

Detection function fitting

Practical 3, Intermediate Distance Sampling workshop, CREEM, 2018

Aims

By the end of this practical you should feel confident doing the following:

- Loading data from ArcGIS .gdb files
- Working on a `data.frame` in R to get it into the correct format for `Distance`
- Fitting a detection function using `ds()`
- Checking detection functions
- Making at goodness of fit plots
- Selecting models using AIC
- Estimating abundance (using R and maths!)

Preamble

First need to load the requisite R libraries

```
library(rgdal)
```

```
## Warning: package 'rgdal' was built under R version 3.4.4
```

```
## Loading required package: methods
```

```
## Loading required package: sp
```

```
## rgdal: version: 1.3-4, (SVN revision 766)
```

```
## Geospatial Data Abstraction Library extensions to R successfully loaded
```

```
## Loaded GDAL runtime: GDAL 2.1.3, released 2017/20/01
```

```
## Path to GDAL shared files: /Library/Frameworks/R.framework/Versions/3.4/Resources/library/rgdal/gdal
```

```
## GDAL binary built with GEOS: FALSE
```

```
## Loaded PROJ.4 runtime: Rel. 4.9.3, 15 August 2016, [PJ_VERSION: 493]
```

```
## Path to PROJ.4 shared files: /Library/Frameworks/R.framework/Versions/3.4/Resources/library/rgdal/p
```

```
## Linking to sp version: 1.3-1
```

```
library(ggplot2)
```

```
library(Distance)
```

```
## Loading required package: mrds
```

```
## This is mrds 2.2.0
```

```
## Built: R 3.4.3; ; 2018-06-25 18:07:22 UTC; unix
```

```
##
```

```
## Attaching package: 'Distance'
```

```
## The following object is masked from 'package:mrds':
```

```
##
```

```
##      create.bins
```

```
library(knitr)
```

Load the data

The observations are located in a “geodatabase” we created in Arc. We want to pull out the “Sightings” table (called a “layer”) and make it into a `data.frame` (so it’s easier for R to manipulate).

```
distdata <- readOGR("Analysis.gdb", layer="Sightings")
```

```
## OGR data source with driver: OpenFileGDB
## Source: "/Users/dill/current/workshop-sites/stand-intermed-2018/practicals/Analysis.gdb", layer: "Si
## with 137 features
## It has 7 fields
```

```
distdata <- as.data.frame(distdata)
```

We can check it has the correct format using `head`:

```
head(distdata)
```

```
##      Survey GroupSize SeaState  Distance      SightingTime SegmentID
## 1 en04395          2      3.0 246.0173 2004/06/28 10:22:21         48
## 2 en04395          2      2.5 1632.3934 2004/06/28 13:18:14         50
## 3 en04395          1      3.0 2368.9941 2004/06/28 14:13:34         51
## 4 en04395          1      3.5  244.6977 2004/06/28 15:06:01         52
## 5 en04395          1      4.0 2081.3468 2004/06/29 10:48:31         56
## 6 en04395          1      2.4 1149.2632 2004/06/29 14:35:34         59
##      SightingID coords.x1 coords.x2
## 1              1  -65.636   39.576
## 2              2  -65.648   39.746
## 3              3  -65.692   39.843
## 4              4  -65.717   39.967
## 5              5  -65.820   40.279
## 6              6  -65.938   40.612
```

The `Distance` package expects certain column names to be used. Renaming is much easier to do in R than ArcGIS, so we do it here.

```
distdata$distance <- distdata$Distance
distdata$object <- distdata$SightingID
distdata$size <- distdata$GroupSize
```

Let’s see what we did:

```
head(distdata)
```

```
##      Survey GroupSize SeaState  Distance      SightingTime SegmentID
## 1 en04395          2      3.0 246.0173 2004/06/28 10:22:21         48
## 2 en04395          2      2.5 1632.3934 2004/06/28 13:18:14         50
## 3 en04395          1      3.0 2368.9941 2004/06/28 14:13:34         51
## 4 en04395          1      3.5  244.6977 2004/06/28 15:06:01         52
## 5 en04395          1      4.0 2081.3468 2004/06/29 10:48:31         56
## 6 en04395          1      2.4 1149.2632 2004/06/29 14:35:34         59
##      SightingID coords.x1 coords.x2  distance object size
## 1              1  -65.636   39.576 246.0173         1     2
## 2              2  -65.648   39.746 1632.3934         2     2
## 3              3  -65.692   39.843 2368.9941         3     1
## 4              4  -65.717   39.967  244.6977         4     1
## 5              5  -65.820   40.279 2081.3468         5     1
## 6              6  -65.938   40.612 1149.2632         6     1
```

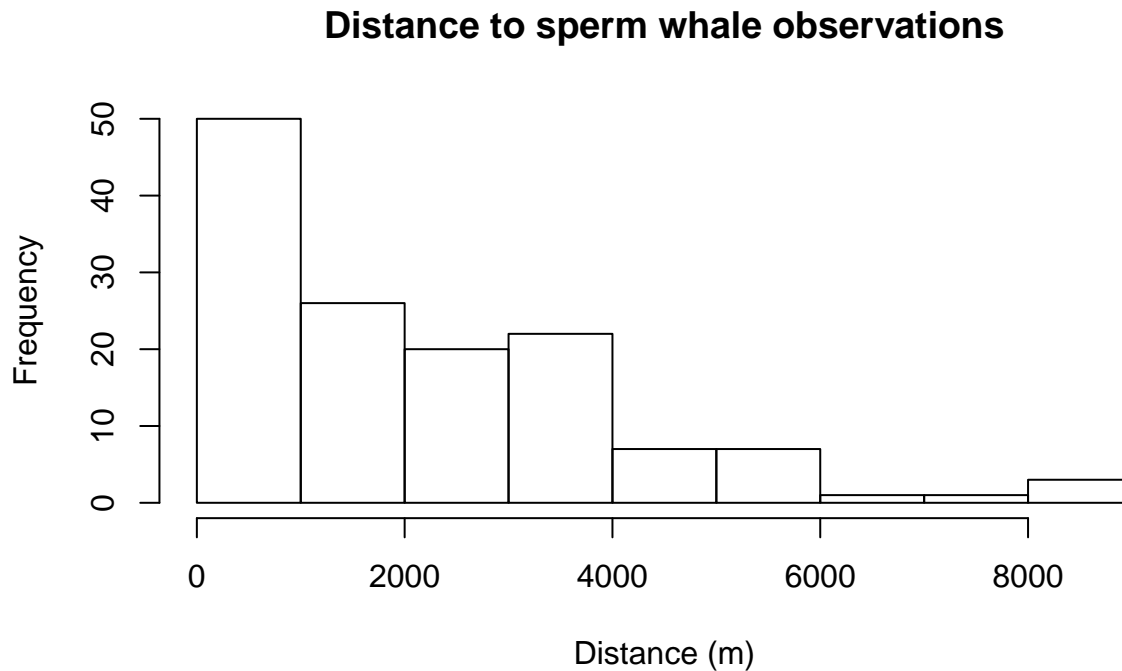


Figure 1: Distribution of observed perpendicular detection distances.

We now have four “extra” columns.

Exploratory analysis

Before setting off fitting detection functions, let’s look at the relationship of various variables in the data.

Don’t worry too much about understanding the code that generates these plots at the moment.

Distances

Obviously, the most important covariate in a distance sampling analysis is distance itself. We can plot a histogram of the distances to check that (1) we imported the data correctly and (2) it conforms to the usual shape for line transect data.

```
hist(distdata$distance, xlab="Distance (m)", main="Distance to sperm whale observations")
```

Size and distance

We might expect that there will be a relationship between the distance at which we see animals and the size of the groups observed (larger groups are easier to see at larger distances), so let’s plot that to help us visualise the relationship.

```
# plot of size versus distance and sea state vs distance, linear model and LOESS smoother overlay
# put the data into a simple format, only selecting what we need
```

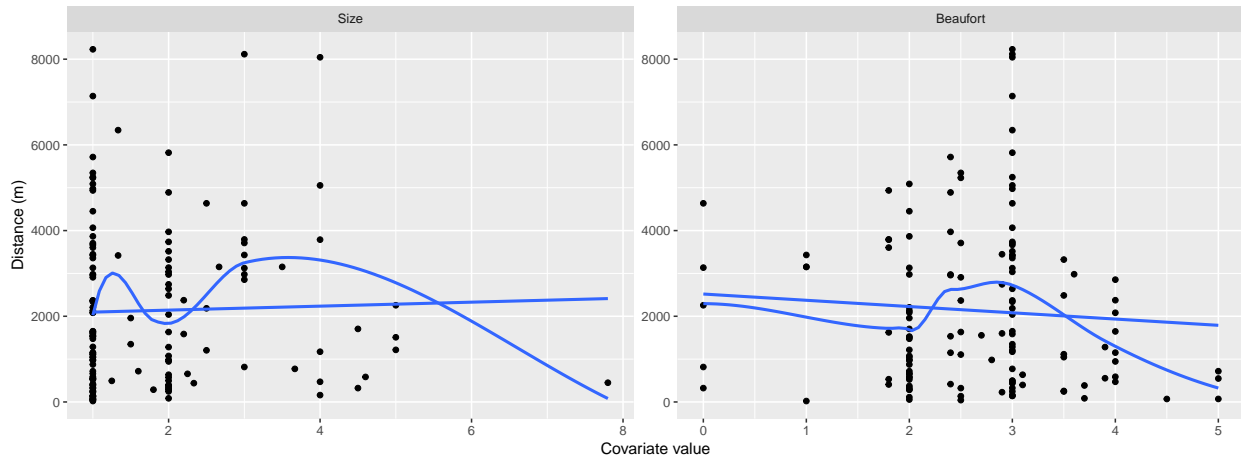


Figure 2: Effect of group size upon detection distances.

```

distplot <- distdata[,c("distance", "size", "SeaState")]
names(distplot) <- c("Distance", "Size", "Beaufort")
library(reshape2)
# "melt" the data to have only three columns (try head(distplot))
distplot <- melt(distplot, id.vars="Distance", value.name="covariate")

# make the plot
p <- ggplot(distplot, aes(x=covariate, y=Distance)) +
  geom_point() +
  facet_wrap(~variable, scale="free") +
  geom_smooth(method="loess", se=FALSE) +
  geom_smooth(method="lm", se=FALSE) +
  labs(x="Covariate value", y="Distance (m)")
print(p)

```

Distance and sea state

We might also expect that increasing sea state would result in a drop in observations. We can plot histograms of distance for each sea state level (making the sea state take only values 0,1,2,4,5 for this).

```

distdata$SeaStateCut <- cut(distdata$SeaState, seq(0,5,by=1), include.lowest=TRUE)
p <- ggplot(distdata) +
  geom_histogram(aes(distance)) +
  facet_wrap(~SeaStateCut) +
  labs(x="Distance (m)", y="Count")
print(p)

```

```
## `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
```

Survey effect

Given we are including data from two different surveys we can also investigate the relationship between survey and distances observed.

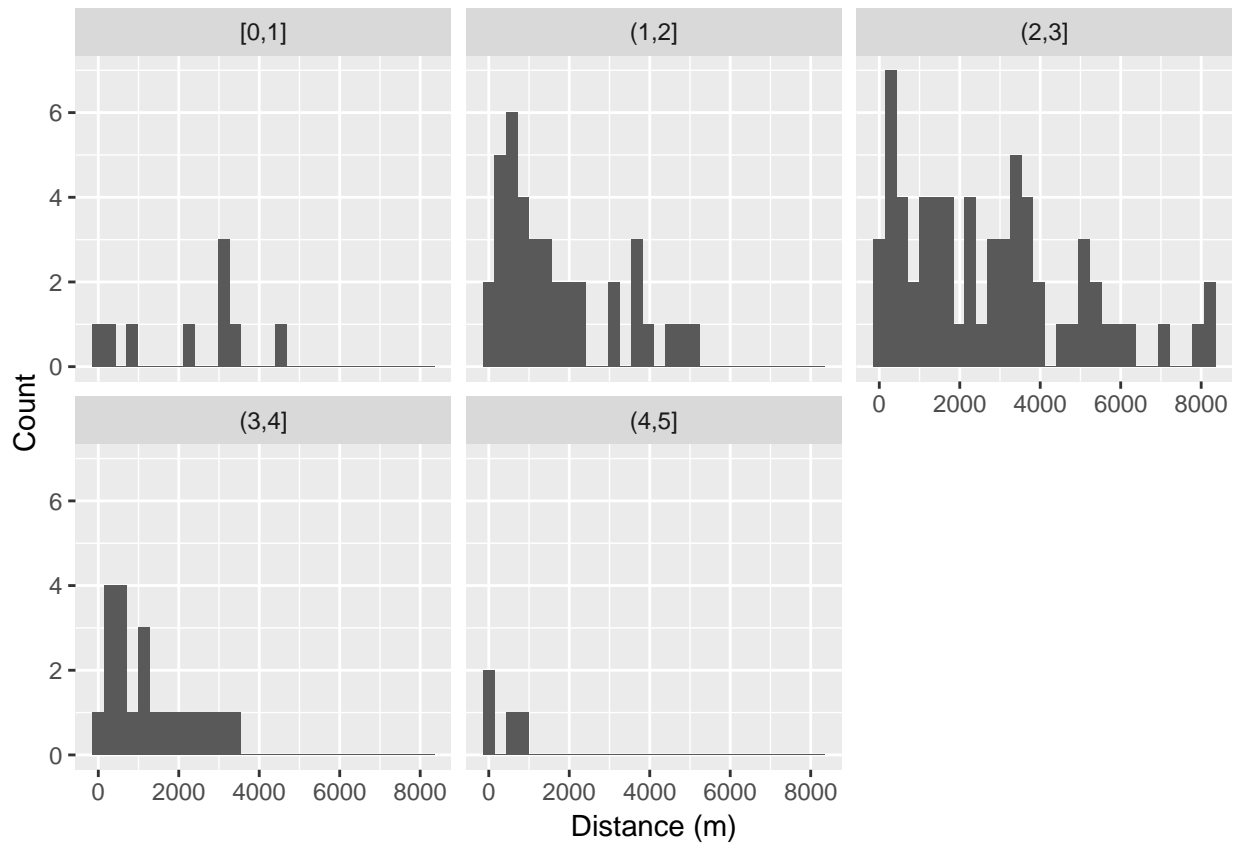


Figure 3: Effect of sea state upon detection distances.

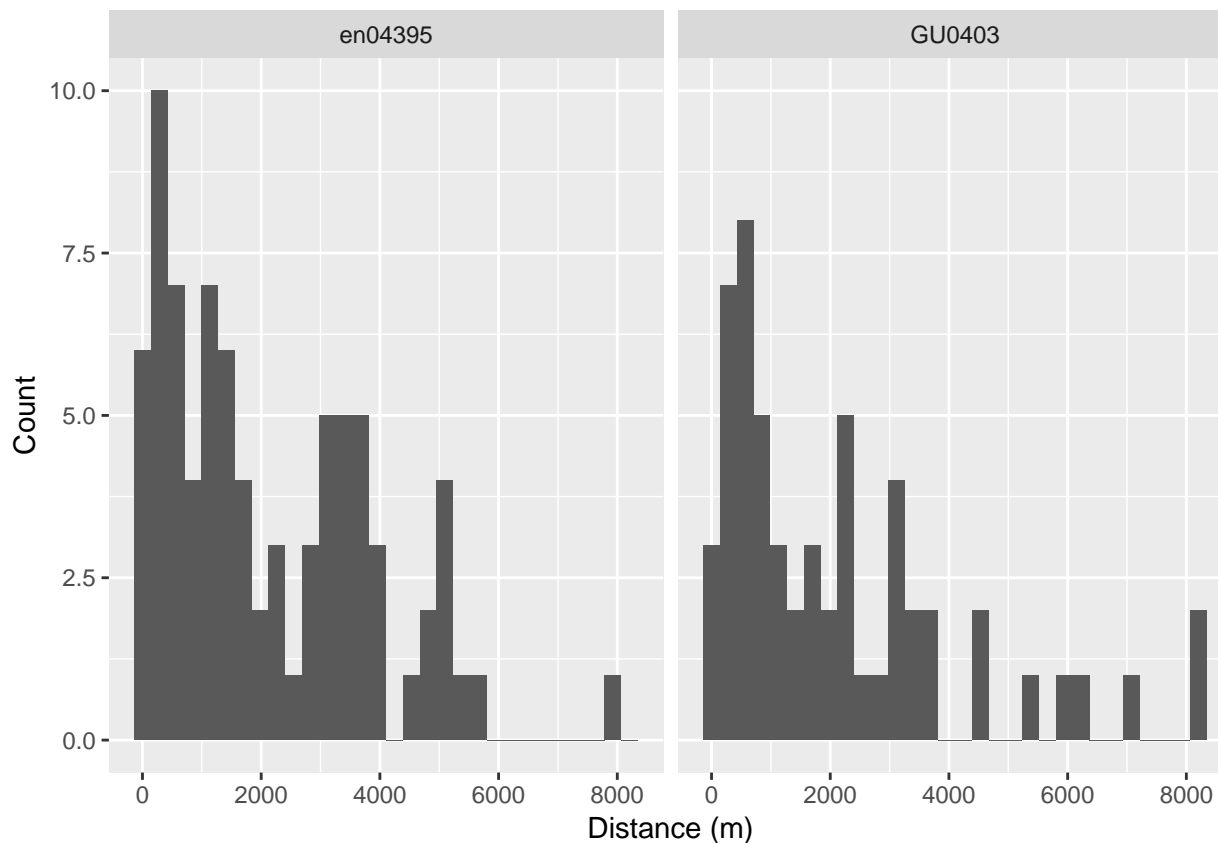


Figure 4: Effect of survey upon detection distances.

```
p <- ggplot(distdata) +
  geom_histogram(aes(distance)) +
  facet_wrap(~Survey) +
  labs(x="Distance (m)", y="Count")
print(p)
```

```
## `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
```

Fitting detection functions

It's now time to fit some detection function models. We'll use the `ds()` function from the `Distance` package to fit the detection function. You can access the help file for the `ds()` function by typing `?ds` – this will give you information about what the different arguments to the function are and what they do.

We can fit a very simple model with the following code:

```
df_hn <- ds(data=distdata, truncation=6000, key="hn", adjustment=NULL)
```

```
## Fitting half-normal key function
```

```
## Key only model: not constraining for monotonicity.
```

```
## AIC= 2252.06
```

```
## No survey area information supplied, only estimating detection function.
```

Let's dissect the call and see what each argument means:

- `data=distdata`: the data to use to fit the model, as we prepared above.
- `truncation=6000`: set the truncation distance. Here, observations at distances greater than 6000m will be discarded before fitting the detection function.
- `key="hn"`: the key function to use for the detection function, in this case half-normal (?ds lists the other options).
- `adjustment=NULL`: adjustment term series to fit. NULL here means that no adjustments should be fitted (again ?ds lists all options).

Other useful arguments for this practical are:

- `formula=`: gives the formula to use for the scale parameter. By default it takes the value `~1`, meaning the scale parameter is constant and not a function of covariates.
- `order=`: specifies the "order" of the adjustments to be used. This is a number (or vector of numbers) specifying the order of the terms. For example `order=2` fits order 2 adjustments, `order=c(2,3)` will fit a model with order 2 and 3 adjustments (mathematically, it only makes sense to include order 3 with order 2). By default the value is NULL which has `ds()` select the number of adjustments by AIC.

Summaries

We can look at the summary of the fitted detection function using the `summary()` function:

```
summary(df_hn)
```

```
##
## Summary for distance analysis
## Number of observations : 132
## Distance range       : 0 - 6000
##
## Model : Half-normal key function
## AIC   : 2252.06
##
## Detection function parameters
## Scale coefficient(s):
##           estimate      se
## (Intercept) 7.900732 0.07884776
##
##           Estimate      SE      CV
## Average p      0.5490484 0.03662569 0.06670757
## N in covered region 240.4159539 21.32287580 0.08869160
```

Goodness of fit

Goodness of fit quantile-quantile plot and test results can be accessed using the `ddf.gof()` function:

```
ddf.gof(df_hn$ddf)
```

```
##
## Goodness of fit results for ddf object
##
## Chi-square tests
##           [0,545] (545,1.09e+03] (1.09e+03,1.64e+03] (1.64e+03,2.18e+03]
## Observed 33.000000    20.00000000    19.00000000    8.000000
## Expected 21.708156    20.84245788    19.21325454    17.005089
## Chisquare 5.873634     0.03405238     0.002366986    4.768669
```

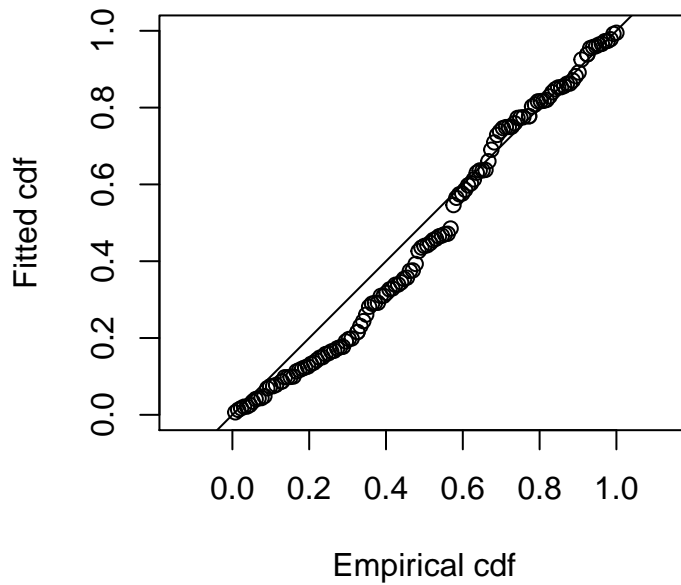


Figure 5: Goodness of fit QQ plot of half-normal detection function.

```
##          (2.18e+03,2.73e+03] (2.73e+03,3.27e+03] (3.27e+03,3.82e+03]
## Observed          9.000000          13.0000000          14.000000
## Expected          14.450499          11.7899695           9.235669
## Chisquare          2.055842           0.1241881           2.457737
##          (3.82e+03,4.36e+03] (4.36e+03,4.91e+03] (4.91e+03,5.45e+03]
## Observed          3.000000          4.0000000          7.000000
## Expected          6.946241          5.0159948           3.477683
## Chisquare          2.241906           0.2057908           3.567525
##          (5.45e+03,6e+03]      Total
## Observed          2.00000000 132.00000
## Expected          2.31498601 132.00000
## Chisquare          0.04285822 21.37457
##
## P = 0.011087 with 9 degrees of freedom
##
## Distance sampling Cramer-von Mises test (unweighted)
## Test statistic = 0.396179 p-value = 0.0739475
```

Note two things here: 1. We use the `$ddf` element of the detection function object 2. We're ignoring the χ^2 test results, as they rely on binning the distances to calculate test statistics whereas Cramer-von Mises and Kolmogorov-Smirnov tests do not.

Plotting

We can plot the models simply using the `plot()` function:

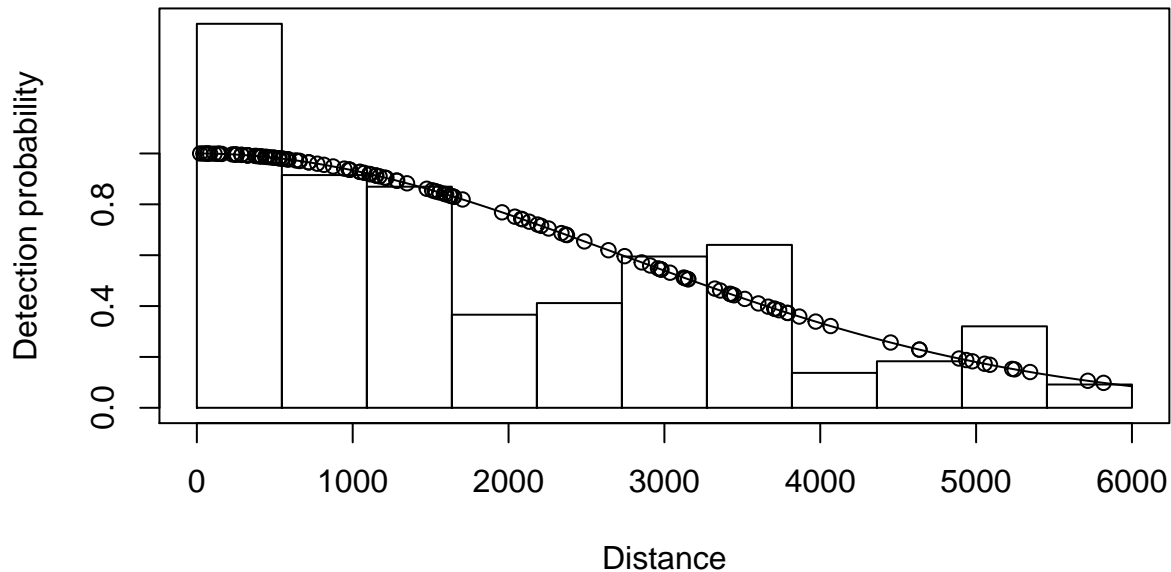


Figure 6: Half-normal detection function.

```
plot(df_hn)
```

The dots on the plot indicate the distances where observations are. We can remove them (particularly useful for a model without covariates) using the additional argument `showpoints=FALSE` (try this out!).

Now you try...

Now try fitting a few models and comparing them using AIC. Don't try to fit all possible models, just try a selection (say, a hazard-rate, a model with adjustments and a couple with different covariates). You can also try out changing the truncation distance.

Here's an example to work from. Some tips before you start:

- You can include as many lines as you like in a given chunk (though you may find it more manageable to separate them up, remembering each time to give the chunk a unique name).
- You can run the current line of code in RStudio by hitting `Control+Enter` (on Windows/Linux; `Command+Enter` on Mac).
- Giving your models informative names will help later on! Here I'm using `df_` to indicate that this is a detection function, then shortened forms of the model form and covariates, separated by underscores, but use what makes sense to you (and future you!).

```
df_hr_ss_size <- ds(distdata, truncation=6000, adjustment=NULL,
                    key="hr", formula=~SeaState+size)
```

```
## Fitting hazard-rate key function
```

```
## AIC= 2249.327
```

Table 1: Comparison of half normal and hazard rate with sea state and group size.

Model	Key function	Formula	C-vM p-value	\hat{P}_a	se(\hat{P}_a)	Δ AIC
df_hr_ss_size	Hazard-rate	SeaState + size	0.880	0.355	0.074	0.000
df_hn	Half-normal	1	0.074	0.549	0.037	2.733

No survey area information supplied, only estimating detection function.

Once you have the hang of writing models and looking at the differences between them, you should move onto the next section.

Model selection

Looking at the models individually can be a bit unwieldy – it’s nicer to put that data into a table and sort the table by the relevant statistic. The function `summarize_ds_models()` in the `Distance` package performs this task for us.

The code below will make a results table with relevant statistics for model selection in it. The `summarize_ds_models()` function takes a series of object names as its first argument. We can do that with the two models that I fitted like so:

```
model_table <- summarize_ds_models(df_hn, df_hr_ss_size)
kable(model_table, digits=3, format="latex", row.names = FALSE, escape=FALSE,
      caption = "Comparison of half normal and hazard rate with sea state and group size.")
```

(You can add the models you fitted above into this list.)

A further note about model selection for the sperm whale data

Note that there is a considerable spike in our distance data. This may be down to observers guarding the trackline (spending too much time searching near zero distance). It’s therefore possible that the hazard-rate model is overfitting to this peak. So we’d like to investigate results from the half-normal model too and see what the effects are on the final spatial models.

Estimating abundance

Just for fun, let’s estimate abundance from these models using a Horvitz-Thompson-type estimator.

We know the Horvitz-Thompson estimator has the following form:

$$\hat{N} = \frac{A}{a} \sum_{i=1}^n \frac{s_i}{p_i}$$

we can calculate each part of this equation in R:

- A is the total area of the region we want to estimate abundance for. This was $A = 5.285e + 11m^2$.
- a is the total area we surveyed. We know that the total transect length was 9,498,474m and the truncation distance. Knowing that $a = 2wL$ we can calculate a .
- s_i are the group sizes, they are stored in `df_hnddfdata$size`.
- p_i are the probabilities of detection, we can obtain them using `predict(df_hn$ddf)$fitted`.

We know that in general operations are vectorised in R, so calculating `c(1, 2, 3)/c(4, 5, 6)` will give `c(1/4, 2/5, 3/6)` so we can just divide the results of getting the s_i and p_i values and then use the `sum()` function to sum them up.

Try out estimating abundance using the formula below using both `df_hn` and your favourite model from above:

Note that in the solutions to this exercise (posted on the course website) I show how to use the function `dht()` to estimate abundance (and uncertainty) for a detection function analysis. This involves some more complex data manipulation steps, so we've left it out here in favour of getting to grips with the mathematics.

Accounting for perception bias

It's common, especially in marine surveys, for animals at zero distance to be missed by observers. There are several ways to deal with this issue. For now, we are just going to use a very simply constant correction factor to inflate the abundance.

From Palka (2006), we think that observations on the track line were such that $g(0) = 0.46$, we can apply that correction to our abundance estimate (in a very primitive way):

This kind of correction works fine when it's appropriate to use a single number to adjust by, in general we'd like to model the perception bias, for example using "mark-recapture distance sampling" techniques.

Save model objects

Save your top few models in an `RData` file, so we can load them up later on. We'll also save the distance data we used to fit out models.

```
save(df_hn, df_hr_ss_size, # add you models here, followed by commas!  
     distdata,  
     file="df-models.RData")
```

You can check it worked by using the `load()` function to recover the models.

References

Palka, D. L. (2006) *Summer Abundance Estimates of Cetaceans in US North Atlantic Navy Operating Areas*. Northeast Fisheries Science Center Reference Document 06-03