secrdesign - sampling design for spatially explicit capture—recapture

Murray Efford 2014-11-27

Contents

Introduction	2	
A simple example	3	
Defining scenarios		
Detector layouts	4	
The scenarios dataframe	4	
Running simulations	6	
Arguments of run.scenarios()	6	
Customising run.scenarios()	7	
More on masks	8	
Summarising simulation output	9	
Extracting estimate tables from fitted models	9	
Choosing the statistics to summarise	10	
Disposing of rogue values	12	
Summary method	12	
Plot method	14	
Additional topics	15	
Parallel processing	15	
Shortcut evaluation of precision	16	
Non-uniform populations	17	
Linear habitat	17	
Splitting data generation and model fitting	18	
Populations with sub-classes or multiple sessions	18	
Limitations, tips and troubleshooting	19	
References	20	

A	Appendix. Examples			
	Multiple grids, varying number of occasions	21		
	Learned trap response	23		
	Non-uniform possums	27		
	Code for linear habitat	30		
	Grouped populations	32		

The R package **secrdesign** is a set of tools to assist the design of studies using spatially explicit capture–recapture (SECR). It provides convenient wrappers for simulation and model fitting functions in package **secr** to emulate many features of the 'Simulator' module of Density 5.0 (Efford 2012). Other tools may be added in future.

This document is a technical guide to **secrdesign**. It assumes an understanding of estimator properties such as bias, precision, and confidence interval coverage, and the use of Monte Carlo simulation to predict the frequentist performance of different sampling designs. Using **secrdesign** can be daunting because it allows for many different combinations of data generation, model fitting and summary statistics. Several examples are given to indicate the range of possibilities.

Introduction

When designing a study we assume you have in mind –

- (i) a population parameter you want to measure (probably density or population size),
- (ii) one or more design variables over which you have some control (number and spacing of detectors, number of sampling occasions etc.),
- (iii) some pilot data, or parameter estimates from published studies, and
- (iv) a criterion by which to evaluate different designs. This is most likely the precision of the estimates, as this in turn determines your ability (power) to recognise changes. Cost or effort may be an explicit criterion, or the designs may be constructed to allocate constant effort in different ways.

We use 'relative standard error' (RSE) for the relative precision of an estimate. This is sometimes called the coefficient of variation (CV) of the estimate, but RSE is more appropriate. We use 'accuracy' in the sense of Williams et al. (2002 p.45) and other authors from the United States. Accuracy combines both systematic error (bias) and precision: one measure is the square root of the mean squared difference between the true value and the estimate (RMSE).

Once you have sorted out (i)–(iv) you can proceed to use Monte Carlo simulation in **secrdesign** to evaluate how alternative sampling scenarios perform with respect to your chosen criterion.

Fig. 1 shows the sequence of steps taken in **secrdesign** to conduct simulations and summarise the results. Each step is described in detail in a later section. Simulations are specified by 'scenarios' stored in a dataframe. The scenario dataframe will usually be constructed with make.scenarios. You may construct it manually, but there is a rigid list of required columns (see The scenarios dataframe).

Typically, you will (i) construct detector array(s), (ii) construct a dataframe of scenarios, (iii) use run.scenarios() to generate data and fit SECR models, (iv) select some statistics with select.stats(), and (v) summarise and plot the results.

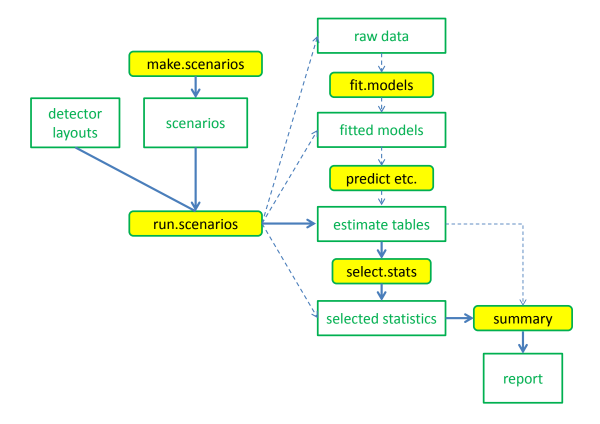


Fig. 1. Core functions in **secrdesign** (yellow) and their main inputs and outputs. Output from the simulation function run.scenarios() may be saved as whole fitted models, predicted values (parameter estimates), or selected statistics. Each form of output requires different subsequent handling. The default path is shown by solid blue arrows.

A simple example

For an introductory example we construct a simple set of scenarios and perform some simulations. The trap layout is a default 6×6 grid of multi-catch traps at 20-m spacing. Density takes one of two levels (5/ha or 10/ha) and detection parameters sigma and g0 are fixed.

The output is an object of class c("estimatetables", "secrdesign", "list"). We use the summary method for estimatetables to view results, and here display only the summary output (omitting a header that describes the simulations).

```
summary(sims1)$0UTPUT
```

```
## $`D = 5`
```

```
##
                      mean
               n
## estimate
               50
                  5.23767 0.24994
## SE.estimate 50
                  1.84409 0.05958
               50 2.70152 0.16476
## lcl
## ucl
               50 10.28534 0.37589
               50 0.04753 0.04999
## RB
## RSE
               50 0.36890 0.00962
## COV
               50 0.92000 0.03876
##
## $`D = 10`
##
                      mean
                                se
                n
               50
                   9.58795 0.36972
## estimate
## SE.estimate 50
                   2.45915 0.05721
## lcl
               50 5.86410 0.27453
## ucl
               50 15.75842 0.47775
## RB
               50 -0.04121 0.03697
## RSE
                  0.26727 0.00668
               50
## COV
               50
                  0.92000 0.03876
```

Later sections show how to customize the summary and plot results.

Defining scenarios

Detector layouts

Detector layouts are specified as **secr** 'traps' objects. These may be input from text files using **read.traps** or constructed according to a particular geometry and spacing with functions such as **make.grid**, **make.circle**, **make.systematic** or **trap.builder**. See the help files for these **secr** functions for further details. The detector type (multi-catch trap, proximity detector etc.) is stored as an attribute of each 'traps' object, which may also include detector-level covariates and detector 'usage' by occasion.

Multiple layouts are combined in a single list object; component names ('grid6x6' etc.) will be used to annotate the output.

This creates square grids with the default detector type 'multi' and default spacing 20 m. See ?secr::make.grid for other options.

The scenarios dataframe

The function make.scenarios() constructs a dataframe in which each row defines a simulation scenario. Its arguments (with defaults) are:

```
make.scenarios (trapsindex = 1, noccasions = 3, nrepeats = 1, D, g0,
    sigma, lambda0, detectfn = 0, recapfactor = 1, popindex = 1,
    detindex = 1, fitindex = 1, groups, crosstraps = TRUE)
```

Each argument except for 'groups' and 'crosstraps' may be used to specify a range of values for a parameter. Four ('trapsindex', 'popindex', 'detindex', 'fitindex') are actually surrogate numerical indices; the index is used to select one component from a list of possibilities later provided as input to run.scenarios().

By default, a scenario is formed from each unique combination of the input values (trapsindex, noccasions, nrepeats, D, g0, sigma, lambda0, detectfn, recapfactor, popindex, and fitindex) using expand.grid. For example,

```
make.scenarios (trapsindex = 1:3, noccasions = 4, D = 5, g0 = 0.2, sigma = c(20,30))
```

```
##
     scenario trapsindex noccasions nrepeats D gO sigma detectfn recapfactor popindex
## 1
                                                 1 5 0.2
                                                              20
                                                                         0
             1
                          1
                                       4
                                                                                       1
                                                                                                 1
             2
                          2
                                       4
                                                                         0
## 2
                                                 1 5 0.2
                                                              20
                                                                                       1
                                                                                                 1
                          3
             3
                                       4
                                                                         0
## 3
                                                 1 5 0.2
                                                              20
                                                                                       1
                                                                                                 1
## 4
             4
                          1
                                       4
                                                 1 5 0.2
                                                              30
                                                                         0
                                                                                       1
                                                                                                 1
                          2
## 5
             5
                                       4
                                                 1 5 0.2
                                                              30
                                                                         0
                                                                                       1
                                                                                                 1
## 6
             6
                          3
                                                 1 5 0.2
                                                              30
                                                                         0
                                                                                       1
                                                                                                 1
##
     detindex fitindex
## 1
             1
                        1
## 2
             1
                        1
## 3
             1
                        1
## 4
             1
                        1
## 5
             1
                        1
## 6
```

In this case three different grids (possibly differing in number of traps) are trapped for the same number of occasions. A more interesting possibility is to vary the number of occasions inversely with the number of traps. However, if we naively set e.g., noccasions = c(8, 4, 2), this would generate all combinations of grid and number of occasions (18 different scenarios).

The alternative is to set crosstraps = FALSE. Then the vectors 'trapsindex', 'noccasions', and 'nrepeats' are locked together (if fewer values are provided in one of the vectors then it is re-used as required), and only the combination is crossed with the remaining parameter scenarios:

```
make.scenarios (trapsindex = 1:3, noccasions = c(8,4,2), D = 5, g0 = 0.2, sigma = c(20,30), crosstraps = FALSE)
```

```
scenario trapsindex noccasions nrepeats D gO sigma detectfn recapfactor popindex
##
## 1
             1
                          1
                                       8
                                                 1 5 0.2
                                                             20
                                                                         0
## 2
                          2
                                       4
                                                                         0
                                                                                       1
             2
                                                 1 5 0.2
                                                              20
                                                                                                 1
## 3
             3
                          3
                                       2
                                                 1 5 0.2
                                                              20
                                                                         0
                                                                                       1
                                                                                                 1
## 4
             4
                          1
                                       8
                                                 1 5 0.2
                                                              30
                                                                         0
                                                                                       1
                                                                                                 1
                          2
             5
## 5
                                       4
                                                 1 5 0.2
                                                              30
                                                                         0
                                                                                       1
                                                                                                 1
             6
                          3
                                       2
                                                 1 5 0.2
                                                                         0
## 6
                                                             30
                                                                                       1
                                                                                                 1
##
     detindex fitindex
## 1
             1
                        1
## 2
             1
                        1
## 3
             1
                        1
## 4
             1
                        1
## 5
             1
                        1
             1
## 6
                        1
```

All arguments except 'fitindex' control the generation of data. Note that 'g0' and 'lambda0' are alternatives: use the one appropriate to the detection function specified with detectfn' (see?secr::detectfn' for codes).

'D' is omitted if an inhomogeneous Poisson distribution is specified using a mask covariate (see Non-uniform populations). D is in animals / hectare (1 ha = 0.01km^2) and sigma is in metres, as in secr.

The 'nrepeats' column refers to the number of notional independent replicates of the particular detector layout. Notional replicates are simulated by (invisibly) multiplying density (D) by this factor, and ultimately dividing it into the estimate. Think of 5 grids of automatic cameras so widely separated that no animal moves between the grids. As detections of different animals are ordinarily modelled as independent, the entire design is equivalent to 5 times the density of animals interacting with one grid. This breaks down, of course, if animals compete for traps (as with single-catch traps), and should not be used in that case except as a rough approximation.

Just as trapsindex serves as a placeholder for entire detector arrays, popindex, detindex and fitindex tell run.scenarios() which set of arguments to select from pop.args, det.args and fit.args for sim.popn, sim.capthist and secr.fit respectively. These are for more advanced use: you may not need them.

When a vector of group identifiers is provided in 'groups', the population in each scenario is a set of independently sampled groups, each defined on a separate row. Groups are initially assigned the same parameter values and other settings: it is up to the user to insert group-specific values (example at Grouped populations).

Running simulations

The function run.scenarios() generates multiple datasets and, if requested, fits an SECR model to each one. In this section we describe its main arguments, with additional detail on habitat masks and customizing the output.

Arguments of run.scenarios()

```
run.scenarios (nrepl, scenarios, trapset, maskset, xsigma = 4,
    nx = 32, pop.args, det.args, fit = FALSE, fit.args, extractfn =
    NULL, multisession = FALSE, ncores = 1, seed = 123)
```

nrepl determines the number of replicate simulations. Make this large enough that the summary statistics have enough precision to answer your question. This is usually a matter for experimentation, remembering that precision (SE) is proportional to the square root of nrepl.

scenarios is the dataframe constructed with make.scenarios() as described in the last section.

trapset is a single 'traps' object or a list of traps objects, as described in 'Detector layouts' above.

maskset is an optional set of habitat masks, usually one per detector layout. If not specified, then masks will be constructed 'on the fly' using secr::make.mask with a 'buffer' of width xsigma × scenarios\$sigma and nx cells in the x dimension.

pop.args provides additional control over sim.popn (see ?secr::sim.popn for more). You may wish, for example, to set pop.args = list(Ndist = "fixed") to override the default of Poisson variation in the total number of simulated animals.

det.args provides additional control over sim.capthist (see ?secr::sim.capthist for more). One use is to
retain the simulated population as an attribute of the capthist object by setting det.args = list(savepopn
= TRUE); another is to set the binomN argument for count detectors. The sim.capthist arguments traps,
popn, detectpar, detectfn and noccasions are defined in the scenario or pop.args and cannot be overridden
by setting det.args.

Use fit = FALSE to generate and summarise detection data without fitting SECR models. This lets you check that your scenarios result in reasonable numbers of detected individuals (n), detections (ndet), and movements (nmov), before launching a full-blown simulation.

fit.args lets you specify how SECR models will be fitted to the simulated data. Most default arguments of secr.fit may be overridden by including them in fit.args. For example, to specify a negative exponential detection function use fit.args = list(detectfn = "EX"). If you wish to compare nspec different model specifications then fit.args should be a list of lists, one per specification, with fitindex taking values in the range 1:nspec.

The use of multisession and ncores is discussed under Additional topics.

Customising run.scenarios()

The output from run.scenarios() is controlled by its argument extractfn. This is a short function that is applied either (i) to each simulated raw dataset (capthist object) (fit = FALSE), or (ii) to each fitted model (fit = TRUE).

The default (builtin) extractfn (below) behaves appropriately for either data type. It is mostly concerned with summarising raw counts when fit = FALSE. A dataframe with no rows is returned when a model fails to fit.

```
extractfn <- function(x) {</pre>
    if (inherits(x, 'capthist')) {
        ## summarised raw data
        counts <- function(CH) {</pre>
            ## for single-session CH
            nmoves <- sum(unlist(sapply(moves(CH), function(y) y>0)))
            ## detectors per animal
            dpa <- if (length(dim(CH)) == 2)</pre>
                mean(apply(abs(CH), 1, function(y) length(unique(y[y>0]))))
                mean(apply(apply(abs(CH), c(1,3), sum)>0, 1, sum))
            c(n=nrow(CH), ndet=sum(abs(CH)>0), nmov=nmoves, dpa = dpa)
        }
        if (ms(x))
            unlist(lapply(x, counts))
        else {
            gp <- covariates(x)$group</pre>
            if (is.null(gp))
                counts(x)
            else
                unlist(lapply(split(x,gp,dropnullocc=TRUE), counts))
        }
    }
    else if (inherits(x,'secr') & (!is.null(x$fit)))
        ## fitted model:
        ## default predictions of 'real' parameters
        predict(x)
    else
        ## null output: dataframe of 0 rows and 0 columns
        data.frame()
}
```

When fit = FALSE, the default output from each replicate is a vector of 4 summary statistics:

- n number of different individuals
- ndet total number of detections (captures' andrecaptures')
- nmov total number of detections at a detector other than the one where an animal was last detected
- dpa detectors per animal (average number of detectors at which each animal was recorded)

When fit = TRUE, the default output from each replicate is the result of applying predict to the fitted model, i.e. a dataframe of 'real' parameter estimates and their standard errors etc. (an empty dataframe is returned if model fitting fails). Nearly the same is achieved by setting extractfn = predict for the 'beta' coefficients set extractfn = coef. For a conditional likelihood fit it may be appropriate to set extractfn = derived. To focus on population size in the masked region, set extractfn = region.N.

The user may also choose to save the entire dataset (fit = FALSE) or the entire fitted model (secr object; fit = TRUE) for each replicate by setting extractfn = identity. For a large analysis there is a risk of exceeding memory limits in R, and saving everything is generally not a good idea. For most purposes it is sufficient to save a trimmed version of the fitted model extractfn = trim (note secr defines the function trim). However, the full model is needed for derived.SL or regionN.SL (see Extracting estimate tables from fitted models).

run.scenarios() sets the class¹ of its output to distinguish among fitted models ("fittedmodels"), estimate tables from predict, coef etc. ("estimatetables"), and numeric values ready for summarisation ("selectd-statistics"). Simulated data saved with fit = FALSE, extractfn = identity have class c("rawdata", "secrdesign", "list"). An attribute 'outputtype' is used to make finer distinctions among these types of output ("secrfit", "predicted", "coef", "capthist", or "numeric"). Output from extractfn = derived is treated as "predicted".

When fit = TRUE, analyses are performed with secr.fit. Other analyses may be specified by setting fit = FALSE and providing the analysis function as the value for extractfn. The function should accept a 'capthist' object as input. For example, conventional closed-population estimates may be obtained with

This applies various nonspatial estimators to simulated spatial samples. Named arguments of extractfn may be included (here, 'estimator'); these are used for all scenarios, unlike fit.args which may vary among scenarios. Summarisation of alternative analyses will usually require careful selection of 'parameter' and 'statistics' in select.stats (see Choosing the statistics to summarise).

An alternative is to write your own code along these lines:

```
sum1 <- function(out) {
    require(abind)
    ## collapse replicates to an array, omitting non-numeric column
    out <- do.call(abind, c(out, along = 3))[,-1,,drop = FALSE]
    ## convert array from character to numeric
    mode(out) <- "numeric"
    ## take the average over replicates (meaningless for some fields)
    apply(out, 1:2, mean, na.rm = TRUE)
    }
lapply(closedNsim$output, sum1)</pre>
```

More on masks

A habitat mask in **secr** is a raster representation of the region near the detectors in which the centres of detected animals may lie. This excludes both nearby non-habitat, and habitat that is so distant that it is

 $^{^{1}}$ the full class is actually c(x, "secrdesign"", "list") where x is as described.

implausible any animal centred there would reach a detector. It is often convenient to define a mask to include all cells whose centre is within a certain distance of a detector - the buffer radius.

Within **secrdesign**, a mask is used both when generating populations of animals with sim.popn and when fitting SECR models with secr.fit. The extent of the mask used to generate populations is important if you are concerned with population size (for example, if you set extractfn = region.N). Then the size of the region determines the true value of the parameter of interest (N), and influences its sampling variance. The extent of the mask is less critical when density is the parameter of interest.

The default behaviour of run.scenarios() is to use a concave buffer of width xsigma × sigma around the particular detector layout. The 'coarseness' of the mask is determined by nx; note that the default for run.scenarios (nx = 32) is coarser than the default for secr::make.mask (nx = 64). This makes for speed, and is fairly safe when the buffer width is well matched to the scale of movement (we know sigma, so the default buffer width is well-matched). The same mask is used both for generating populations and fitting models.

Users may specify their own masks in the 'maskset' list argument. If the number of masks in maskset is one or a number equal to the number of detector layouts, then a column 'maskindex' is added automatically to the scenarios dataframe (all 1, or equal to trapsindex, in the two cases). Otherwise, the user must have manually added a maskindex column to scenarios to clarify which mask should be used with which scenario.

Summarising simulation output

run.scenarios usually takes a long time to run, but having saved its output you can quickly extract and summarise the results in many different ways.

We saw in the previous section and Fig. 1 that the output from run.scenarios for each replicate may be a fitted model, a table of parameter estimates, or a numeric vector. Summarisation across replicates (the summary method) requires output in 'selected statistics' form, so each of the other forms must be processed first²

Look again at Fig. 1: you will see that the primary input to the summary function is in the form of selected statistics. A secondary route is to automatically extract statistics from estimates tables, as shown by the dashed line in Fig. 1 – we used this in A simple example. We now address how other forms of output from run.scenarios can be processed into 'selected statistics' form.

Extracting estimate tables from fitted models

The methods predict and coef for the 'fittedmodels' class and the function derived.SL are provided to extract estimates of 'real' parameters from each fitted model. These are direct analogues of predict, coef and derived in secr. Here, they apply across all replicates of all scenarios and return an object of class estimatetables. regionN.SL is another possibility.

In each case, the result is a dataframe or list of dataframes for each replicate. Rows correspond to estimated parameters (or 'R.N' and 'E.N' for regionN.SL) and columns to the respective estimates, standard errors, and confidence limits (with some variations).

The ... argument of the functions predict, coef, derived.SL and regionN.SL lets you pass arguments such as alpha to the corresponding secr function (e.g., predict.secr or derived).

We can skip this step for the output from our simple example as it is already in 'estimatetables' form.

²Processing happens silently using default settings of select.stats() when summary is applied directly to 'estimate tables' output.

Choosing the statistics to summarise

Given tabular output from predict() or derived.SL(), we must select replicate-specific numerical quantities for further summarisation.³ This is the role of select.stats(), which has arguments –

```
select.stats(object, parameter = "D", statistics)
```

The parameter of interest defaults to density ("D"); others such as "g0" or "sigma" may be substituted, so long as they appear in the input object. To check which parameters are available use

```
find.param(object)
```

The task of select.stats() is to reduce each replicate to a vector of numeric values - we can think of the result as a replicate × values matrix for each scenario (Fig. 2). A later step (see Summary method) computes statistics ('fields') such as mean and SE for each column in the matrix.

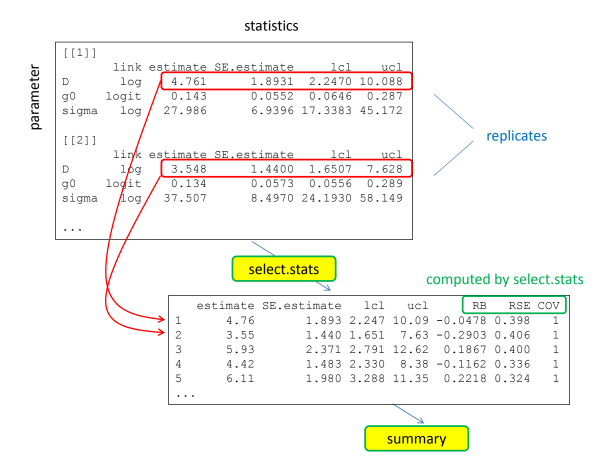


Fig. 2. Operation of select.stats for one scenario. Each replicate contributes one row to a replicates \times statistics matrix.

³If run.scenarios() has been used with fit = FALSE, then the output from each replicate is probably already in the form of selected statistics (the default raw data summaries 'n', 'ndet', 'nmov' and 'dpa') and select.stats() is not relevant. The same may also apply with a user-provided extractfn when fit = TRUE.

Here we describe the replicate-specific statistics that form the numeric vector. These may be simply 'estimate', 'SE.estimate', 'lcl' and 'ucl' as output from predict.secr. Additionally, when fit = TRUE, we can include statistics derived from the estimates of a parameter (Table 1). To describe these we use 'true' to stand for the known value of a real parameter, and 'estimate' for the estimate from a particular replicate.

Table 1. Computed statistics available in select.stats()

Statistic	Short name	Value
Relative bias ¹	RB	(estimate – true) / true
Relative SE^2	RSE	SE.estimate / estimate
Absolute deviation	ERR	abs(estimate - true)
Coverage indicator	COV	(estimate $>$ lcl) & (estimate $<$ ucl)

- 1. Also called 'normalised bias'
- 2. Also called 'coefficient of variation'

We use ('lcl', 'ucl') to represent a confidence interval for 'estimate'. Usually these are 95% intervals, but the level may be varied by setting the argument 'alpha' in predict (e.g., alpha = 0.1 for a 90% interval). Intervals from predict.secr are symmetrical on the link scale, and hence asymmetrical on the natural scale. Note also the argument loginterval in derived; the default loginterval = TRUE gives an asymmetrical interval on the natural scale.

The coverage indicator COV is a binary value (0/1); this becomes interesting later when averaged over a large number of replicates to give a coverage proportion. The absolute deviation ERR also comes into its own later as the basis for RMSE. In a sense the same is true of replicate-specific RB: RB should be reported only as an average over a large number of replicates.

Returning to our simple example, we apply select.stats() to focus on the density parameter "D".

```
stats1 <- select.stats(sims1, parameter = "D", statistics = c("estimate",
    "lcl", "ucl", "RB", "RSE", "COV"))
lapply(stats1$output, head, 4)</pre>
```

```
## $`1`
##
     estimate
                 lcl
                       ucl
                                 RB
                                       RSE COV
## 1
        4.563 2.320 8.976 -0.0873 0.3557
                                              1
## 2
        3.626 1.767 7.439 -0.2749 0.3794
                                              1
## 3
        3.861 1.838 8.109 -0.2279 0.3926
                                              1
## 4
        4.455 2.396 8.285 -0.1090 0.3246
                                              1
##
## $\2\
                                         RSE COV
##
     estimate
                  lcl
                        ucl
                                   RB
## 1
       16.330 11.211 23.79
                             0.63297 0.1937
## 2
               5.695 16.19 -0.03992 0.2713
        9.601
                                                1
## 3
       10.306
               6.417 16.55
                             0.03060 0.2453
                                                1
## 4
       12.629
               8.371 19.05 0.26288 0.2122
```

The two scenarios yield two replicates × statistic matrices, from which we display the first 4 rows.

Disposing of rogue values

Simulation output may contain rogue values due to idiosyncracies of model fitting. For example, nonidentifiability due to inadequate data can result in spurious extreme estimates of the sampling variance. The median (chosen as a field value in summary) is recommended as a robust alternative to the mean when there are some extreme estimates. Another way to deal with the problem is to set statistics to NA when a simulation fails. The function validate sets selected 'target' statistics to NA for replicates in which another test statistic is out-of-range or NA:

```
x <- validate (x, test, validrange = c(0, Inf), targets = test)
```

The permissable bounds are usually arbitrary, and the method should be used with care. The keyword "all" may be used for targets to indicate all columns.

validate accepts a selected statistics object (x) as input and returns a modified selected statistics object as output. See Learned trap response for an application.

Summary method

The summary method for 'selected statistics' objects reports both header information on the simulation scenarios and user-selected summaries of the pre-selected statistics.

```
summary (object, dec = 5, fields = c("n", "mean", "se"), alpha = 0.05,
type = c("list", "dataframe", "array"), ...)
```

Here the summary statistics are called 'fields' to distinguish them from the 'statistics' in each column of the numeric replicate × value matrix for each scenario (see Choosing the statistics to summarise). The task of the summary method is to compute the 'field' value for each 'statistic', summarising across replicates to give a 'statistic' × 'field' matrix for each scenario. The choice of 'fields' is shown in Table 2.

Table 2. Statistic fields available in the summary method for selected statistics objects.

Field	Description
n	number of non-missing values
mean	mean
se	standard error
sd	sample standard deviation
\min	minimum
max	maximum
lcl	lower $100(1-\alpha)$ % confidence limit
ucl	upper $100(1-\alpha)$ % confidence limit
${ m rms}$	root mean square
median	median
qxxx	xxx/1000 quantile
qyyy	yyy/1000 quantile

The summary fields 'lcl' and 'ucl' are for a simple Wald interval $(\hat{\mu} + z_{\alpha/2}\widehat{SE}(\hat{\mu}), \hat{\mu} + z_{1-\alpha/2}\widehat{SE}(\hat{\mu}))$ where z_{α} is the 100α -percentile of a standard normal distribution (e.g., $z_{0.975} = 1.96$). [Do not confuse these with the confidence limit statistics of the same name that are symmetrical only on the link scale].

Quantiles are specified as 'qxxx' and 'qyyy' where xxx and yyy are integers between 1 and 999 corresponding to quantiles 0.001 to 0.999. For example, 'q025' refers to the 2.5% quantile.

Applying the 'rms' field to the absolute deviation of an estimate (ERR) provides the root-mean-square-error 'RMSE'.

To recap – a summary value is reported for each combination of a selected statistic, computed for each replicate, and a 'field' that summarises the statistic across replicates, potentially resulting in a table with this structure:

Cells are left blank for combinations that are unlikely to be meaningful. 'rms' is useful with ERR (i.e. RMSE), but not when applied to other statistics. 'n', 'mean' and 'se' summarise the COV indicator, but other potential summaries are (almost) meaningless.

Apply this to the selected statistics from our simple example:

```
summary(stats1, c('n', 'mean', 'se', 'median'))
```

```
## run.scenarios(nrepl = 50, scenarios = scen1, trapset = traps1,
##
       fit = TRUE, seed = 345)
##
## Replicates
                 50
## Started
                 21:49:17 23 Nov 2014
## Run time
                 1.78 minutes
## Output class
                 selectedstatistics
##
## $constant
##
               value
## trapsindex
## noccasions
                    3
## nrepeats
                    1
                  0.2
## g0
## sigma
                   25
## detectfn
                    0
## recapfactor
                    1
## popindex
                    1
## detindex
## fitindex
                    1
## maskindex
##
```

```
## $varying
##
    scenario
             D
##
           1
             5
##
           2 10
##
## $detectors
    trapsindex trapsname
##
##
             1
                  traps1
##
## OUTPUT
   D = 5
##
                   mean
                                  median
## estimate 50
                5.23767 0.24994
                                 5.04535
## lcl
                                 2.50031
            50
               2.70152 0.16476
            50 10.28534 0.37589 10.01498
## ucl
## RB
            50
               0.04753 0.04999
                                 0.00907
## RSE
            50
               0.36890 0.00962
                                 0.36154
## COV
                0.92000 0.03876
                                 1.00000
##
## $`D = 10`
##
                                  median
             n
                   mean
                             se
## estimate 50
               9.58795 0.36972
                                 9.86434
            50 5.86410 0.27453 5.91509
## lcl
            50 15.75842 0.47775 16.14298
## ucl
            50 -0.04121 0.03697 -0.01357
## RB
## RSE
               0.26727 0.00668
                                 0.25726
## COV
            50 0.92000 0.03876 1.00000
```

Plot method

Use the plot method to visualize the distributions of 'selected statistics' that you have simulated. You may plot either (i) histograms of the selected statistics (type = "hist") or (ii) the estimate and confidence interval for each replicate (type = "CI"). One histogram is plotted for each combination of scenario and statistic—you may want to select a subset of scenarios and statistics, and use the graphics options mfcol or mfrow to control the layout. For type = "CI" the statistics must include 'estimate', 'lcl' and 'ucl' (or 'beta', 'lcl' and 'ucl' if outputtype = "coef").

```
par(mfrow = c(2,2))
plot(stats1, type = "hist", statistic = "estimate")
plot(stats1, type = "CI")
```

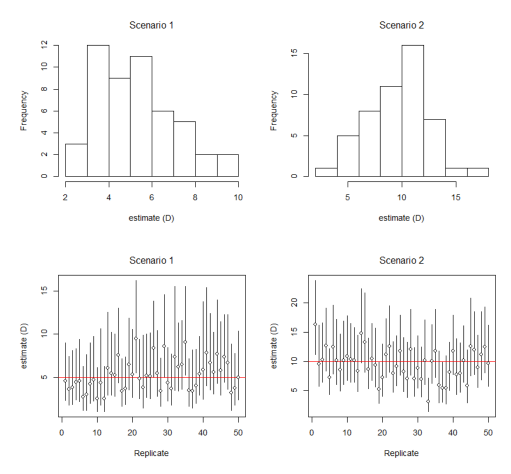


Fig. 3. Plot method applied to a 2-scenario 'selected statistics' object with type = "hist" (top) and type = "CI" (bottom)

Additional topics

Parallel processing

Setting ncores > 1 causes run.scenarios() to run separate scenarios on separate cores. This uses the R package parallel. Technically, it relies on Rscript, and communication between the master and worker processes is via sockets. As stated in the R parallel documentation "Users of Windows and Mac OS X may expect pop-up dialog boxes from the firewall asking if an R process should accept incoming connections". It appears to be safe to accept these.

Use parallel::detectCores() to get an idea of how many cores are available on your machine; this may (in Windows) include virtual cores over and above the number of physical cores. If you use the maximum available cores for run.scenarios() then expect any other processes on the machine to slow down!

Running one scenario per core is suboptimal if scenarios differ widely in how long they take to run: the system waits for the slowest. There is no way around this limitation in **secrdesign**.

Random number generation for multiple cores uses the "L'Ecuyer-CMRG" random number generator as described in ?RNG.

Shortcut evaluation of precision

The asymptotic variance (and hence RSE) of a maximum likelihood estimate is typically obtained from the curvature of the likelihood computed numerically at the fitted value of the parameter(s) (i.e., at the MLE). Fitting SECR models is slow. An alternative estimate of the RSE that is sufficient for most purposes may be got from the curvature of the likelihood computed at the known 'true' value(s) of the parameter(s). This is much faster as it does not require the model to be fitted.

secr.fit may be 'tricked' into providing this variance estimate by setting method = "none" and providing the true values as the start vector. run.scenarios() makes this easy by assuming that if you specify method = "none" you wish to use start = "true". However, this works only when there is a 1:1 relationship between 'beta' and 'real' parameters; it does not work when 'recapfactor' is specified.

```
sims2 <- run.scenarios(nrepl = 50, trapset = traps1, scenarios = scen1,
    fit = TRUE, fit.args = list(method = "none"))</pre>
```

summary(sims2)

```
## run.scenarios(nrepl = 50, scenarios = scen1, trapset = traps1,
##
       fit = TRUE, fit.args = list(method = "none"))
##
## Replicates
                 50
## Started
                 21:51:04 23 Nov 2014
## Run time
                 0.381 minutes
## Output class selectedstatistics
##
## $constant
##
               value
## trapsindex
                    1
## noccasions
                    3
## nrepeats
                    1
                  0.2
## g0
## sigma
                  25
## detectfn
                    0
## recapfactor
                    1
## popindex
## detindex
                    1
## fitindex
                    1
## maskindex
                    1
##
## $varying
##
    scenario D
##
           1
             5
##
           2 10
##
## $detectors
##
    trapsindex trapsname
##
             1
                  traps1
##
## $fit.args
    fitindex method
##
##
           1
               none
##
## OUTPUT
```

```
## $`D = 5`
##
                n
                      mean
                                 se
## estimate
                   5.0000 0.00000
               50
                    1.8888 0.04280
## SE.estimate 49
## 1c1
                49
                    2.4603 0.03457
                49 10.2701 0.16263
## ucl
## RB
                 0
                        NA
## RSE
                49
                    0.3777 0.00856
## COV
                 0
                        ΝA
                                 NA
##
## $`D = 10`
##
                      mean
## estimate
                50 10.0000 0.00000
                   2.6670 0.05948
## SE.estimate 50
               50 6.0032 0.06291
## 1c1
## ucl
               50 16.7560 0.19230
## RB
                0
                        NA
## RSE
                50
                    0.2667 0.00595
## COV
                 0
                        NΑ
                                 NΑ
```

Each estimate of RSE is essentially the same as before (see summary(stats1) in Summary method), but the run time is reduced by nearly 80%. Note the true value of density appears as a constant 'estimate' in the summary. Care is needed with this method as its performance in extreme cases has not been investigated fully.

Non-uniform populations

The simulated population by default has a uniform (homogeneous) Poisson distribution. To generate and sample from a spatially inhomogeneous population we use the 'IHP' option for argument model2D in secr::sim.popn. This involves three steps:

- 1. Create a habitat mask object with the desired extent.
- 2. Add to the mask a covariates dataframe with one or more columns defining pixel-specific densities.
- 3. In run.scenarios() specify a list of pop.args including model2D = "IHP" and D = "XX" where XX is the name of the particular mask covariate you wish to use for density, and name your mask in the 'maskset' argument.

A full demonstration is given in the Appendix (Non-uniform possums).

To visualize simulated populations you should set **savepopn = TRUE** in det.args and later extract the population the capthist object (for example, with a custom extractfn).

To compare several inhomogeneous distributions, specify several pop.args lists and use the popindex argument in make.scenarios(). The distribution may be varied simply by using the sim.popn() argument D to select different covariates of one mask.

The columns 'nrepeats' and 'D' in the scenarios argument of run.scenarios are ignored when model2D = "IHP". 'D' is replaced by the average density over the mask, which is used as the 'true' value of density in computing RB, RSE etc. in summaries. For stratified analyses you will have to define your own extractfn.

Linear habitat

secrdesign may be used to simulate sampling of populations in linear habitats as implemented in R package secrlinear. The procedure is similar to that for non-uniform (inhomogeneous Poisson) populations as

described in the previous section: one or more masks must be provided, but in this case they will be of type 'linearmask'.

The steps are:

- 1. Create linear habitat mask objects with the desired extent.
- 2. Create detector layouts and a scenario dataframe as usual.
- 3. Add a 'maskindex' column to the scenarios dataframe identifying which mask is to be used in each scenario (may be omitted for a single mask).
- 4. In run.scenarios() specify your mask(s) in the maskset argument.

Density may be specified in the scenario dataframe as a constant number of animals per km, and in this case the 'nrepeats' column is respected.

Density also may be modelled as inhomogeneous, i.e. varying along the length of the linear mask. The mechanism for this is like that for two dimensions: use a list of pop.args including model2D = "linear" and D = "XX" where XX is the name of the particular mask covariate you wish to use for density. In this case, the columns 'nrepeats' and 'D' in the scenarios dataframe are ignored, as for the 'IHP' option.

With a linear mask, run.scenarios defaults to secrlinear::networkdistance for the distance function (secr.fit argument details\$userdist).

Splitting data generation and model fitting

Each new detector layout or new model specification (in a fit.args list) defines a new scenario. The default procedure is to generate new data (both animal locations and simulated detection histories) for each scenario. To compare different models applied to the same dataset, save raw data from an initial call to run.scenarios() with fit = FALSE, extractfn = identity, and separately fit a list of models with fit.models. You can also peek at the raw data with the summary method.

Here, scen3 describes two scenarios, and in the call to fit.models each of these is split into two new scenarios, one for each component of fit.args.

It is not possible within **secrdesign** precisely to evaluate the application to the same animal distribution (population) of differing detector layouts or specifications for the fitted model (cf Fewster and Buckland 2004). Comparisons inevitably include variance from the varying number and placement of animals, and the sampling process; this variance may be reduced by fixing the number of individuals (pop.args = list(Ndist = "fixed")).

Populations with sub-classes or multiple sessions

Simulation of structured populations is introduced in **secrdesign** 2.2.0 and is still experimental.

The 'groups' argument of make.scenarios replicates rows so that within a scenario there is one row for each group. Group-specific parameter values are inserted by the user.

Rows sharing the same scenario number are recognised by run.scenarios as subclasses (groups). Each subclass is generated as a separate capthist object. The argument multisession determines whether the capthist objects corresponding to subclasses are pooled for analysis (using secr::rbind.capthist) or treated as multiple distinct sessions (using secr::MS.capthist).

The original sub-class of each individual is recorded as an individual covariate named "group". This is a factor. It may ignored in the fitted model, or used in such secr.fit arguments as 'groups' and 'hcov', or included in models directly as an individual covariate when CL = TRUE. (If the output from predict.secr is not a single dataframe then you will have to write a custom extractfn).

An example is given in the Appendix (Grouped populations).

Limitations, tips and troubleshooting

secrdesign has some limitations (Surprise!).

- 1. A progress message is output only on the completion of each scenario, which can be annoying, and when using multiple cores even this message is lost. It is strongly recommended that you start by generating summaries of raw data only (run.scenarios() with fit = FALSE), and confirm that your scenarios are realistic by reviewing the simulated number of detected individuals, total number of detections, etc. If these are inadequate or unrealistically large then there's no point going on. Then, try fitting with just a few replicates to be sure you have specified the model you intended and to assess the likely run time. Only then submit a run with a large number of replicates.
- 2. Only 2-parameter detection functions are allowed for data generation. This excludes the hazard-rate function, the cumulative gamma, and some others.
- 3. The default extractfn does does not handle models that produce more than one estimates table per replicate (e.g., finite mixture models). A custom extractfn is needed; it should either produce a numeric vector of 'selected statistics' or mimic single-dataframe output from predict().
- 4. The function secr::sim.capthist that generates detection histories for secrdesign has limited capacity for simulating temporal, behavioural or other heterogeneity in detection probability. Heterogeneity may be simulated as discrete subclasses (see preceding section). Only a simple permanent learned response is allowed in run.scenarios ('recapfactor').
- 5. As noted before, the same mask is used for generating populations and fitting models. It would be possible to replace the maskset component of a 'rawdata' object before running fit.models, but this is not recommended.
- 6. It is easy to forget the random number seed. Consider replacing the default value.
- 7. The method for fitting a fixed-N model (distribution = binomial) is somewhat fragile: it can fail when given a start value for D that is less than the minimum density observed (i.e. the number of distinct individuals divided by the mask area). This can easily happen when a population is simulated with pop.args = list(Ndist = "poisson") (the default) and sampled with high detection probability, but secr.fit is called with (distribution = "binomial"). The solution is to use pop.args = list(Ndist = "fixed").
- 8. If your summaries do not include enough significant digits, increase the 'dec' argument of summary.selectedstatistics!

References

Borchers, D. L. and Efford, M. G. (2008) Spatially explicit maximum likelihood methods for capture–recapture studies. *Biometrics* **64**, 377–385.

Cooch, E. and White, G. (eds) (2014) *Program MARK: A Gentle Introduction*. 13th edition. Available online at http://www.phidot.org/software/mark/docs/book/.

Efford, M. G. (2012) DENSITY 5.0: software for spatially explicit capture—recapture}. Department of Mathematics and Statistics, University of Otago, Dunedin, New Zealand http://www.otago.ac.nz/density.

Efford, M. G., Borchers D. L. and Byrom, A. E. (2009) Density estimation by spatially explicit capture–recapture: likelihood-based methods. In: D. L. Thomson, E. G. Cooch, M. J. Conroy (eds) *Modeling Demographic Processes in Marked Populations*. Springer. Pp 255–269.

Efford, M. G., Dawson, D. K. and Borchers, D. L. (2009) Population density estimated from locations of individuals on a passive detector array. *Ecology* **90**, 2676–2682.

Efford, M. G. and Fewster, R. M. (2013) Estimating population size by spatially explicit capture–recapture. *Oikos* 122, 918–928.

Fewster, R. M. and Buckland, S. T. (2004) Assessment of distance sampling estimators. In: S. T. Buckland, D. R. Anderson, K. P. Burnham, J. L. Laake, D. L. Borchers and L. Thomas (eds) *Advanced distance sampling*. Oxford University Press, Oxford, U. K. Pp. 281–306.

Williams, B. K., Nichols, J. D. and Conroy, M. J. (2002) Analysis and management of animal populations. Academic Press, San Diego

Appendix. Examples

Here we give some annotated examples of simulation code and selected output. Running this code with reduced nrepl, and viewing the output, will give you an idea of how secrdesign works.

Multiple grids, varying number of occasions

This is the example from the main text, slightly extended

```
traps4 <- list(grid6x6 = make.grid(6,6),</pre>
               grid8x9 = make.grid(8,9),
               grid12x12 = make.grid(12,12))
scen4 \leftarrow make.scenarios (trapsindex = 1:3, noccasions = c(8,4,2), D = 5,
    g0 = 0.2, sigma = c(20,30), crosstraps = FALSE)
sims4 <- run.scenarios(nrepl = 500, trapset = traps4, scenarios =</pre>
     scen4, fit = FALSE, ncores = 3)
class(sims4)
                    ## just peeking
## [1] "selectedstatistics" "secrdesign"
                                                   "list"
find.stats(sims4)
                    ## just peeking
## [1] "n"
              "ndet" "nmov" "dpa"
summary(sims4)
## run.scenarios(nrepl = 500, scenarios = scen4, trapset = traps4,
##
      fit = FALSE, ncores = 3)
##
## Replicates
                 500
## Started
                 21:06:10 23 Nov 2014
## Run time
                 0.369 minutes
## Output class selectedstatistics
##
## $constant
##
               value
## nrepeats
## D
                   5
## g0
                 0.2
## detectfn
                   0
## recapfactor
                   1
## popindex
                   1
## detindex
## fitindex
##
## $varying
## scenario trapsindex noccasions sigma maskindex
##
                           8
           1
                      1
                                       20
```

```
20
##
                                   20
##
          3
                     3
                                2
                                                3
##
          4
                     1
                                   30
                                                4
##
          5
                     2
                                4
                                   30
                                                5
                                2
##
          6
                     3
                                     30
                                                6
##
## $detectors
## trapsindex trapsname
##
            1
                grid6x6
##
            2 grid8x9
##
            3 grid12x12
##
## OUTPUT
## $`trapsindex = 1, noccasions = 8, sigma = 20, maskindex = 1`
         n mean
## n
       500 14.378 0.17151
## ndet 500 50.546 0.68825
## nmov 500 30.782 0.48020
## dpa 500 2.785 0.01952
## $`trapsindex = 2, noccasions = 4, sigma = 20, maskindex = 2`
         n mean
                       se
       500 21.342 0.20958
## n
## ndet 500 48.200 0.52306
## nmov 500 23.542 0.30967
## dpa 500 2.045 0.00915
##
## $`trapsindex = 3, noccasions = 2, sigma = 20, maskindex = 3`
         n mean
       500 32.404 0.25352
## n
## ndet 500 46.904 0.38394
## nmov 500 12.860 0.15695
## dpa 500 1.397 0.00382
##
## $`trapsindex = 1, noccasions = 8, sigma = 30, maskindex = 4`
         n mean
## n
       500 23.190 0.20912
## ndet 500 92.086 0.97648
## nmov 500 62.306 0.75562
## dpa 500 3.327 0.01948
## $`trapsindex = 2, noccasions = 4, sigma = 30, maskindex = 5`
         n mean
                       se
       500 31.648 0.23747
## n
## ndet 500 82.594 0.67822
## nmov 500 47.164 0.44615
## dpa 500 2.422 0.00874
##
## $`trapsindex = 3, noccasions = 2, sigma = 30, maskindex = 6`
         n mean
## n
       500 45.41 0.29922
## ndet 500 73.90 0.50501
## nmov 500 26.77 0.23381
## dpa 500 1.59 0.00347
```

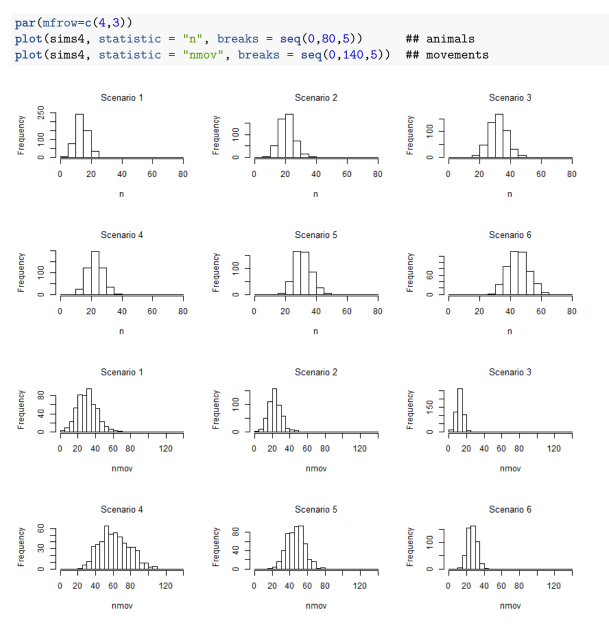


Fig. 4. Numbers of individuals (n) and movements (nmov) from six scenarios differing in trap number, number of sampling occasions and scale of movement.

Learned trap response

Here we assess the bias in \hat{D} caused by ignoring a learned trap response.

```
## select statistics and throw out any replicates with SE > 100
## (there is one -- see reduced n in output for scenario 11)
stats5 <- select.stats(sims5)
stats5 <- validate(stats5, "SE.estimate", c(0,100), "all")
sum5 <- summary(stats5, fields = c("n", "mean", "se", "lcl", "ucl", "median"))</pre>
```

```
## plot
plot(c(0.5,6.5), c(-0.2,0.4), type = "n", xlab = "Scenario", ylab = "RB(D-hat)")
for (i in 1:12) {
    xv <- if (i<=6) i else (i-6)+0.05
    segments (xv, sum5$0UTPUT[[i]]["RB","lcl"], xv, sum5$0UTPUT[[i]]["RB","ucl"])
    ptcol <- if (i<=6) "white" else "black"
    points(xv, sum5$0UTPUT[[i]]["RB","mean"], pch = 21, bg = ptcol)
}
abline(h = 0, col="red")
text(c(1.5,3.5,5.5), rep(0.38,3), paste("recapfactor", c(0.5,1,2), sep = " = "))</pre>
```

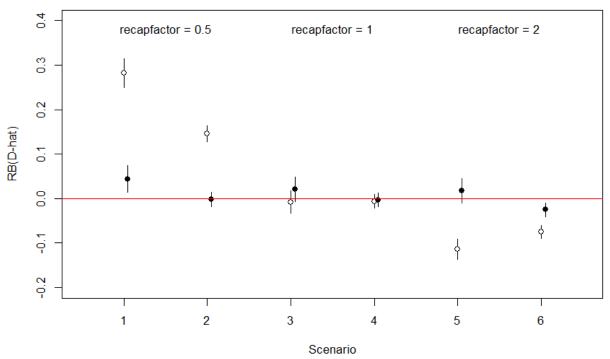


Fig. 5 Relative bias of SECR density estimate from null model (filled circles) and g0∼b model (open circles) when data were generated with negative, zero, or positive learned response.

```
## look at extended output
sum5
```

```
## run.scenarios(nrepl = 500, scenarios = scen5, trapset = traps5,
## fit = TRUE, fit.args = list(list(model = g0 ~ 1), list(model = g0 ~
## b)), ncores = 6)
##
## Replicates 500
## Started 18:54:34 23 Nov 2014
## Run time 103.4 minutes
## Output class selectedstatistics
```

```
##
## $constant
##
              value
## noccasions
## nrepeats
                  1
## D
                  5
## g0
                0.2
## sigma
                 25
## detectfn
                  0
## popindex
                  1
## detindex
                  1
##
## $varying
   scenario trapsindex recapfactor fitindex maskindex
##
           1
                                0.5
                                           1
                      1
           2
                      2
##
                                0.5
                                           1
                                                     2
##
           3
                      1
                                1.0
                                           1
                                                     1
                      2
##
           4
                                1.0
                                           1
##
           5
                      1
                                2.0
                                           1
                                                     1
                      2
##
           6
                                2.0
                                           1
                                                     2
##
          7
                      1
                                0.5
                                           2
                                                     1
##
           8
                      2
                                0.5
                                           2
          9
                                           2
##
                      1
                                1.0
                                                     1
##
          10
                      2
                                1.0
                                           2
                                                     2
                                           2
##
          11
                                2.0
                                                     1
                      1
##
          12
                      2
                                2.0
                                           2
                                                     2
##
## $detectors
##
   trapsindex trapsname
##
            1
                grid6x6
##
             2 grid10x10
##
## $fit.args
##
  fitindex model
##
           1 ~ g0 1
##
           2 \sim g0 b
##
## OUTPUT
## $`trapsindex = 1, recapfactor = 0.5, fitindex = 1, maskindex = 1`
##
                      mean
                                       lcl
                                               ucl median
                                se
                n
## estimate
               500 6.4126 0.08204 6.2518 6.5733 6.2543
## SE.estimate 500 1.9965 0.02054 1.9562 2.0368 1.9330
               500 3.5507 0.05517 3.4426 3.6589 3.3630
## lcl
## ucl
               500 11.6784 0.12692 11.4296 11.9271 11.3254
## RB
               500 0.2825 0.01641 0.2504 0.3147 0.2509
               500 0.3214 0.00251 0.3165 0.3263 0.3132
## RSE
## COV
               500 0.8600 0.01553 0.8296 0.8904 1.0000
##
## $`trapsindex = 2, recapfactor = 0.5, fitindex = 1, maskindex = 2`
                n mean
                               se
                                     lcl
                                          ucl median
## estimate
               500 5.7277 0.04459 5.6403 5.8151 5.6665
## SE.estimate 500 1.0221 0.00456 1.0132 1.0311 1.0175
               500 4.0501 0.03642 3.9787 4.1215 4.0027
## lcl
## ucl
               500 8.1067 0.05348 8.0019 8.2115 8.0313
```

```
500 0.1455 0.00892 0.1280 0.1630 0.1333
## RB
## RSE
              500 0.1809 0.00069 0.1796 0.1823 0.1794
## COV
              500 0.8800 0.01455 0.8515 0.9085 1.0000
##
## $`trapsindex = 1, recapfactor = 1, fitindex = 1, maskindex = 1`
##
                    mean
                               se lcl
                                              ucl
                                                   median
               n
              500 4.96161 0.06381 4.83654 5.08669 4.98843
## estimate
## SE.estimate 500 1.47525 0.01175 1.45223 1.49828 1.47518
              500 2.81512 0.04459 2.72772 2.90251
                                                   2.78540
              500 8.79952 0.08750 8.62803 8.97102 8.84495
## ucl
              500 -0.00768 0.01276 -0.03269 0.01734 -0.00231
              500 0.30992 0.00240 0.30521 0.31463 0.29930
## RSE
              500 0.95000 0.00976 0.93088 0.96912 1.00000
## COV
##
## $`trapsindex = 2, recapfactor = 1, fitindex = 1, maskindex = 2`
##
               n mean se
                                 lcl ucl median
              500 4.9675 0.03917 4.89071 5.04426 4.92346
## estimate
## SE.estimate 500 0.8810 0.00375 0.87364 0.88832 0.88014
             500 3.5202 0.03224 3.45698 3.58334 3.47659
## lcl
              500 7.0161 0.04641 6.92515 7.10708 6.97002
## ucl
## RB
              500 -0.0065 0.00783 -0.02186 0.00885 -0.01531
## RSE
              500 0.1801 0.00073 0.17866 0.18151 0.17812
## COV
              500 0.9600 0.00877 0.94281 0.97719 1.00000
##
## $`trapsindex = 1, recapfactor = 2, fitindex = 1, maskindex = 1`
               n
                  mean
                             se
                                  lcl
                                          ucl median
## estimate
              500 4.4284 0.05842 4.3139 4.54287 4.4158
## SE.estimate 500 1.2996 0.00981 1.2803 1.31878 1.3012
## lcl
              500 2.5323 0.04180 2.4504 2.61425 2.4967
## ucl
              500 7.8019 0.07721 7.6506 7.95325 7.8078
## R.B
              500 -0.1143 0.01168 -0.1372 -0.09143 -0.1168
## RSE
              500 0.3070 0.00243 0.3023 0.31178 0.2972
## COV
              500 0.9440 0.01029 0.9238 0.96417 1.0000
##
## $`trapsindex = 2, recapfactor = 2, fitindex = 1, maskindex = 2`
               n
                    mean
                             se lcl
                                              ucl median
## estimate
              500 4.62434 0.03824 4.54939 4.69929 4.59226
## SE.estimate 500 0.81600 0.00362 0.80891 0.82309 0.81839
             500 3.28267 0.03149 3.22094 3.34440 3.23926
## ucl
              500 6.52046 0.04528 6.43172 6.60920 6.49095
              500 -0.07513 0.00765 -0.09012 -0.06014 -0.08155
              500 0.17943 0.00076 0.17795 0.18092 0.17822
## RSE
              500 0.93200 0.01127 0.90991 0.95409 1.00000
## $`trapsindex = 1, recapfactor = 0.5, fitindex = 2, maskindex = 1`
##
               n mean se lcl ucl median
## estimate
              500 5.22167 0.07578 5.07315 5.37019 5.18323
## SE.estimate 500 1.74583 0.02123 1.70423 1.78744 1.71095
## lcl
              500 2.78488 0.05062 2.68566 2.88410 2.70792
              500 9.94459 0.12380 9.70195 10.18723 9.75850
## ucl
## R.B
              500 0.04433 0.01516 0.01463 0.07404 0.03665
             500 0.34976 0.00350 0.34290 0.35661 0.33759
## RSE
## COV
             500 0.94200 0.01046 0.92149 0.96251 1.00000
##
```

```
## $`trapsindex = 2, recapfactor = 0.5, fitindex = 2, maskindex = 2`
##
                       mean
                                        lcl
                                                      median
                 n
                                 se
                                                1107
## estimate
               500
                    4.98864 0.04090
                                     4.9085 5.06880
## SE.estimate 500
                   0.92341 0.00434 0.9149 0.93192
                                                      0.92743
## lcl
               500
                    3.48357 0.03322
                                     3.4184 3.54868
                                                      3.47396
## ucl
               500 7.15203 0.04916 7.0557 7.24838
                                                     7.15163
               500 -0.00227 0.00818 -0.0183 0.01376 -0.00455
## RB
## RSE
               500
                    0.18824 0.00084 0.1866 0.18989
                                                      0.18550
## COV
               500 0.94600 0.01012 0.9262 0.96583
##
## $`trapsindex = 1, recapfactor = 1, fitindex = 2, maskindex = 1`
##
                 n
                       mean
                                 se
                                         lcl
                                                  ucl median
## estimate
               500
                   5.10488 0.06876
                                     4.97011
                                              5.23966
                                                        4.9510
## SE.estimate 500
                   1.72420 0.08918
                                     1.54942
                                              1.89899
                                                        1.5818
               500 2.78061 0.04719
                                     2.68812
## 1c1
                                              2.87311
                                                        2.6784
## ucl
               500 10.08840 0.61405
                                     8.88488 11.29193
                                                       9.1970
## RB
               500 0.02098 0.01375 -0.00598
                                              0.04793 -0.0098
## RSE
               500 0.34471 0.01301
                                     0.31922
                                              0.37021
                                                       0.3212
                                              0.97560
## COV
               500 0.95800 0.00898 0.94040
                                                       1.0000
##
## $`trapsindex = 2, recapfactor = 1, fitindex = 2, maskindex = 2`
                 n
                      mean
                                se
                                       lcl
                                                ucl
                                                      median
                   4.9840 0.03952
## estimate
               500
                                    4.9065 5.06145
                                                    4.95613
                    0.9066 0.00427
## SE.estimate 500
                                    0.8983 0.91502
                                                    0.90778
## 1cl
               500 3.5012 0.03202 3.4385 3.56397
                                                    3.48533
## ucl
               500 7.1013 0.04781 7.0076 7.19497
                                                    7.08731
## RB
               500 -0.0032 0.00790 -0.0187 0.01229 -0.00877
               500 0.1846 0.00076 0.1831 0.18609
## RSE
                                                    0.18262
               500 0.9500 0.00976 0.9309 0.96912
## COV
                                                    1.00000
##
## $`trapsindex = 1, recapfactor = 2, fitindex = 2, maskindex = 1`
##
                                        lcl
                                                 ucl median
                      mean
                                se
                 n
## estimate
               499 5.08846 0.06975
                                    4.95175 5.22517 5.08003
                                    1.55528 1.62122 1.57657
## SE.estimate 499 1.58825 0.01682
               499 2.81611 0.04749
                                    2.72302 2.90920 2.80196
## lcl
               499 9.28585 0.10583 9.07842 9.49327 9.22943
## ucl
## RB
               499 0.01769 0.01395 -0.00965 0.04503 0.01601
## RSE
               499 0.32504 0.00269 0.31976 0.33031 0.31165
## COV
               499 0.93587 0.01098 0.91436 0.95739 1.00000
##
## $`trapsindex = 2, recapfactor = 2, fitindex = 2, maskindex = 2`
##
                 n
                       mean
                                 se
                                         lcl
                                                  ucl
                                                         median
## estimate
               500
                    4.87325 0.04075
                                     4.79337
                                              4.95313
                                                        4.82339
## SE.estimate 500 0.89038 0.00427
                                     0.88201
                                              0.89876
                                                       0.88879
## 1c1
               500
                    3.41845 0.03312
                                     3.35353
                                              3.48337
                                                        3.39162
## ucl
               500
                    6.95444 0.04902
                                     6.85836
                                              7.05052
                                                        6.91494
## RB
               500 -0.02535 0.00815 -0.04133 -0.00937 -0.03532
## RSE
               500 0.18580 0.00080
                                     0.18422
                                              0.18737
                                                        0.18290
## COV
               500 0.94800 0.00994
                                    0.92852
                                              0.96748
                                                       1.00000
```

Non-uniform possums

Code to illustrate the use of homogeneous and inhomogeneous density models.

```
## add covariates to builtin secr object possummask
## D1 is homogeneous density
## D2 is artificial SW - NE gradient in density
xy <- apply(possummask,1,sum) / 500</pre>
covariates(possummask)[, "D1"] <- 2</pre>
covariates(possummask)[, "D2"] <- xy - mean(xy) + 2.5</pre>
## Note that this object already had a covariates dataframe
## -- if it didn't we would use
## covariates(possummask) <- data.frame ( D1 = ..., D2 = ...)</pre>
## specify scenarios
## anticipate two different sets of arguments for sim.popn
## with popindex = 1:2
scen6 <- make.scenarios (g0 = 0.2, sigma = 45, noccasions = 5,
    popindex = 1:2)
## specify alternate models for distribution of animals
poplist <- list(list(model2D = "IHP", D = "D1"),</pre>
                list(model2D = "IHP", D = "D2"))
## run scenarios and summarise
## we use the trap layout from the builtin secr object possumCH
sims6 <- run.scenarios (500, scen6, traps(possumCH), possummask,</pre>
   pop.args = poplist)
```

summary(sims6)

```
## run.scenarios(nrepl = 500, scenarios = scen6, trapset = traps(possumCH),
      maskset = possummask, pop.args = poplist)
##
##
## Replicates
                500
## Started
                18:06:10 23 Nov 2014
## Run time
                0.812 minutes
## Output class selectedstatistics
##
## $constant
##
              value
## trapsindex
## noccasions
## nrepeats
                 1
## g0
                0.2
## sigma
                 45
## detectfn
                 1
## recapfactor
## detindex
## fitindex
                  1
## maskindex
##
```

```
## $varving
   scenario
             D popindex
           1 2.0
##
           2 2.5
                        2
##
##
## $detectors
   trapsindex trapsname
##
                  traps1
##
## $pop.args
   popindex model2D D
##
           1
                 IHP D1
           2
                 IHP D2
##
##
## OUTPUT
## ^D = 2, popindex = 1
##
               mean
          n
## n
        500 111.866 0.46103
## ndet 500 272.892 1.12915
## nmov 500 143.050 0.67248
## dpa 500
              2.186 0.00429
## ^D = 2.5, popindex = 2
##
          n
               mean
        500 139.704 0.48521
## n
## ndet 500 330.404 1.12189
## nmov 500 169.374 0.68597
## dpa 500
              2.128 0.00358
```

To visualize individual realisations of the distribution of animals, use fit = FALSE (the default), det.args = list(savepopn = TRUE), and save the entire capthist object (extractfn = identity). Here we create a single replicate.

```
sims6a <- run.scenarios (1, scen6, traps(possumCH), possummask,
    pop.args = poplist, det.args = list(savepopn = TRUE),
    extractfn = identity)</pre>
```

```
## sims6a$output is now a list (one component per scenario) of lists
## (one component per replicate) of simulated capthist objects, each
## with its 'popn' object embedded as an attribute

pop1 <- attr(sims6a$output[[1]][[1]], "popn")
pop2 <- attr(sims6a$output[[2]][[1]], "popn")
par(mfrow = c(1,2), mar=c(1,1,1,6))
plot(possummask, covariate = "D1", dots = FALSE, breaks = 0:6)
plot(traps(possumCH), detpar = list(col = 'green', pch = 15), add = TRUE)
plot(pop1, frame = FALSE, add = TRUE, col = "blue", pch = 16, cex = 0.6)
plot(traps(possumCH), detpar = list(col = 'green', pch = 15), add = TRUE)
plot(pop2, frame = FALSE, add = TRUE, col = "blue", pch = 16, cex = 0.6)</pre>
```

```
## click on map to display height; Esc to exit
spotHeight(possummask, prefix = "D2")
```

```
pop1 <- attr(sims6a$output[[1]][[1]], "popn")
pop2 <- attr(sims6a$output[[2]][[1]], "popn")
png(file='d:/density secr 2.9/secrdesign/vignettes/secrdesign-fig6.png',
    width=850, height=400)
par(mfrow = c(1,2), mar=c(1,1,1,6), cex=1.25)
plot(possummask, covariate = "D1", dots = FALSE, breaks = 0:6)
plot(traps(possumCH), detpar = list(col = 'green', pch = 15), add = TRUE)
plot(pop1, frame = FALSE, add = TRUE, col = "blue", pch = 16, cex = 0.6)
plot(possummask, covariate = 'D2', dots = FALSE, breaks = 0:6)
plot(traps(possumCH), detpar = list(col = 'green', pch = 15), add = TRUE)
plot(pop2, frame = FALSE, add = TRUE, col = "blue", pch = 16, cex = 0.6)
dev.off()</pre>
```

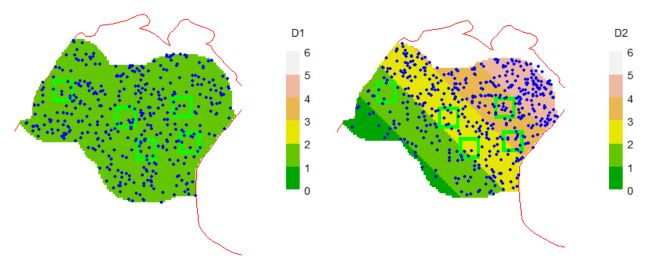


Fig. 6. Simulated homogeneous (left) and inhomogeneous (right) distributions of brushtail possums at Waitarere, New Zealand. Traps in green (each hollow grid 180 m square).

Code for linear habitat

Code to illustrate the use of linear habitat models. This assumes you have the package **secrlinear**, which is not yet on CRAN.

summary(sims7)

```
## run.scenarios(nrepl = 500, scenarios = scen7, trapset = trplist,
##
      maskset = linmask, det.args = list(det.arg), fit = FALSE,
##
      seed = 345)
## Replicates
                500
## Started
                18:02:42 23 Nov 2014
## Run time
                2.244 minutes
## Output class selectedstatistics
##
## $constant
##
              value
## noccasions
## nrepeats
                  1
## g0
                0.2
## sigma
                 25
## detectfn
## recapfactor
## popindex
## detindex
## fitindex
## maskindex
##
## $varying
## scenario trapsindex
##
         1 1 50
##
          2
                    2 50
                    1 200
##
          3
##
          4
                     2 200
##
## $detectors
## trapsindex trapsname
##
            1 spacing30
##
            2 spacing60
##
## $det.args
```

```
detindex
               userdist
##
           1 userdistfn
##
## OUTPUT
## $`trapsindex = 1, D = 50`
##
          n
              mean
                        se
        500 90.90 0.40609
## n
## ndet 500 152.00 0.78590
  nmov 500
             40.92 0.33327
  dpa 500
              1.45 0.00294
##
## $`trapsindex = 2, D = 50`
##
          n
              mean
        500 59.172 0.34544
## n
## ndet 500 75.112 0.46533
## nmov 500 5.996 0.10661
## dpa 500 1.102 0.00174
##
## $`trapsindex = 1, D = 200`
##
          n
               mean
## n
        500 357.760 0.84627
## ndet 500 600.050 1.58310
## nmov 500 161.766 0.62869
## dpa 500
              1.452 0.00144
##
## $`trapsindex = 2, D = 200`
##
          n
               mean
## n
        500 236.124 0.6774
## ndet 500 300.358 0.9301
## nmov 500
             24.186 0.2252
## dpa 500
              1.102 0.0009
```

Grouped populations

This example demonstrates the simulation of a structured population - nominally females and males with a 2:1 sex ratio.

First we form a scenarios dataframe with 2 groups and two levels of 'noccasions', and manually adjust the 'male' parameter values:

```
scen8 <- make.scenarios (D = 8, g0 = 0.3, sigma = 30, noccasions = c(4,8), groups = c('F','M'))
male <- scen8$group == 'M'
scen8$D[male] <- 4
scen8$g0[male] <- 0.2
scen8$sigma[male] <- 40
scen8[,1:8]</pre>
```

```
##
     scenario group trapsindex noccasions nrepeats D
## 1
             1
                   F
                                           4
                                                      1 8 0.3
                               1
                                            4
## 2
             1
                   М
                               1
                                                      1 4 0.2
                                                                  40
## 3
             2
                   F
                               1
                                           8
                                                      1 8 0.3
                                                                  30
             2
## 4
                   М
                                1
                                           8
                                                      1 4 0.2
                                                                  40
```

Next we set up a trapping grid, a habitat mask, and a customized extract function for multi-class output from a hybrid mixture model:

```
grid <- make.grid(8, 8, spacing = 30)
mask <- make.mask(grid, buffer = 160, type = 'trapbuffer')
## extracts total density and proportion from output for the first group (F)
exfn <- function(x) {
    if (inherits(x, 'secr') & !is.null(x$fit)) {
        pred <- predict(x)
        pred[[1]][c('D', 'pmix'),]
    }
    else data.frame()
}</pre>
```

It is desirable to check the raw simulations. We specify the mask, rather than relying on one constructed automatically, to ensure the same mask is used for both females and males.

```
## raw8 <- run.scenarios(20, scen8, trapset = list(grid), fit = FALSE, maskset = list(mask))
summary(raw8)
## run.scenarios(nrepl = 20, scenarios = scen8, trapset = list(grid),
##
       maskset = list(mask), fit = FALSE)
##
                 20
## Replicates
                 20:59:21 25 Nov 2014
## Started
## Run time
                 0.026 minutes
## Output class selectedstatistics
##
## $constant
##
               value
## trapsindex
                   1
## nrepeats
                   1
## detectfn
                   0
## recapfactor
## popindex
                   1
## detindex
## fitindex
## maskindex
##
## $varying
    scenario group noccasions D gO sigma
                            4 8 0.3
##
           1
                 F
                             4 4 0.2
                                        40
##
           1
                 М
##
           2
                 F
                             8 8 0.3
                                        30
##
                             8 4 0.2
                                        40
##
## $detectors
##
    trapsindex trapsname
##
             1
                  traps1
##
## OUTPUT
## ^{r} group = F, noccasions = 4, D = 8, g0 = 0.3, sigma = 30 +
```

group = M, noccasions = 4, D = 4, g0 = 0.2, sigma = 40

```
##
          n
               mean
## F.n
            77.700 1.94679
         20
## F.ndet 20 203.500 6.10371
## F.nmov 20 107.950 4.21930
## F.dpa 20
              2.268 0.02970
## M.n
         20 44.350 1.04195
## M.ndet 20 110.900 3.44804
## M.nmov 20 60.900 2.73756
## M.dpa 20
              2.308 0.04501
##
## $`group = F, noccasions = 8, D = 8, g0 = 0.3, sigma = 30 +
     group = M, noccasions = 8, D = 4, g0 = 0.2, sigma = 40
##
               mean
          n
## F.n
          20 84.600 2.17207
## F.ndet 20 376.400 10.42805
## F.nmov 20 247.150 7.21339
## F.dpa 20
              3.340 0.04695
## M.n
         20 53.650
                     1.56487
## M.ndet 20 236.100 6.22850
## M.nmov 20 165.850 4.89160
## M.dpa 20
              3.672 0.07370
```

Now fit the models and check the summary output for density ('D') and sex ratio (proportion female 'pmix') without repeating the header information.

summary(select.stats(sims8,'D'))\$OUTPUT

```
## $`group = F, noccasions = 4, D = 8, g0 = 0.3, sigma = 30, maskindex = 1 +
     group = M, noccasions = 4, D = 4, g0 = 0.2, sigma = 40, maskindex = 2
##
               n
                      mean
## estimate
               20 12.21220 0.17656
## SE.estimate 20 1.18185 0.01178
## lcl
              20 10.10681 0.15528
## ucl
               20 14.75645 0.20020
               20 0.01768 0.01471
## RB
## RSE
              20 0.09690 0.00051
## COV
              20 1.00000 0.00000
## $`group = F, noccasions = 8, D = 8, g0 = 0.3, sigma = 30, maskindex = 1 +
     group = M, noccasions = 8, D = 4, g0 = 0.2, sigma = 40, maskindex = 2
##
               n
                      mean
               20 11.74179 0.22664
## estimate
## SE.estimate 20 1.04618 0.01089
              20 9.86404 0.20609
## lcl
              20 13.97771 0.24764
## ucl
              20 -0.02152 0.01889
## RB
## RSE
              20 0.08940 0.00085
              20 1.00000 0.00000
## COV
```

summary(select.stats(sims8,'pmix'))\$OUTPUT

```
## Warning: assuming first scenario row corresponds to requested pmix
## Warning: assuming first scenario row corresponds to requested pmix
## $`group = F, noccasions = 4, D = 8, gO = 0.3, sigma = 30, maskindex = 1 +
## group = M, noccasions = 4, D = 4, g0 = 0.2, sigma = 40, maskindex = 2
##
               n
                     mean
## estimate
              20 0.66197 0.00987
## SE.estimate 20 0.04485 0.00041
              20 0.56929 0.01015
## lcl
              20 0.74358 0.00893
## ucl
## RB
              20 -0.00705 0.01480
              20 0.06818 0.00155
## RSE
              20 0.95000 0.05000
## COV
## $`group = F, noccasions = 8, D = 8, g0 = 0.3, sigma = 30, maskindex = 1 +
## group = M, noccasions = 8, D = 4, g0 = 0.2, sigma = 40, maskindex = 2
##
               n
                    mean
## estimate
              20 0.67468 0.00898
## SE.estimate 20 0.04044 0.00053
             20 0.59102 0.00949
## lcl
## ucl
              20 0.74846 0.00808
## RB
              20 0.01202 0.01347
## RSE
              20 0.06028 0.00142
## COV
              20 0.95000 0.05000
```