Lab 6

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Learning goals

You will learn how to:

- 1. Program the likelihood function of a model.
- 2. Estimate the parameters of a model through maximum likelihood.
- 3. Estimate the confidence intervals of the model parameters through profiling and the quadratic approximation.
- (4). Estimate parameters in a Bayesian framework and how parameter uncertainty can be assessed (time permitting and when of interest to you).

Fitting models to data

In this exercise you will learn how to fit models to data through means of maximum likelihood and compare the likelihood of different models (hypotheses). Fitting a model to data through likelihood requires that you take four steps:

- 1. Specify how the dependent variable depends on the independent variable, i.e. specify a function how the mean of y depends on the value of x.
- 2. Specify a probability distribution to describe the deviations of the observations from the mean
- 3. Choose the parameters of the deterministic model and the probability model such that the negative log likelihood is lowest.
- 4. Compare the likelihood of alternative models (change the deterministic function or the stochastic function) and compare with AIC(c) or BIC which model is most parsimonious.

To fit a model through means of maximum likelihood you need to specify a function that calculate the negative log likelihood (NLL) based on the data and the parameter values. For example to calculate the NLL of a linear model and a normal distribution the following function works:

```
nll = function(par,y,x){
  a = par[1]
  b = par[2]
  sd = par[3]
  # this calculates the mean y for a given value of x: the deterministic function
  mu = a+b*x
  # this calculates the likelihood of the function given the probability
  # distribution, the data and mu and sd
  nll = -sum(dnorm(y,mean=mu,sd=sd,log=T))
  return(nll)
}
```

Note that -sum(log(dnorm(y,mean=mu,sd=sd))) should not be used as it may lead to underflow (the computer cannot store very very small probabilities) and therefore to optimisation problems.

Next we specify a function to find the maximum likelihood estimate

```
par=c(a=1,b=1,c=1) # initial parameters
opt1 = optim(par=par,nll,x=x,y=y) # y represents the data, x the independent variable
```

It can also be done through ${\tt mle2}$

```
nll.mle = function(a,b,sd){
    # this calculates the mean y for a given value of x: the deterministic function
    mu = a+b*x
    # this calculates the likelihood of the function given the probability
    # distribution, the data and mu and sd
    nll = -sum(dnorm(y,mean=mu,sd=sd,log=T))
    return(nll)
}

# the data should be supplied through data and the parameters through list().
mle2.1 = mle2(nll.mle,start=list(a=1,b=1,sd=1),data=data.frame(x,y))
summary(mle2.1)
```

Made-up data: negative binomial

The simplest thing to do to convince yourself that your attempts to estimate parameters are working is to simulate the "data" yourself and see if you get close to the right answers back.

Start by making up some negative binomial 'data'': first, set the random-number seed so we get consistent results across differentR' sessions:

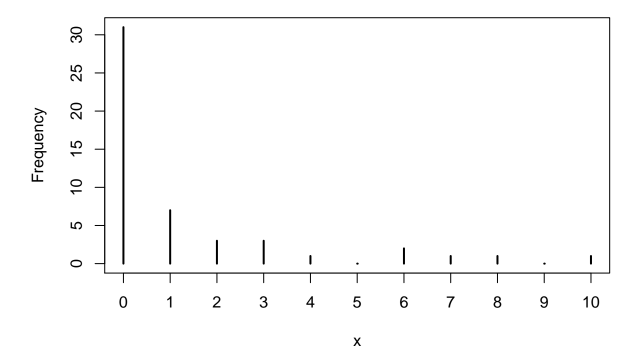
```
set.seed(1001)
```

Generate 50 values with $\mu = 1$, k = 0.4 (save the values in variables in case we want to use them again later, or change the parameters and run the code again):

```
mu.true=1
k.true=0.4
x = rnbinom(50,mu=mu.true,size=k.true)
```

Take a quick look at what we got:

```
plot(table(factor(x,levels=0:max(x))),
    ylab="Frequency",xlab="x")
```



(reminder: I won't always draw the pictures, but it's good to make a habitat of examining your variables (with summary() etc. and graphically) as you go along to make sure you know what's going on!)

Negative log-likelihood function for a simple draw from a negative binomial distribution: the first parameter, p, will be the vector of parameters, and the second parameter, dat, will be the data vector (in case we want to try this again later with a different set of data; by default we'll set it to the 'x} vector we just drew randomly).

```
NLLfun1 = function(p,dat=x) {
  mu=p[1]
  k=p[2]
  -sum(dnbinom(x,mu=mu,size=k,log=TRUE))
}
```

Calculate the negative log-likelihood for the true values. I have to combine these values into a vector with c() to pass them to the negative log-likelihood function. Naming the elements in the vector is optional but will help keep things clear as we go along:

```
nll.true=NLLfun1(c(mu=mu.true,k=k.true)); nll.true
```

The NLL for other parameter values that I know are way off ($\mu = 10, k = 10$):

```
NLLfun1(c(mu=10,k=10))
```

```
## [1] 291.4351
```

Much higher negative log-likelihood, as it should be.

Find the method-of-moments estimates for μ and k:

```
m = mean(x)
v = var(x)
mu.mom = m
k.mom = m/(v/m-1)
```

Negative log-likelihood estimate for method of moments parameters:

```
nll.mom=NLLfun1(c(mu=mu.mom,k=k.mom)); nll.mom
```

```
## [1] 72.08996
```

Despite the known bias, this estimate is better (lower negative log-likelihood) than the "true" parameter values. The Likelihood Ratio Test would say, however, that the difference in likelihoods would have to be greater than $\chi_2^2(0.95)/2$ (two degrees of freedom because we are allowing both μ and k to change):

```
ldiff=nll.true-nll.mom;ldiff
```

```
## [1] 0.5576733
```

```
qchisq(0.95,df=2)/2
```

```
## [1] 2.995732
```

So — better, but not significantly better at p = 0.05. pchisq(2*ldiff,df=2,lower.tail=FALSE) would tell us the exact p-value if we wanted to know.)

But what is the MLE? Use optim with the default options (Nelder-Mead simplex method) and the method-of-moments estimates as the starting estimates (par):

```
01 = optim(fn=NLLfun1,par=c(mu=mu.mom,k=k.mom),hessian=TRUE);
01
```

```
## $par
##
                      k
          mu
## 1.2602356 0.2884793
##
## $value
  [1] 71.79646
##
##
## $counts
## function gradient
##
         45
##
## $convergence
##
  [1] 0
##
## $message
## NULL
##
## $hessian
##
               mu
## mu 7.387808317
                   0.004901576
## k 0.004901576 97.372581394
```

The optimization result is a list with elements:

- the best-fit parameters (01\$par, with parameter names because we named the elements of the starting vector—see how useful this is?);}
- the minimum negative log-likelihood (01\$value);

- information on the number of function evaluations (O1\$counts; the gradient part is NA because we didn't specify a function to calculate the derivatives (and the Nelder-Mead algorithm wouldn't have used them anyway)
- information on whether the algorithm thinks it found a good answer O1\$convergence, which is zero if R thinks everything worked and uses various numeric codes (see ?optim for details) if something goes wrong;
- O1\$message which may give further information about the when the fit converged or how it failed to converge;
- because we set hessian=TRUE, we also get O1\$hessian, which gives the (finite difference approximation of) the second derivatives evaluated at the MLE.

The minimum negative log-likelihood (round(01\$value,2)) is better than either the negative log-likelihood corresponding to the method-of-moments parameters (round(nll.mom,2)) or the true parameters (round(nll.true,2)), but all of these are within the LRT cutoff.

Now let's find the likelihood surface, the profiles, and the confidence intervals.

The likelihood surface is straightforward: set up vectors of μ and k values and run for loops, set up a matrix to hold the results, and run for loops to calculate and store the values. Let's try μ from 0.4 to 3 in steps of 0.05 and k from 0.01 to 0.7 in steps of 0.01. (I initially had the μ vector from 0.1 to 2.0 but revised it after seeing the contour plot below.)

```
muvec = seq(0.4,3,by=0.05)
kvec = seq(0.01,0.7,by=0.01)
```

The matrix for the results will have rows corresponding to μ and columns corresponding to k:

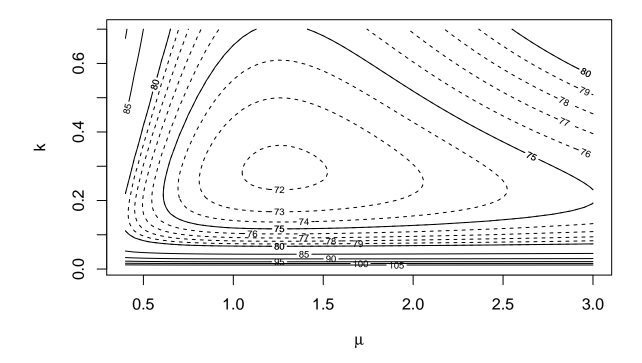
```
resmat = matrix(nrow=length(muvec),ncol=length(kvec))
```

Run the for loops:

```
for (i in 1:length(muvec)) {
  for (j in 1:length(kvec)) {
    resmat[i,j] = NLLfun1(c(muvec[i],kvec[j]))
  }
}
```

Drawing a contour: the initial default choice of contours doesn't give us fine enough resolution (it picks contours spaced 5 apart to cover the range of the values in the matrix), so I added levels spaced 1 log-likelihood unit apart from 70 to 80 by doing a second contour plot with add=TRUE.

```
contour(muvec,kvec,resmat,xlab=expression(mu),ylab="k")
contour(muvec,kvec,resmat,levels=70:80,lty=2,add=TRUE)
```



Maximum likelihood and covariates

The following steps will lead you through the model fitting procedure when we have a covariate.

1. Take the first dataset (or another one from the six that we have been working with before) and tweak the above functions such that it matches with the deterministic and stochastic model that you have chosen. In case you got stuck in the previous exercises where you had to choose a deterministic function and a stochastic function see next page for suggestions.

hint: In a previous exercise you have eyeballed the parameter values of the functions, you can use these as starting values.

hint: In case you get convergence problems, further adapt your starting values, or choose a different optimizer. For example Nelder-Mead is a robust one, e.g. method = "Nelder-Mead".

- 2. Change the determinstic function for a possible alternative determinstic function
- 3. Compare the likelihoods of the data given both models
- 4. Apply model selection criteria and conclude which model fits that data best.
- 5. Does the model makes sense from a biological perspective?

Optional and time permitting:

6. Repeat the above procedure for the other 5 datasets

Parameter uncertainty in the estimated parameters

The next exercise will be about assessing the uncertainty in the estimated parameters. Bolker recommends to use the likelihood profile for assessing the uncertainty in the parameters because this one is more accurate than the one based on the Hessian matrix. We take a number of steps to get a feel for the uncertainty in the parameter estimates.

Take the first dataset and assume that the function was generated by the monomolecular function $a(1-e^{(-bx)})$. Fit this model with normally distributed errors through this data with mle2 and inspect the coefficients.

- 1. Calculate for the parameters a and b the NLL for a number of combinations of a and b and plot the NLL surface using contour plot. What is your conclusion about the relationship between the two parameters? Are they fully independent? hint: you can use a for a double for-loop to run over all parameters hint: see the example on the made-up data for inspiration
- 2. Calculate the confidence intervals of the parameters through constructing the likelihood profile and the quadratic approximation. Consult page 193ff of Bolker for inspiration and use the following pseudocode to achieve this:
 - a. Adapt the likelihood function such that one parameter is not optimised but chosen by you, say parameter a.
 - b. Vary a of a range and optimise the other parameteters.
 - c. Plot the NLL as a function of parameter a.
 - d. Find the values of a that enclose $-L + \chi^2(1-\alpha)/2$
 - e. Compare your results with the function confint()
- 3. Calculate the confidence intervals through the quadratic approximation. Take the following steps to achieve this:
 - a. Get the standard error of the parameter estimates through vcov. Note that vcov return the variance/covariance matrix
 - b. Calculate the interval based on the fact that the 95% limits are 1.96 (qnorm(0.975,0,1)) standard deviation units away from the mean.
- 4. Plot the confidence limits of the both method and compare the results. What is your conclusion?

Bayesian exercise... Alejandro

Hints for choosing deterministic functions and stochastic functions

1. Deterministic functions

dataset 1 light response curve. There are a number of options of functions to choose from, depending on the level of sophistication: $\frac{ax}{(b+x)}$, $a(1-e^{(-bx)})$, $\frac{1}{2\theta}(\alpha I + p_{max} - \sqrt{(\alpha I + p_{max})^2 - 4\theta I p_{max}})$ see page 98. A parameter d can be added in all cases to shift the curve up or down.

dataset 2 The dataset describes a functional responses. Bolker mentions four of those min(ax, s) $\frac{ax}{(b+x)}$, $\frac{ax^2}{(b^2+x^2)}$, $\frac{ax^2}{(b+cx+x^2)}$

dataset 3 Allometric relationships generally have the form ax^b

dataset 4 This could be logistic growth $n(t) = \frac{K}{1 + (\frac{K}{n_0})e^{-rt}}$ or the gompertz function $f(x) = e^{-ae^{-bx}}$

dataset 5 What about a negative exponential? ae-bx or a power function ax^b

dataset 6 Species reponse curves are curves that describe the probability of presence as a function of some factor. A good candidate good be a unimodel response curve. You could take the equation of the normal distribution without the scaling constant: e.g. $ae^{\frac{-(x-\mu)^2}{2\sigma^2}}$

2. Stochastic functions/Probability distributions

dataset 1 y represents real numbers and both positive and negative numbers occur. This implies that we should choose a continuous probability distribution. In addition, the numbers seems unbound. Within the family of continuous probability distributions, the normal seems a good candidate distribution because this one runs from -inf to +inf. In contrast the Gamma and the Lognormal only can take positive numbers, so these distributions cannot handle the negative numbers. In addition, the beta distribution is not a good candidate because it runs from 0-1.

dataset 2 y represents real numbers and only positive numbers occur. The data represents a functional response (intake rate of the predator), and it is likely that you can only measure positive numbers (number of prey items per unit of time). This implies that we should choose a continuous probability distribution. Within the family of continuous probability distributions, the Gamma and the Lognormal could be taken as candidate distributions because they can only take positive numbers (beware that the Gamma cannot take 0). However, you could try to use a normal as well.

dataset 3 y seems represents counts (this is the cone dataset that is introduced in ch. 6.). Given that it contains counts we can pick a distribution from the family of discrete distributions. The Poisson and the Negative Binomial could be good candidates to describe this type of data.

dataset 4 y represents population size over time. From looking at the data, they seems to represent counts. Given that it contains counts we can pick a distribution from the family of discrete distributions. The Poisson and the Negative Binomial could be good candidates to describe this type of data.

dataset 5 No information is given on y. The data clearly seems to represent counts. Thus the same reasoning applies here as to the two previous datasets.

dataset 6 The data (y) represents species occurences (presence/absence). The binomial model would be a good model to predict the probability of presence.