Rdistance Tutorial for Beginners

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April 16, 2015

This tutorial is meant to be a beginner's guide to using Rdistance. It is assumed that you have some familiarity with using Program R, but not necessarily with distance-sampling analysis. This beginning tutorial focuses on input data requirements, fitting a detection function, and estimating abundance (or density). Here, we make use of the example datasets already contained within Rdistance (i.e., line transect surveys of sparrows), so you can complete this tutorial without having any data of your own. This tutorial is current as of version 1.2.2 of Rdistance.

1: Install and load Rdistance

If you haven't already done so, install the latest version of Rdistance. In the R console, issue install.packages("Rdistance"). After the package is installed, it can be loaded into the current session as follows:

require(Rdistance)

```
## Loading required package: Rdistance
## Rdistance (version 1.2.2)
```

2: Read in input data

Rdistance requires two input datasets. These can be prepared outside of R and read in as data.frames using, for example, read.csv. In the following sections, we make use of the sparrow example datasets already contained within Rdistance.

The first required dataset is a detection data.frame, with a row for each detection, and the following required columns, named as follows:

- siteID = Factor, the site or transect where the detection was made.
- groupsize = Numeric, the number of individuals within the detected group.
- dist = Numeric, the perpendicular distance (also known as off-transect distance) from the transect to the detected group.

If the observers recorded sighting distance and sighting angle instead of perpendicular distance (as is often common in line transect surveys), you can use the perpendicular distances based on the sighting distances and sighting angles.

The second required dataset is a transect data.frame, with a row for each transect surveyed, and the following required columns, named as follows:

- siteID = Factor, the site or transect surveyed.
- length = Numeric, the length of the transect. Use the same units as the detection distances.
- ... = Any additional transect-level covariate columns.

3: Fit a detection function

After prepping the input data, the first step is to explore your data and fit a detection function. First, load the example dataset of sparrow detections and their distances from the package using data(sparrow.detections). Be sure that you have installed and loaded Rdistance prior to issuing the following commands:

```
data(sparrow.detections)
head(sparrow.detections)
```

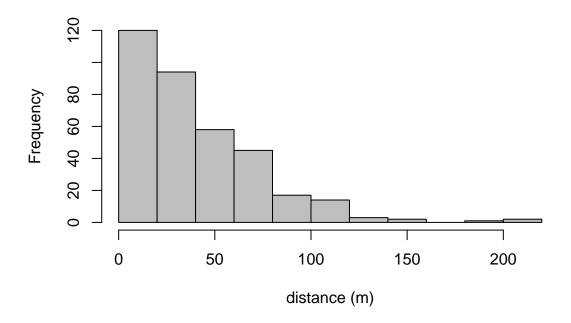
```
##
     siteID groupsize sightdist sightangle
## 1
                     1
         A1
                               65
## 2
                     1
                               70
         Α1
                                           10
## 3
         A1
                     1
                               25
                                           75
## 4
                     1
                               40
                                            5
         Α1
                               70
                                           85
## 5
         Α1
                     1
## 6
                     1
                                           90
         Α1
                               10
```

Note that the observers recorded sighting distances and sighting angles. Use the perp.dists function to calculate the perpendicular distance from each detected group to the transect, then remove the sightdist and sightangle columns.

```
##
     siteID groupsize dist
## 1
                     1 16.8
         Α1
## 2
                     1 12.2
         Α1
                     1 24.1
## 3
         Α1
## 4
         A1
                     1 3.5
                     1 69.7
## 5
         Α1
## 6
         A1
                     1 10.0
```

Explore the distribution of distances.

```
hist(sparrow.detections$dist, col="grey", main="", xlab="distance (m)")
```



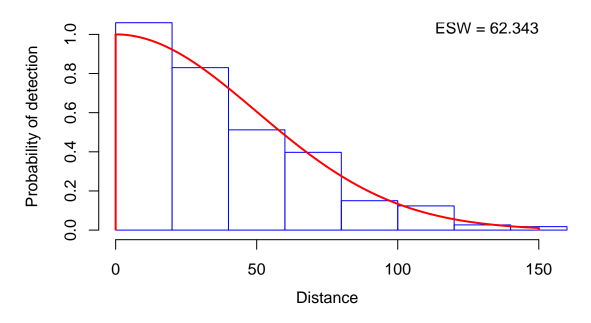
summary(sparrow.detections\$dist)

```
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 0.00 14.15 30.75 39.64 57.35 207.00
```

Next, fit a detection function (plotted as a red line) using F.dfunc.estim. For now, we will proceed using the half-normal likelihood as the detection function, but in Section 5 of this tutorial, we demonstrate how to run an automated process that fits multiple detection functions and compares them using AICc. Note that distances greater than 150 m are quite sparse, so here we right-truncate the data, tossing out detections where dist > 150.

```
dfunc <- F.dfunc.estim(sparrow.detections, likelihood="halfnorm", w.hi=150)
plot(dfunc)</pre>
```

halfnorm, 0 expansions



dfunc

Log likelihood: 1630.716

AIC: 3263.443

The effective strip width (ESW) is the key information from the detection function that will be used to next estimate abundance (or density). The ESW is calculated by integrating under the detection function. A survey with imperfect detection and ESW equal to X effectively covers the same area as a study with perfect

4: Estimate abundance given the detection function

detection out to a distance of X. See the help documentation for ESW for details.

Estimating abundance requires the additional information contained in the second required dataset, described earlier, where each row reprensents one transect. Load the example dataset of surveyed sparrow transects from the package.

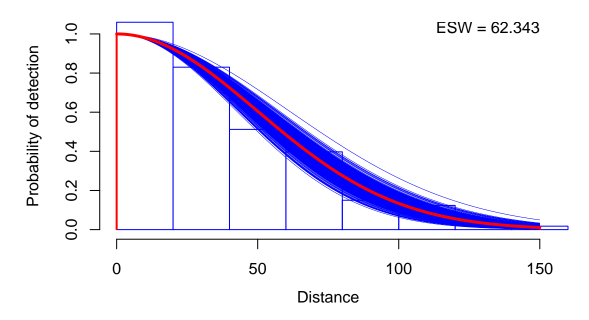
```
data(sparrow.transects)
head(sparrow.transects)
```

```
##
     siteID length sage
## 1
          A1
                500 High
## 2
          A2
                500 High
## 3
          ΑЗ
                500 High
## 4
          A4
                500 High
          B1
                500 High
## 6
                500 High
          B2
```

Next, estimate abundance (or density in this case) using F.abund.estim. If area=1, then density is given in the squared units of the distance measurements – in this case, sparrows per square meter. Instead, we set area=10000 in order to convert to sparrows per hectare (1 ha == 10,000 m²). The equation used to calculate the abundance estimate is detailed in the help documentation for F.abund.estim.

Confidence intervals for abundance are calculated using a bias-corrected bootstrapping method (see F.abund.estim), and the detection function fit in each iteration of the bootstrap is plotted as a blue line (if plot.bs=TRUE). Note that, as with all bootstrapping procedures, there may be slight differences in the confidence intervals between runs due to so-called 'simulation slop'. Increasing the number of bootstrap iterations (R = 500 used here) may be necessary to stabilize CI estimates.

halfnorm, 0 expansions



Computing bootstrap confidence interval on $\mathbb{N}\dots$

```
fit
```

```
## Call: F.dfunc.estim(dist = sparrow.detections, likelihood = "halfnorm",
                                                                                w.hi = 150)
##
## Coefficients:
      Sigma
##
## 49.87415
##
## Convergence: Success
## Function: HALFNORM
## Strip: 0 to 150
## Effective strip width: 62.34334
## Scaling: g(0) = 1
## Log likelihood: 1630.716
## AIC: 3263.443
##
## Abundance estimate: 0.8265162; 95% CI=(0.6610107 to 1.010256)
```

Results of interest (such as the abundance estimate and confidence interval) can be extracted from the resulting object (here called fit).

```
fit$n.hat

## [1] 0.8265162

fit$ci

## 1.921159% 96.78156%
## 0.6610107 1.0102564
```

5: Use AICc to select a detection function and estimate abundance

Alternatively, steps 3 (fitting a detection function) and 4 (estimating abundance) can be automated using the function F.automated.CDA. This function attempts to fit multiple detection functions, uses AICc (by default, but see help documentation for AIC.dfunc for other options) to find the 'best' detection function, then proceeds to estimate abundance using that detection function. By default, F.automated.CDA tries dozens of detection functions, but you can restrict the process to fewer detection functions if you choose (see help documentation for F.automated.CDA). Specifying plot=TRUE would return a plot of each detection function. In this example, we won't restrict the number of detection functions attempted, and we won't plot each (plot=FALSE).

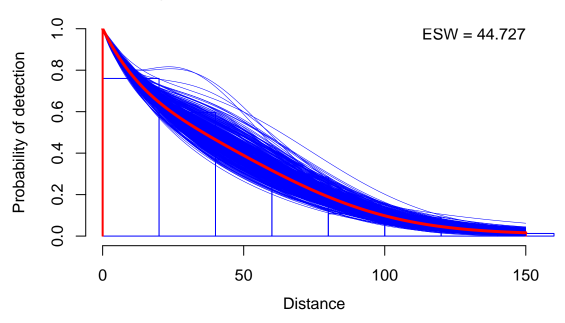
```
auto <- F.automated.CDA(detection.data=sparrow.detections, transect.data=sparrow.transects, w.hi=150, plot=FALSE, area=10000, R=500, ci=0.95, plot.bs=TRUE)
```

```
## Likelihood
                                Converged? Scale?
                Series
                        Expans
## halfnorm cosine
                        Yes
                                0k 3263.4429
## halfnorm cosine
                                0k
                                    3261.585
                        Yes
## halfnorm hermite 1
                        Yes
                                Πk
                                    3262.8066
                                    3262.1526
## halfnorm simple
                        Yes
## halfnorm cosine
                                0k
                                    3263.3155
                        Yes
```

```
## halfnorm hermite 2
                               0k
                                   3263.593
                       Yes
## halfnorm simple 2
                               0k
                                   3263.4741
                       Yes
## halfnorm cosine 3
                       Yes
                               0k
                                   3265.1285
## halfnorm hermite 3
                                   3265.5499
                       Yes
                               Ok
## halfnorm simple 3
                       Yes
                               0k
                                   3265.216
## hazrate
               cosine 0
                           Yes
                                   0k 3267.6246
## hazrate
               cosine 1
                           Yes
                                   Ok 3263.3092
## hazrate
                           No
                                   NA NA
               hermite 1
## hazrate
               simple 1
                           No
                                   NA
                                       NA
## hazrate
               cosine 2
                           Yes
                                   0k 3265.0704
## hazrate
               hermite 2
                           Yes
                                   Ok 3268.9682
                                   0k 3265.5796
## hazrate
               simple 2
                           Yes
                                   0k 3267.117
## hazrate
               cosine 3
                           Yes
## hazrate
               hermite 3
                           Bad
                                   NA NA
## hazrate
               simple 3
                           No
                                   NA NA
## uniform
               cosine 0
                           Yes
                                   0k
                                       3260.7318
## uniform
               cosine 1
                           {\tt Bad}
                                   NA
                                       NA
## uniform
               hermite 1
                           Yes
                                   0k 3262.736
                                   0k 3262.2602
## uniform
               simple 1
                           Yes
               cosine 2
## uniform
                           Bad
                                   NA NA
## uniform
               hermite 2
                           No
                                   NA NA
## uniform
               simple 2
                           Bad
                                   NA NA
## uniform
                                   NA NA
               cosine 3
                           Bad
## uniform
               hermite 3
                           No
                                   NA
                                       NA
## uniform
               simple 3
                           Bad
                                   NA NA
## negexp
               cosine 0
                           Yes
                                   Ok 3263.8153
## negexp
                cosine 1
                           Yes
                                   0k 3260.0985
## negexp
               hermite 1
                           Yes
                                   0k
                                       3260.3108
                                   0k 3261.0097
## negexp
               simple 1
                           Yes
                                   Ok 3262.0306
## negexp
               cosine 2
                           Yes
## negexp
               hermite 2
                           No
                                   NA
                                       NA
## negexp
               simple 2
                           Yes
                                   Ok
                                       3262.2878
                                   Ok 3263.4898
## negexp
                cosine 3
                           Yes
               hermite 3
                                   NA NA
## negexp
                           No
                                   0k 3264.2055
## negexp
                simple 3
                           Yes
## Gamma
                   0
                       Yes
                               0k 3586.8919
```

Note: Some models did not converge or had parameters at their boundaries.

negexp, cosine expansion, 1 expansions



```
## Computing bootstrap confidence interval on N...
##
##
  ----- Final Automated CDS Abundance Estimate -----
  Call: F.dfunc.estim(dist = dist, likelihood = fit.table$like[1], w.lo = w.lo,
                                                                                  w.hi = w.hi, expan
##
##
  Coefficients:
##
         Beta
   0.02754839
               -0.26542176
##
## Convergence: Success
## Function: NEGEXP with 1 expansion(s) of COSINE series
## Strip: 0 to 150
## Effective strip width: 44.72749
## Scaling: g(0) = 1
## Log likelihood: 1628.032
## AIC: 3260.098
##
## Abundance estimate: 1.152038; 95% CI=(0.8563271 to 1.494071)
```

You can see that the detection function with the lowest AICc value (and thus selected as the 'best') is the negative exponential likelihood, with one cosine expansion.

6: Conclusion

Note that the detection function that you select has a large influence on the resulting abundance estimate. In sections 3 and 4, we fit a half-normal detection function and used that function to estimate sparrow density.

Our estimate was 0.83 sparrows per ha (95% CI = 0.66 - 1.01). In section 5, we used AICc to determine the best-fitting detection function and used that function to estimate sparrow density again. Our new estimate was 1.15 sparrows per ha (95% CI = 0.86 - 1.49). (Note, recall that your estimates may vary slightly from these due to minor 'simulation slop' inherent in bootstrapping methods). Thus we see that choosing an appropriate detection function is critical to accurately estimating abundance. The F.automated.CDA function can help you select a detection function that fits your data well.

That concludes this Rdistance tutorial. You are now ready to read in your own data, fit a detection function, and estimate abundance.