# Distance sampling online workshop

Analysis in R: Analysis of multi-species surveys

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## 1 More complex analyses

This practical is based on the Montrave songbird case study in Buckland et al. (2015, Section 5.2.2.3), with computer code under Montrave songbird case study. Both point and line transect surveys were conducted and here we use the data from the **line transect** data, although the issues (and solutions) will be similar.

These data are provided in a 'flat file' format (i.e. it contains all the necessary columns to estimate a detection function, density and abundance). While both formats are equally valid, the 'flat file' approach has a particular idiosyncrasy which we exploit here to introduce more functions and data manipulation.

Several species of birds were identified but not all species were detected on all transects. If a simple data selection is performed to select records for a particular species, then not all of the transects will be included in the resulting data (because that species may not have been seen). This doesn't matter if we are only interested in fitting detection functions, but will matter if we wish to estimate density and abundance because the effort will be too low since some of the transects are missing. To correct for this, some data frame manipulation is required. There is generally more than one way to do something in R (R Core Team, 2018) - for an alternative way see the computer code 'Montrave song bird case study' associated with Buckland et al. (2015), as well as Section 9 below.

# 2 Objectives of the practical

- 1. Data frame selection and manipulation
- 2. Extracting estimates from dht object
- 3. Customising detection function plots
- 4. Improve re-usability of code with functions

## 3 Importing the data

The data is in a 'flat file' format and contains the following columns:

- · Region.Label name of study
- Area size of study region (hectares)
- · repeats number of visits to transect
- · Sample.Label line transect identifier
- · Effort length of transect (km)
- · distance perpendicular distance (m)
- species species of bird (c=chaffinch, g=great tit, r=robin and w=wren)
- visit on which visit bird was detected.



European robin (*Erithacus rubecula*); one of the species in the Montrave study (Buckland, 2006).



Aerial view of Montrave study area. White diagonal lines represent transects walked for data analysed here.

Use the following command to import the data from the website associated with Buckland et al. (2015) and then use the head command to examine it.

**Question:** Explore the data. How many transects are there?

```
length(unique(birds$Sample.Label))
```

For now, save the transect labels to a new object as we will use them later on:

```
tran.lab <- unique(birds$Sample.Label)</pre>
```

The table command is a quick way to determine how many detections there are of each species:

```
table(birds\species)
```

As a hint of things to come, create a two-way table showing the number of detections by transect and by species. If there are zeroes in this table, it will create a challenge.

```
with(birds, table(species, Sample.Label))
```

Each of the line transects was visited twice which is not taken into account at present. However, it is straightforward to do so:

```
birds$Effort <- birds$Effort * birds$repeats
```

#### 4 Manipulating the robin data

robins <- birds[birds\$species=="r", ]</pre>

For the purposes of this practical, we are interested in estimating the density of robins and so we select only these records:

**Question:** On how many transects were robins detected?

```
length(unique(robins$Sample.Label))
```

If we were to use the robins data as it is at present to estimate density, then density would be **incorrect** because the search effort associated with three transects is missing. Adding these missing transects to the robins data, requires several steps:

- 1. identify the missing transects,
- 2. select the information for the missing transects,
- 3. get the missing information in the correct format,
- 4. add the missing information to the robins data.

The following commands identifies the missing transects. After each command, type the name of the object which has been created to see what each command has done.

```
robin.lab <- unique(robins$Sample.Label)</pre>
miss.lab <- tran.lab[!is.element(el=tran.lab, set=robin.lab)]</pre>
```

To understand what the second command has done, unpeel it starting at its centre:

- elements of tran.lab are selected using []
- the is.element function (without the ! symbol) selects the elements in tran.lab, which are also in the set argument (i.e. robin.lab)
- the ! is used to select the elements in tran.lab that are NOT in robin.lab.

Now we know which transects are missing, we can select these records from the birds data frame:

```
miss.data <- birds[is.element(birds$Sample.Label, miss.lab), ]</pre>
```

However, the information about the transects are repeated in this new data frame because we have just selected all records for these transects. A quick check of the number of rows will confirm this:

```
length(miss.data$Sample.Label)
```

To get rid of rows where Sample. Label is duplicated use the command:

```
miss.data <- miss.data[!duplicated(miss.data$Sample.Label), ]</pre>
```

This command has selected the records from miss.data for which the transect label is not duplicated.

We only want to keep the information about search effort and so data in the distance, species and visit columns are set to missing:

```
miss.data$distance <- rep(NA, length(miss.lab))
miss.data$species <- rep("NA", length(miss.lab))</pre>
miss.data$visit <- rep(NA, length(miss.lab))</pre>
```

Examine miss.data.

The final thing to do is to add the missing data (miss.data) to the robins data frame using the rbind function (this combines data frames with the same columns).

```
robins <- rbind(robins, miss.data)</pre>
```

Let's see the result of all this manipulation:

```
tail(robins, n=4)
```

If we wanted to be very tidy, then the data frame could be sorted so that the transect labels were in order:

```
robins <- robins[order(robins$Sample.Label), ]</pre>
```

#### 5 Analysis

Before we fit any models, have a quick look at the histogram of distances:

```
hist(robins$distance, breaks=20)
```

Consistent with Buckland et al. (2015), three detection functions are fitted using the ds() function in the R package Distance (Miller, 2017):

**Question:** What is the preferred model for the robin data?

```
summarize_ds_models(robin.hn.herm, robin.uni.cos, robin.haz.simp)
```

## 6 Examining the dht object

The fitted model object (e.g. robin.uni.cos) is made up of two parts; the detection function in the ddf part and the estimates in the dht part. In this section, we look at the dht part.

To list the elements that are contained in dht, use the names function:

```
names(robin.uni.cos$dht)
```

Detections were of individual birds and so group size was not included in these data - if it had been included (in a column called size), then as well as individuals there would have been elements clusters and Expected.S.

The estimates stored in the individuals object can be listed in a similar manner:

```
names(robin.uni.cos$dht$individuals)
```

To collect together the density estimates (and estimates of precision) from all the fitted models, we can use the following command:

Question: Examine the three sets of density estimates to see if the previous suggestion (that the density estimates are similar) is confirmed

#### **Goodness of fit**

Here we look at goodness of fit test with unequal bin intervals and just consider one of the fitted models. First we specify the required bin intervals.

```
robin.brks \leftarrow c(0, 12.5, 22.5, 32.5, 42.5, 52.5, 62.5, 77.5, 95.0)
```

Perform the tests using both exact distance data for the Cramer-von Mises test and specified breakpoints for  $\chi^2$  test for the uniform-cosine model that had the (slightly) smallest AIC score.

```
gof_ds(robin.uni.cos, breaks=robin.brks, chisq = TRUE,
       main="QQ plot unif-cos for robins")
```

## **Customising the detection function plot**

The plot function provides a basic plot of the fitted detection function overlaid onto the scaled distribution of distances:

```
plot(robin.uni.cos)
```

However, the plot can be customised for reporting:

```
plot(robin.uni.cos, showpoints=FALSE, black.white=TRUE, pl.den=50,
     lwd=2, breaks=robin.brks, main="Uniform-cosine", xlab="Distance (m)")
```

The arguments are:

- showpoints logical indicating whether observed distances are shown
- lwd line width (1=default)
- pl.den density of shading of histogram (0=no shading)

For other options see help(plot.ds) (Note plot is a generic function which selects a relevant type of plot based the object).

## 9 Advanced: modularising R code to work with multiple species

When analysing a multi-species survey, it is likely that the invesigator will want to analyse all (or at least many) of the species encountered during the survey. This will necessitate some repetitive calculation, such as accounting for transects without detections and fitting multiple detection functions for each species.

To facilitate the repetitive nature of such analyses, it is useful to take advantage of the programmatic nature of the R language to create functions that can be called repeatedly with arguments to accommodate changing circumstances. The code below demonstrates such a modular approach whereby two functions augment.empty.transects() and fit.hn.uni.haz() are defined to aide in the repeated analyses.

The first function augment.empty.transects() performs the data manipulation described in Section 4, with two arguments: the data frame containing the full survey data and the species code on which to subset the

```
augment.empty.transects <- function(survey, species) {</pre>
  # Purpose: find transects on which species not detected
              adjust data file to correct effort
  # Input: raw data file, species on which to subset data
  # Output: data frame with correct effort for model fitting
  # Rexstad August 2018
  num.transects <- length(unique(survey$Sample.Label))</pre>
  holetab <- as.matrix(table(survey$species, survey$Sample.Label))==0
  holes <- which(holetab, arr.ind = TRUE)
  if (length(holes[rownames(holes)==species])==0) {
    adj.survey <- survey[survey$species==species, ]</pre>
  } else {
    alltranlen <- vector(mode="numeric", length=num.transects)</pre>
    for (i in 1:num.transects) {
      alltranlen[i] <- survey$Effort[survey$Sample.Label==i][1]
    }
    empty.transects <- NULL
    for (i in 1:length(holes[rownames(holes)==species,2])) {
      empty.label <- holes[rownames(holes)==species,2][i]</pre>
      empty.length <- alltranlen[holes[rownames(holes)==species,2]][i]</pre>
      empty.record <- cbind(survey[1,1:3], empty.label, empty.length)</pre>
      empty.transects <- rbind(empty.transects, empty.record)</pre>
    }
    empty.transects[ ,c("a", "b", "c")] <- NA</pre>
    names(empty.transects) <- names(survey)</pre>
    adj.survey <- survey[survey$species==species, ]</pre>
    adj.survey <- rbind(adj.survey, empty.transects)</pre>
    adj.survey <- adj.survey[order(adj.survey$Sample.Label), ]
  }
  return(adj.survey)
}
```

The second function, fit.hn.uni.haz() fits three candidate models to a dataset provided as the first argument. The second argument is the truncation distance. The final argument determines whether the summarize\_ds\_models() table is printed.

```
fit.hn.uni.haz <- function(data, trunc, print=TRUE) {</pre>
  # Purpose: fit three key functions to transect data,
              perform model selection and
               print model selection table
  # Input: data to analyse, truncation distance, print flag
  # Output: fitted model object (class 'dsmodel')
  # Rexstad August 2018
  hn.herm <- ds(data, trun=trunc, key="hn", adj="herm", con=.1)
```

```
uni.cos <- ds(data, trun=trunc, key="unif", adj="cos", con=.1)
 haz.simp <- ds(data, trun=trunc, key="hr", adj="poly", con=.1)
 mods <- summarize_ds_models(hn.herm, uni.cos, haz.simp, output="plain")</pre>
 if(print) print(knitr::kable(mods))
 names(mods) <- c("mod","key","form","fit","pa","sepa","daic")</pre>
 if(mods[1,1] == "hn.herm") {
    result <- hn.herm
 } else {
   if(mods[1,1]=="uni.cos") {
    result <- uni.cos
 } else {
    result <- haz.simp
 }
 }
 return(result)
}
```

The two functions are used in tandem in the calling code below. Note the for loop that iterates through three of the four species detected in the Montrave survey (great tit not analysed because there were few detections).

```
for(species in c("r", "c", "w")) {
  best.model <- fit.hn.uni.haz(augment.empty.transects(birds, species),</pre>
                                100, print=FALSE)
  plot(best.model,
       main=paste("Montrave lines, species ", species,
       "\nD-hat=", round(best.model$dht$individuals$D$Estimate,4),
       "SE=", round(best.model$dht$individuals$D$se, 4)))
}
```

#### References

https://www.R-project.org/.

```
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```