

# Package ‘LCMCR’

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**Type** Package

**Title** Bayesian Non-Parametric Latent-Class Capture-Recapture

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**Author** Daniel Manrique-Vallier

**Description** Bayesian population size estimation using non parametric latent-class models.

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**License** GPL (>= 2)

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LCMCR-package

*Capture-Recapture Estimation using Bayesian Nonparametric latent-class models*

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## Description

This package implements a fully Bayesian multiple-recapture method for estimating the unknown size of a population using non-parametric latent class models. This is an implementation of the method described in Manrique-Vallier (2016). The estimation algorithm is based on Markov Chain Monte Carlo sampling.

## Details

Package: LCMCR  
Type: Package  
Version: 0.4.13  
Date: 2023-05-23  
License: GPL >= 2

## Author(s)

Daniel Manrique-Vallier <dmanriqu@indiana.edu>

## References

Manrique-Vallier, D. (2016) "Bayesian Population Size Estimation Using Dirichlet Process Mixtures", *Biometrics*.

## Examples

```
library('LCMCR')

###Using Kosovo data.###
data(kosovo_aggregate)

###Example 1: Capture-Recapture estimation using convenience functions###
#Create and initialize an LCMCR object for MCMC sampling#
## Not run:
sampler <- lcmCR(captures = kosovo_aggregate, tabular = FALSE, in_list_label = '1',
  not_in_list_label = '0', K = 10, a_alpha = 0.25, b_alpha = 0.25,
  seed = 'auto', buffer_size = 10000, thinning = 100)
#Obtain 1000 samples from the posterior distribution of N#
N <- lcmCR_PostSampl(sampler, burnin = 10000, samples = 1000, thinning = 100, output = FALSE)

#Posterior quantiles#
```

```

quantile(N, c(0.025, 0.5, 0.975))

###Example 2: Capture-Recapture estimation using the lcm_CR_Basic object directly###
#Create and initialize an LCMCR object for MCMC sampling#
sampler <- lcmCR(captures = kosovo_aggregate, tabular = FALSE, in_list_label = '1',
               not_in_list_label = '0', K = 10, a_alpha = 0.25, b_alpha = 0.25,
               seed = 'auto', buffer_size = 1000, thinning = 100)

#Run 10000 iterations as burn-in
sampler$update(10000, output = FALSE)

#List all parameters from the model
sampler$get_param_list()

#Set parameter 'n0' for tracing
sampler$set_trace('n0')

#List currently traced parameters.
sampler$get_trace_list()

#Activate tracing
sampler$activate_tracing()

#Run the sampler 100000 times
sampler$update(100000, output = FALSE)

#Get the 1000 samples from the posterior distribution of N
N <- sampler$get_trace('n0') + sampler$n

#Plot the trace of N
plot(N, type = 'l')

#Compute posterior quantiles
quantile(N, c(0.025, 0.5, 0.975))

## End(Not run)

```

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kosovo\_aggregate

*Killings in the Kosovo war from March 20 to June 22, 1999.*

---

### Description

Capture pattern data for  $J = 4$  independently collected lists that jointly document  $n = 4400$  observed killings in the Kosovo war between March 20 to June 22, 1999.

### Usage

```
data("kosovo_aggregate")
```

**Format**

A data frame with 4400 observations on the following 4 variables.

EXH a factor with levels 0 1

ABA a factor with levels 0 1

OSCE a factor with levels 0 1

HRW a factor with levels 0 1

**Details**

This data set was analyzed by Ball et al. (2002).

**References**

Ball, P., Betts, W., Scheuren, F., Dudukovic, J., and Asher, J. (2002), "Killings and Refugee Flow in Kosovo, March/June, 1999," Report to ICTY.

**Examples**

```
data(kosovo_aggregate)
```

---

lcmCR

*Bayesian Nonparametric Latent Class Capture-Recapture*


---

**Description**

Create and initialize an object of class lcm\_CR\_Basic.

**Usage**

```
lcmCR(captures, tabular = FALSE, in_list_label = "1", not_in_list_label = "0",
      K = 5, a_alpha = 0.25, b_alpha = 0.25, buffer_size = 10000, thinning = 10,
      seed = "auto", verbose = TRUE)
```

**Arguments**

<code>captures</code>	input dataset. A data frame with the multiple-recapture data. See 'Details' for input formats.
<code>tabular</code>	a logical value indicating whether or not the data is tabulated. See 'Details'.
<code>in_list_label</code>	factor label that indicates that individual is in list (e.g. 'Yes')
<code>not_in_list_label</code>	factor label that indicates that individual is in not list (e.g. 'No')
<code>K</code>	maximum number of latent classes. Indicates the truncation level of the stick-breaking process.
<code>a_alpha</code>	shape parameter of the prior distribution of concentration parameter of the stick-breaking process.

b_alpha	inverse scale parameter of the prior distribution of concentration parameter of the stick-breaking process.
buffer_size	size of the tracing buffer.
thinning	thinning interval for the tracing buffer
seed	integer seed of the internal RNG.
verbose	Generate progress messages?

### Details

Input data must be provided as a data frame. The first J columns are two-level factors representing the multiple-recapture lists. Arguments `in_list_label` and `not_in_list_label` indicate the labels that represent inclusion and exclusion from the lists. This function supports two input formats:

- When `tabular=FALSE` each row represents a single individual's capture history. The number of rows must match the size of the observed population. Rows indicating no capture in all list simultaneously are illegal.
- When `tabular=TRUE` each row represents a unique capture pattern. This format requires an additional numeric column at the right, called "Freq", indicating the count corresponding to such pattern.

### Value

An object of class `lcm_CR_Basic` initialized and ready to use.

### Author(s)

Daniel Manrique-Vallier

### See Also

[lcm\\_CR\\_Basic](#), [lcm\\_CR\\_Basic\\_generator](#)

### Examples

```
require('LCMCR')
data(kosovo_aggregate)
sampler <- lcmCR(captures = kosovo_aggregate, tabular = FALSE, in_list_label = '1',
  not_in_list_label = '0', K = 10, a_alpha = 0.25, b_alpha = 0.25,
  seed = 'auto', buffer_size = 10000, thinning = 100)
sampler
N <- lcmCR_PostSampl(sampler, burnin = 10000, samples = 1000, thinning = 100, output = FALSE)
quantile(N, c(0.025, 0.5, 0.975))
```

---

lcmCR_PostSampl	<i>Generate Samples from the Posterior Distribution of Population Size under a LCMCR Model</i>
-----------------	--

---

### Description

Convenience function for generate samples from the posterior distribution of the population size using an initialized `lcm_CR_Basic` object.

### Usage

```
lcmCR_PostSampl(object, burnin = 10000, samples = 1000, thinning = 10,
  clear_buffer = FALSE, output = TRUE)
```

### Arguments

object	an initialized <code>lcm_CR_Basic</code> object.
burnin	number of burn in iterations.
samples	Number of samples to be generated. Note that this is not the same as the number of iterations for the sampler. Samples are saved one every thinning iterations.
thinning	subsampling interval. Samples are saved one every thinning iterations.
clear_buffer	logical. Clear the tracing buffer before sampling?
output	logical. Print messages?

### Value

A vector with the samples posterior samples of the population size parameter.

### Warning

Invoking this function deletes the content of the object's tracing buffer.

### Note

To create and initialize the `lcm_CR_Basic` object use `lcmCR` or `lcm_CR_Basic_generator`. The user is responsible to check whether the chain has reached the stationary distribution or not.

### Author(s)

Daniel Manrique-Vallier

### Examples

```
data(kosovo_aggregate)
sampler <- lcmCR(captures = kosovo_aggregate, tabular = FALSE, in_list_label = '1',
  not_in_list_label = '0', K = 10, a_alpha = 0.25, b_alpha = 0.25, seed = 'auto')
N <- lcmCR_PostSampl(sampler, burnin = 10000, samples = 1000, thinning = 100, output = FALSE)
quantile(N, c(0.025, 0.5, 0.975))
```

---

lcm\_CR\_Basic-class      *Class "lcm\_CR\_Basic"*


---

### Description

MCMC sampler for the Bayesian non-parametric latent class capture-recapture model.

### Extends

Class "[MCMCenviro](#)n", directly. All reference classes extend and inherit methods from "[envRefClass](#)".

### Fields

All fields are read-only.

**pointer:** external pointer to the C++ object.

**blobsize:** size (in bytes) of the raw object data for serialization. (currently not implemented.)

**local\_seed:** seed of the internal random number generator.

**J:** number of lists in the Capture-Recapture data.

**K:** maximum number of latent classes in the model (truncation level of the stick-breaking process).

**n:** observed number of individuals.

**Captures:** original provided data.

### Methods

`initialize(data_captures, K, a_alpha, b_alpha, in_list_symbol, len_buffer, subsamp):`  
Class constructor.

#### Arguments:

**data\_captures:** input dataset. A data frame with the multiple-recapture data.

**K:** maximum number of latent classes. Indicates the truncation level of the stick-breaking process.

**a\_alpha:** shape parameter of the prior distribution of concentration parameter of the stick-breaking process.

**b\_alpha:** inverse scale parameter of the prior distribution of concentration parameter of the stick-breaking process.

**in\_list\_symbol:** factor label that indicates that individual is in list (e.g. 'Yes')

**buffer\_size:** Size of the tracing buffer.

**subsamp:** thinning interval for the tracing buffer.

**verbose:** logical. Generate progress messages?

The following methods are inherited (from the corresponding class): `Change_SubSamp` ("MCMCenviro"), `Set_Trace` ("MCMCenviro"), `Change_Trace_Length` ("MCMCenviro"), `initialize` ("MCMCenviro"), `Get_Iteration` ("MCMCenviro"), `Get_Param` ("MCMCenviro"), `Reset_Traces` ("MCMCenviro"), `Get_Status` ("MCMCenviro"), `Update` ("MCMCenviro"), `Get_Trace_Size` ("MCMCenviro"), `Get_Trace` ("MCMCenviro"), `Get_Trace_List` ("MCMCenviro"), `Get_Param_List` ("MCMCenviro"), `Init_Model` ("MCMCenviro"), `Activate_Tracing` ("MCMCenviro"), `Deactivate_Tracing` ("MCMCenviro"), `Set_Seed` ("MCMCenviro"), `show` ("MCMCenviro")

**Note**

Use the convenience function `lcmCR` to create objects of this class. This class inherits most of its functionality from "`MCMCenvirom`".

**Author(s)**

Daniel Manrique-Vallier

**See Also**

`lcmCR`, `MCMCenvirom`.

**Examples**

```
showClass("lcm_CR_Basic")
```

---

`lcm_CR_Basic_generator`

*Generator for Class lcm\_CR\_Basic*

---

**Description**

Generator function for class `lcm_CR_Basic`.

**Usage**

```
lcm_CR_Basic_generator(...)
```

**Arguments**

... arguments to be passed to `lcm_CR_Basic` constructor.

**Value**

An object of class `lcm_CR_Basic`.

**Note**

The convenience function `lcmCR` provides a simpler mechanism to create `lcm_CR_Basic` objects.

**Author(s)**

Daniel Manrique-Vallier.

**See Also**

`lcmCR`



**Examples**

```
data(kosovo_aggregate)
x <- lcm_CR_Basic_generator(data_captures=kosovo_aggregate, K=10, a_alpha=0.25, b_alpha=0.25,
                           len_buffer=10000, subsamp=500, in_list_symbol = '1')
x$Get_Status()
```

---

MCMCenviro-class      *Class "MCMCenviro"*


---

**Description**

A generic interface for MCMC sampler objects implementing Bayesian models. Methods provide access to underlying functionality implemented in C++. The underlying implementation provides basic functionality for controlling the chain, and a 'tracing buffer' for storing and retrieving the samples.

**Extends**

All reference classes extend and inherit methods from "[envRefClass](#)".

**Fields**

(All fields are read-only.)

**pointer:** external pointer to the C++ object

**blobsize:** size (in bytes) of the raw object data for serialization. (currently not implemented.)

**seed:** seed of the internal random number generator.

**Methods****GENERAL METHODS**

**Init\_Model(output = TRUE, seed=c('auto', 'r.seed')):** Initializes the sampler.

**Arguments:**

**output:** logical. Print messages to the screen?

**seed:** integer. Seed of the internal RNG. Additionally, `seed='auto'` autogenerates the seed from the internal clock; `seed='r.seed'` autogenerates the seed from the current state of the `.Random.seed` variable.

**Update(num\_iter, output = TRUE):** Runs `num_iter` iterations of the sampler. Set `output = FALSE` to suppress console output.

**Get\_Iteration():** Retrieves the current number of iterations the sampler.

**Get\_Param\_List():** Retrieves the names of the parameters of the model.

**Get\_Param(param):** Retrieves the current value of the parameter `param`.

**Set\_Seed(seed):** Seeds the internal random number generator. It does not affect R's internal RNG.

Get\_Status(): Retrieves the current state of the chain

**Value:**

iteration numeric. Current iteration

initialized logical. Is the sampler initialized?

buffer\_size numeric. Capacity (in samples) of the tracing buffer.

buffer\_used numeric. Number of samples currently stored in the tracing buffer.

tracing character. Names of the variables currently traced.

thinning numeric. Thinning interval of the tracing buffer.

**METHODS FOR CONTROLLING THE TRACING BUFFER**

Get\_Trace\_List(): Retrieves the names of the parameters being currently traced.

Activate\_Tracing(): Activates the tracing buffer. Traced variables will be stored in the buffer when generated with Update().

Deactivate\_Tracing(): Deactivates the tracing buffer. Calls to Update() will not store samples in the buffer.

Set\_Trace(traces): Adds parameters to tracer.

**Arguments:**

param: character vector. Names of the parameters to trace. To list the available parameters for tracing use the Get\_Param\_List() method.

Get\_Trace(param): Retrieves samples stored in the tracing buffer.

**Arguments:**

param: character. Name of the parameter to retrieve.

**Value:** An array. The first dimension indexes the sample; the rest correspond to the original dimensions of the parameter as defined in the model.

Reset\_Traces(): Deletes the content of the tracing buffer.

Change\_SubSamp(new\_subsamp): Changes the sub-sampling period (thinning) of the tracing buffer.

**Warning:** This operation deletes the current content of the tracing buffer.

Get\_Trace\_Size(): Retrieves the size (in number of samples) of the trace buffer.

Change\_Trace\_Length(new\_length): Changes the size (in number of samples) of the tracing buffer.

**Warning:** This operation deletes the current content of the tracing buffer.

**Note**

This class is not designed to be used directly, but as a generic interface for samplers implementing specific models.

**Author(s)**

Daniel Manrique-Vallier

**Examples**

```
showClass("MCMCenvirom")
```

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