide our data dat to a command fitdist in the R library fitdistrplus - it fits your observed data to a normal model and returns estimates of the model parameters (μ and σ):

```
fitdistrplus::fitdist(data = dat$weight_g, distr = "norm")
```

```
## Fitting of the distribution ' norm ' by maximum likelihood
## Parameters:
## estimate Std. Error
## mean 3.4242893 0.004553051
## sd 0.1439801 0.003218794
```

How does the fitdist model estimates for mean (μ) and standard deviation (σ) compare to the true values used to produce the data? Quite well.

C. Estimating parameters of a normal model using ABC (useful when you have more complex models) Let's learn to use ABC (Approximate Bayesian Computation) to estimate the most likely values of the model parameters given the observe data. This is most useful when (1) you have a very complex model and (2) you don't know the underlying values of the parameters that produced the observed data. ABC works differently:

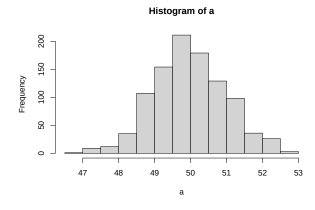
- data is simulated from an underlying model
- summary statistics are calculated from each simulation
- the values of these summary statistics are compared to observed values
- a 'rejection algorithm' is implemented, where simulations with summary statistics that are very far from the observed values are discarded
- the remaining accepted simulations are used to calculate a 'posterior distribution' for underlying model parameters

Let's look at each step more closely:

Step 1. Data is simulated from an underlying model

We stated previously we believe our data is drawn from a normal distribution. We simulate 10000 datasets from a normal model, and we randomly choose parameter values for each simulation from the normal model. We can simulate draws from a normal distribution using the command rnorm

```
# Randomly draw 1000 values from a normal distribution with
# mean 50, standard deviation 1
a <- rnorm(1000, 50, 1)
hist(a)</pre>
```



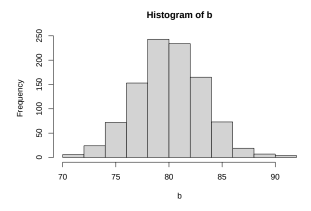
We wish to repeat this process of simulating our insect data (1000 data points) 10,000 times. We do not know the underlying values of the model parameters (μ and σ) that produced our data, so we run our simulations with randomly chosen values for these.

```
mu_rand <- runif(10000, -100, 100) # What is this command doing?
sd_rand <- runif(10000, 0, 20)</pre>
```

Create a histogram (hist) for each of these - do they look as you would expect?

Now for each set of parameters, we simulate data that's meant to mimic our observed insect data using a normal model. How can we do this? Using the rnorm command - we simulate data under a random normal model with parameters μ and σ from each of the 10,000 values we generated above. This works for a single parameter set like this:

```
b <- rnorm(1000, mu_rand[1], sd_rand[1])
hist(b)</pre>
```

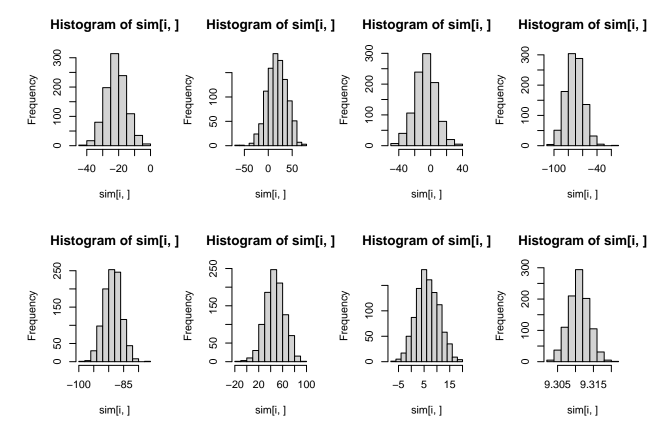


We can do this for all of the parameter sets this way:

```
sim <- array(NA, dim = c(10000, 1000))
for (i in 1:10000) {
    sim[i, ] <- rnorm(1000, mu_rand[i], sd_rand[i])
}</pre>
```

Every single dataset is different:

```
d <- sample.int(10000, 8)
par(mfrow = c(2, 4))
for (i in d) {
    hist(sim[i, ])
}</pre>
```



Step 2. Summary statistics are calculated from each simulation

Our focal summary statistics are (1) the mean of the dataset and (2) the logarithm of the standard deviation of the dataset. We save those to a new variable called stat.obs:

```
stat.obs <- c(mean(dat$weight_g), log(sd(dat$weight_g)))
```

Our simulated summary statistics can be obtained by calculating the mean and log(sd) for each of the 10000 simulations:

Step 3. The values of these summary statistics are compared to observed values (using either a rejection algorithm or a neural network - we use a neural network)

We do this step using the R package abc:

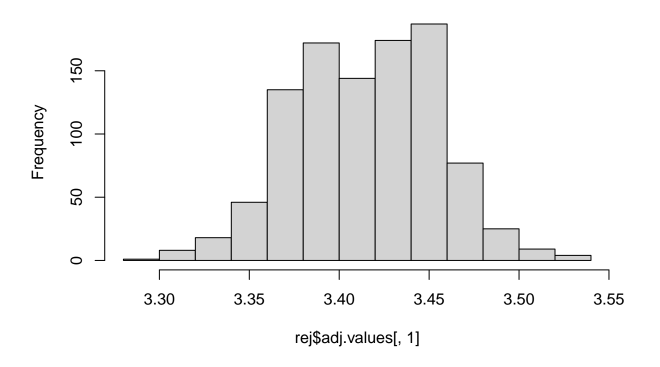
```
## 12345678910
## 12345678910
```

Step 4. The remaining accepted simulations are used to calculate a 'posterior distribution' for underlying model parameters

We can visualize this by plotting the values of the accepted simulations (the ones that are closest to our observed summary statistics):

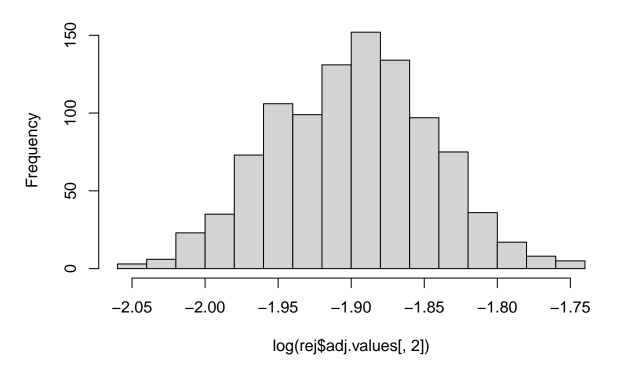
```
par(mfrow = c(1, 1))
hist(rej$adj.values[, 1], main = "posterior distribution of mean")
```

posterior distribution of mean



hist(log(rej\$adj.values[, 2]), main = "posterior distribution of standard deviation")

posterior distribution of standard deviation



How well do these compare to the observed values in the original data (obs.stat)?

D. Estimating parameters of a discrete-time Lotka Volterra model using ABC Let's repeat using ABC (Approximate Bayesian Computation) to estimate the most likely values of the model parameters given the observe data. We can take Gause's *Paramecium* data and try to estimate the underlying parameters from a discrete-time Lotka Volterra model.

Step 1a. Gather your observed data

```
# load competition data
data("gause_1934_science_f02_03")

# subset out data from species grown in mixture
mixturedat <- gause_1934_science_f02_03[gause_1934_science_f02_03$Treatment ==
        "Mixture", ]

# extract time and species data
time <- mixturedat$Day
species <- data.frame(mixturedat$Volume_Species1, mixturedat$Volume_Species2)
colnames(species) <- c("P_caudatum", "P_aurelia")</pre>
```

Step 1b. Arrange your observed data similar to stat.obs above The variable with the observed data, species, has dimensions 23, 2. You should arrange the observed data as a vector of length 46. That means, please arrange the data in species to be 1 vector of length 46. Save that to a new variable called stat.obs Hint: you can try the reshape2::melt command, or you can create your own new variable as stat.obs <- c(species[,1],species[,2])

Make sure your variable is correct by running this command and getting the same results:

```
length(stat.obs)
```

[1] 46

Step 2. Create simulation of data from an underlying model

Our next goal is to use a discrete-time Lotka-Volterra model that can simulate data of the kind observed in Gause's Paramecium. The model should consider the processes - growth and competition - that are important to produce the observed Paramecium time series.

We use the following model:

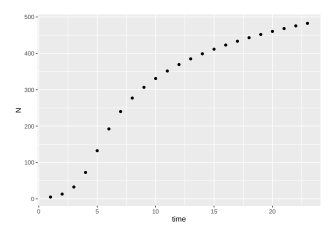
$$\begin{split} N_{1,t+1} &= N_{1,t} + N_{1,t} \cdot (\lambda_1 e^{(-\alpha_{11}N_{1,t} - \alpha_{12}N_{2,t})}) \\ N_{2,t+1} &= N_{2,t} + N_{2,t} \cdot (\lambda_2 e^{(-\alpha_{22}N_{2,t} - \alpha_{21}N_{1,t})}) \end{split}$$

The parameters we need to estimate are:

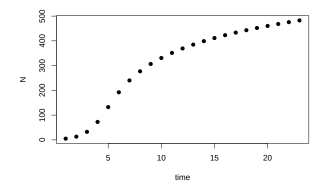
```
\lambda_1, \lambda_2, \alpha_{11}, \alpha_{22}, \alpha_{12}, \alpha_{21}
```

Please create a simulation of this model. I show here a model for a single species. Please modify that to also simulate growth in a second species. For the missing parameters, you can start by using values of $\lambda_2 = 1.5$, $\alpha_{22} = 0.005$, $\alpha_{12} = 0.03$, $\alpha_{21} = 0.007$, $N_{2,0} = 3$. Change the name of the function to disc_lv.

```
# Parameter values to use for simulation
lambda_1 <- 1.7
alpha_11 <- 0.01
N1 0 <- 5
t <- 23
# model function
disc_log <- function(lambda_1, N1_0, alpha_11) {</pre>
    Nt1 <- N1_0 + N1_0 * (lambda_1 * exp(-alpha_11 * N1_0))
    return(Nt1)
# Simulation of model for t time steps
N \leftarrow rep(NA, t)
N[1] <- N1_0
for (i in 2:t) {
    N[i] <- disc_log(lambda_1, N1_0, alpha_11)
    N1_0 \leftarrow N[i]
}
# Plot simulation: ggplot
dat <- as.data.frame(N)</pre>
dat$time <- as.numeric(rownames(dat))</pre>
ggplot2::ggplot(dat, ggplot2::aes(time, N)) + ggplot2::geom_point()
```



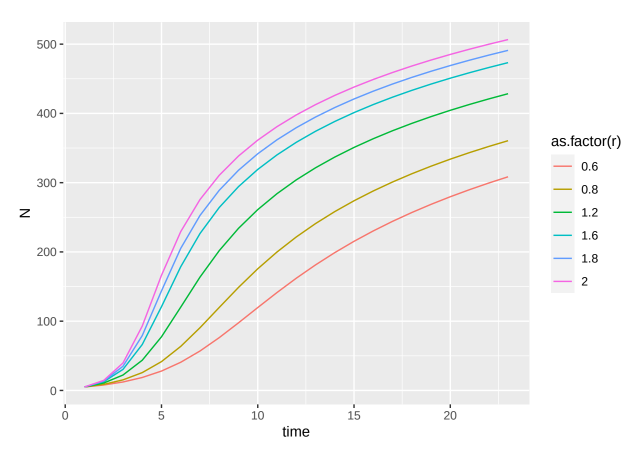
```
# Plot simulation: base R
plot(N, xlab = "time", ylab = "N", pch = 19, col = "black")
```



Step 3. Create random simulations of data, using prior distributions for model parameters Do you recall an exercise where we looked at logistic growth across a range of values for the growth rate r? We used code that looks like this (using disc_log from above):

```
# Simulation of model for t time steps
lambda_1_range <- c(0.6, 0.8, 1.2, 1.6, 1.8, 2.0)
N.g <- numeric()
N1_0 <- 5
for(p in 1:length(lambda_1_range)){
    N <- rep(NA,t)
    N[1] <- N1_0
    for(i in 2:t){
        N[i] <- disc_log(lambda_1_range[p],N1_0,alpha_11)
        N1_0 <- N[i]
    }
    dat <- as.data.frame(N)
    dat$time <- as.numeric(rownames(dat))
    dat$r <- rep(lambda_1_range[p],t)
    N.g <- rbind(N.g,dat)
    N1_0 <- 5
}</pre>
```

```
# Plot in ggplot
ggplot2::ggplot(N.g,ggplot2::aes(x=time,y=N,col=as.factor(r))) + ggplot2::geom_line()
```



In this exercise, you will choose a *prior distribution* for each of the model parameters to estimate, and you will use a random draw from that prior distribution for each repetition of the simulation. This is how we create the variables par.sim and stat.sim needed to use ABC, to estimate the unknown parameter values for Gause's Paramecium. I demonstrate how this works for a discrete-time logistic example, using data from Gause's Paramecium.

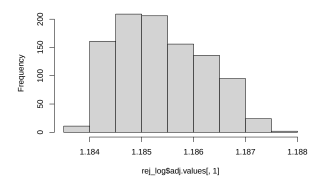
```
## 1. Get observed data
#load competition data
data("gause_1934_science_f02_03")
#subset out data from species grown alone (not in mixtire)
alone_dat <- gause_1934_science_f02_03[gause_1934_science_f02_03$Treatment=="Pc",]
#extract time and species data
time <- alone_dat$Day
species <- data.frame(alone_dat$Volume_Species1)
colnames(species) <- "P_caudatum"

## 2. Arrange observed data for ABC
stat.obs <- species$P_caudatum

## 3. Create a *random* simulation of disc_log, with draws from prior distributions
## for all model parameters. Save the time series values and values for the
## parameters in variables par.sim and stat.sim for ABC.
rand_disc_log <- function(N1_0=N1_0, disc_log, t){</pre>
```

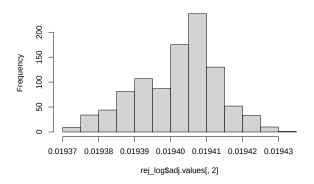
```
## Choose random values of lambda_1 from a prior distribution
  lambda_1r <- runif(1,min=1.2,max=3) # Random uniform distribution between 0-3
  alpha 11r <- rbeta(1,shape1=1,shape2=50) # Random beta distribution, bounded 0-1
  #biased towards low values
  N \leftarrow rep(NA,t)
  N[1] \leftarrow N1 O
  for(i in 2:t){
    N[i] <- disc_log(lambda_1r,N1_0=N1_0,alpha_11r)</pre>
    N1 0 <- N[i]
  }
  return(list(N=N,lambda_1r=lambda_1r,alpha_11r=alpha_11r))
}
## 4. Run this function 1000 times, and save all of the time series and the
## associated parameter values for later ABC
num iter <- 1000000
num_par <- 2 # to estimate: lambda_11 and alpha_11</pre>
par_name <- c("lambda_11","alpha_11")</pre>
par.sim <- array(NA,dim=c(num_iter,num_par),dimnames=list(NULL,par_name)) # to save values of parameter
stat.sim <- array(NA,dim=c(num_iter,t)) # to save values of N over time
colnames(stat.sim) <- paste("S",1:t,sep="")</pre>
for(k in 1:num iter){
  N1_0 <- species$P_caudatum[1]
  simul <- rand_disc_log(N1_0,disc_log,t)</pre>
  par.sim[k,1] <- simul$lambda_1r</pre>
 par.sim[k,2] <- simul$alpha_11r</pre>
  stat.sim[k,] <- simul$N</pre>
}
## 5. Use ABC to estimate the parameter values in the discrete-time growth model
colnames(stat.sim) <- paste("S",1:23,sep="")</pre>
rej_log <- abc::abc(target=stat.obs, param=par.sim, sumstat=stat.sim, tol=.001, method ="neuralnet")
## Warning: All parameters are "none" transformed.
## 12345678910
## 12345678910
## 6. Visualize the posterior distributions of the model parameters
par(mfrow=c(1,1))
hist(rej_log$adj.values[,1],main="posterior distribution of lambda_11")
```

posterior distribution of lambda_11



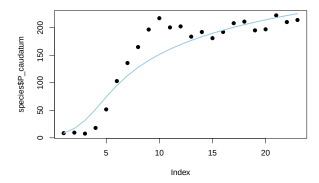
hist(rej_log\$adj.values[,2],main="posterior distribution of alpha_11")

posterior distribution of alpha_11



```
## 7. Add a curve to the observed data that shows the ABC-predicted parameter values
# 7a. Simulate data with the average 'accepted' parameter values
lambda_1 <- mean(rej_log$adj.values[,1])</pre>
alpha_11 <- mean(rej_log$adj.values[,2])</pre>
N1_0 <- species$P_caudatum[1]
t <- 23
# model function
disc_log <- function(lambda_1,N1_0,alpha_11){</pre>
  Nt1 <- N1_0 + N1_0*(lambda_1*exp(-alpha_11*N1_0))</pre>
  return(Nt1)
}
# Simulation of model for t time steps
N \leftarrow rep(NA,t)
N[1] <- N1_0
for(i in 2:t){
  N[i] <- disc_log(lambda_1,N1_0,alpha_11)</pre>
  N1_0 \leftarrow N[i]
plot(species$P_caudatum,pch=19,col="black")
lines(N,lwd=2,col="lightgray")
# 7b. Simulate data with 95% highest and lowest values for each time point
lambda_1 <- quantile(rej_log$adj.values[,1],probs=c(0.025,.975))</pre>
```

```
alpha_11 <- quantile(rej_log$adj.values[,2],probs=c(0.025,.975))</pre>
# 2.5%
N1_0 <- species$P_caudatum[1]
N \leftarrow rep(NA,t)
N[1] <- N1_0
for(i in 2:t){
  N[i] <- disc_log(lambda_1[1],N1_0,alpha_11[1])</pre>
  N1_O <- N[i]
}
lines(1:23,N,lwd=0.5,col="skyblue")
# 97.5%
N1_0 <- species$P_caudatum[1]
N <- rep(NA,t)
N[1] <- N1_0
for(i in 2:t){
  N[i] <- disc_log(lambda_1[2],N1_0,alpha_11[2])</pre>
  N1_0 <- N[i]
lines(1:23,N,lwd=0.5,col="skyblue")
```



Your goal is to repeat this ABC procedure and estimate the most likely values for the 2 Paramecium species from the dataset in Step 1a. The code above is your 'template'm but using only 1 species. Add in the model and parameters for Species #2, and be sure to add in draws of parameter values from the prior distributions for those new parameters.

Warning: All parameters are "none" transformed.

