# Package 'banocc'

December 13, 2024

Type Package

Title Bayesian ANalysis Of Compositional Covariance

Version 1.31.0

Date 2022-04-20

**Maintainer** George Weingart <george.weingart@gmail.com>, Curtis Huttenhower <chuttenh@hsph.harvard.edu>

Description BAnOCC is a package designed for compositional data, where each sample sums to one. It infers the approximate covariance of the unconstrained data using a Bayesian model coded with 'rstan'. It provides as output the 'stanfit' object as well as posterior median and credible interval estimates for each correlation element.

License MIT + file LICENSE

**Depends** R (>= 3.5.1), rstan (>= 2.17.4)

**Imports** coda (>= 0.18.1), mvtnorm, stringr

Suggests knitr, rmarkdown, methods, testthat, BiocStyle

VignetteBuilder knitr

RoxygenNote 5.0.1

biocViews ImmunoOncology, Metagenomics, Software, Bayesian

git\_url https://git.bioconductor.org/packages/banocc

git\_branch devel

git\_last\_commit eef3ada

git\_last\_commit\_date 2024-10-29

**Repository** Bioconductor 3.21

**Date/Publication** 2024-12-13

**Author** Emma Schwager [aut, cre], Curtis Huttenhower [aut] 2 banocc\_model

# **Contents**

bano	cc	bano tion	cc:	A	pα	ıck	age	e fo	or.	Βa	ıye	sic	an	A	Va	lys	sis	oj	f C	on	np	os	iti	on	al	C	'or	re	la	-
Index																														11
	run_banocc							•	•	•		•	•		•							•	•			•	•			8
	get_banocc_output																													
	counts_pos_spike																													7
	counts_null																													6
	counts_neg_spike																													
	counts_hard_null	-																												
	compositions_pos_																													
	compositions_neg_ compositions_null	_																												
	compositions_hard																													
	banocc_model																													
	banocc																													

## **Description**

BAnOCC is a package for inferring correlations between features in compositional data, where each sample sums to one. It provides one object, banocc\_model and one function, run\_banocc

# banocc objects

 $banocc\_model\ has\ the\ stan\ model\ code\ to\ be\ compiled\ using\ rstan::stan.$ 

### banocc functions

run\_banocc takes a compiled model, and returns the 'stanfit' object resulting from a call to rstan::sampling get\_banocc\_output takes a 'stanfit' object or the output of run\_banocc and returns a list with the posterior median and credible interval estimates

banocc_model	The stan model used in the Bayesian fit	
--------------	---	--

# Description

This is the literal model used for fitting in Stan

# Usage

banocc\_model

compositions\_hard\_null

## **Format**

An object of class character of length 1.

## Value

The BAnOCC model as a string to be compiled with rstan::stan\_model

# **Examples**

```
data(compositions_null)
## Not run:
   compiled_banocc_model <- rstan::stan_model(model_code = banocc_model)
## End(Not run)</pre>
```

compositions\_hard\_null

Simulated compositional data with no feature correlations

# Description

These are the normalized samples corresponding to counts\_hard\_null. They should have a very different correlation structure from the counts. In particular, there should be one strong, positive association which is not present in the count correlation structure.

#### Usage

```
compositions_hard_null
```

## **Format**

A data frame with 1000 rows (compositional samples) and 9 variables (the features)

## Value

A data frame with 1000 compositional samples from 9 features, generated by dividing each row of counts\_hard\_null by its sum.

4 compositions\_null

compositions\_neg\_spike

Simulated compositional data with a negative count correlation

#### **Description**

These are the normalized data corresponding to counts\_neg\_spike. The count data have one negative feature correlation, but the compositional correlation structure should be different.

## Usage

compositions\_neg\_spike

#### **Format**

A data frame with 1000 rows (compositional samples) and 9 variables (the features)

#### Value

A data frame with 1000 compositional samples from 9 features, generated by dividing each row of counts\_neg\_spike by its sum.

compositions\_null

Simulated compositional data with no feature correlations

## **Description**

These are the normalized samples corresponding to counts\_null. They should have a similar (but not identical) correlation structure.

#### Usage

compositions\_null

## **Format**

A data frame with 1000 rows (compositional samples) and 9 variables (the features)

#### Value

A data frame with 1000 compositional samples from 9 features, generated by dividing each row of counts\_null by its sum.

compositions\_pos\_spike

Simulated compositional data with a positive count correlation

#### **Description**

These are the normalized data corresponding to counts\_pos\_spike. The count data have one positive feature correlation, but the compositional correlation structure should be different.

#### Usage

compositions\_pos\_spike

#### **Format**

A data frame with 1000 rows (compositional samples) and 9 variables (the features)

#### Value

A data frame with 1000 compositional samples from 9 features, generated by dividing each row of counts\_pos\_spike by its sum.

counts\_hard\_null

Simulated count data with no feature correlations

### **Description**

Nine features are draw independently from very different log-normal distributions whose means and variances are positively correlated. This means that the compositions generated from this dataset (see compositions\_hard\_null) should be have a correlation structure very different from that of these counts.

#### Usage

counts\_hard\_null

#### **Format**

A data frame with 1000 rows (samples) and 9 variables (the features)

## Value

A data frame with 1000 unconstrained samples from 9 features.

6 counts\_null

counts\_neg\_spike

Simulated count data with one negative feature correlation

## **Description**

Nine features are drawn from a log-normal distribution with one negative correlation. The resulting compositions are in compositions\_neg\_spike

#### Usage

```
counts_neg_spike
```

#### **Format**

A data frame with 1000 rows (samples) and 9 variables (the features)

#### Value

A data frame with 1000 unconstrained samples from 9 features.

counts\_null

Simulated count data with no feature correlations

## **Description**

Nine features are drawn independently from similar log-normal distributions to generate null count data. Because the feature distributions are very similar, the compositions generated from this dataset (see compositions\_null), should have a correlation structure similar to that of the counts.

## Usage

counts\_null

#### Format

A data frame with 1000 rows (the samples) and 9 variables (the features)

#### Value

A data frame with 1000 unconstrained samples from 9 features.

counts\_pos\_spike 7

counts_pos_spike Simulated count data with one positive feature correlation	
---	--

# Description

Nine features are drawn from a log-normal distribution with one positive correlation. The resulting compositions are in compositions\_pos\_spike.

## Usage

```
counts_pos_spike
```

## **Format**

A data frame with 1000 rows (samples) and 9 variables (the features)

#### Value

A data frame with 1000 unconstrained samples from 9 features.

get_banocc_output	Takes a model fit from BAnOCC, evaluates convergence and generates appropriate convergence metrics and inference

# Description

Takes a model fit from BAnOCC, evaluates convergence and generates appropriate convergence metrics and inference

## Usage

```
get_banocc_output(banoccfit, conf_alpha = 0.05, get_min_width = FALSE,
  calc_snc = TRUE, eval_convergence = TRUE, verbose = FALSE,
  num_level = 0)
```

## **Arguments**

banoccfit	Either a stanfit object (the Fit element returned by run_banocc), or the list returned by a call to run_banocc.
conf_alpha	The percentage of the posterior density outside the credible interval. That is, a $1-conf_alpha*100\%$ credible interval will be returned.
get_min_wi	A boolean value: should the minimum CI width that includes zero be calculated?
calc_snc	Boolean: should the scaled neighborhood criterion be calculated?

run\_banocc

eval\_convergence

Boolean: if 'TRUE', convergence will be evaluated using the Rhat statistic, and the fit output (estimates, credible intervals, etc.) will be missing if this statistic

does not indicate convergence.

verbose Print informative statements as the function executes?

num\_level The number of indentations to add to the output when verbose = TRUE.

#### Value

Returns a named list with the following elements:

CI The 1-conf\_alpha \* 100% credible intervals

Estimates.median The correlation estimates, which are the marginal posterior medians

*Min.width* Only present if the get\_min\_width argument is TRUE. The minimum CI width that includes zero for each correlation.

SNC Only present if the calc\_snc argument is TRUE. The scaled neighborhood criterion for each correlation.

*Fit* The stanfit object returned by the call to run\_banocc.

**Data** Only present if the banoccfit argument is specified as the output of a call to run\_banocc. It will be missing if banoccfit is specified as a stanfit object.

#### See Also

vignette("banocc-vignette") for more examples.

## **Examples**

run\_banocc

Runs BAnOCC to fit the model and generate appropriate convergence metrics and inference.

#### Description

Runs BAnOCC to fit the model and generate appropriate convergence metrics and inference.

run\_banocc 9

# Usage

```
run_banocc(compiled_banocc_model, C, n = rep(0, ncol(C)), L = 10 *
    diag(ncol(C)), a = 0.5, b = 0.01, cores = getOption("mc.cores", 1L),
    chains = 4, iter = 50, warmup = floor(iter/2), thin = 1,
    init = NULL, control = NULL, verbose = FALSE, num_level = 0)
```

# Arguments

guments	
compiled_banoc	c_model
	The compiled stan model (as with stan_model(model_code = banocc_model)).
С	The dataset as a data frame or matrix. This should be N by P with N samples as the rows and P features as the columns.
n	The prior mean for m; vectors of length less than P (the number of features/columns of C) will be recycled.
L	The prior variance-covariance for m (must be positive-definite with dimension PxP where P=number of features/columns in C), or a vector of length p of variances for m. If a vector of length less than P is given, it will be recycled.
a	The shape parameter of a gamma distribution (the prior on the shrinkage parameter lambda)
b	The rate parameter of a gamma distribution (the prior on the shrinkage parameter lambda)
cores	Number of cores to use when executing the chains in parallel, which defaults to 1 but we recommend setting the mc.cores option to be as many processors as the hardware and RAM allow (up to the number of chains).
chains	A positive integer specifying the number of Markov chains. The default is 4.
iter	A positive integer specifying the number of iterations for each chain (including warmup). The default is 2000.
warmup	A positive integer specifying the number of warmup (aka burnin) iterations per chain. If step-size adaptation is on (which it is by default), this also controls the number of iterations for which adaptation is run (and hence these warmup samples should not be used for inference). The number of warmup iterations should not be larger than iter and the default is iter/2.
thin	A positive integer specifying the period for saving samples. The default is 1, which is usually the recommended value.
init	The initial values as a list (see sampling in the rstan package). Default value is NULL, which means that initial values are sampled from the priors for parameters m and lambda while O is set to the identity matrix.
control	A named list of parameters to control the sampler's behavior. See the details in the documentation for the control argument in stan.
verbose	Print informative statements as the function executes?
num_level	The number of indentations to add to the output when verbose = TRUE.

run\_banocc

# Value

Returns a named list with the following elements:

**Data** The data formatted as a named list that includes the input data (C) and the prior parameters (n, L, a, b)

Fit The stanfit object returned by the call to sampling

## See Also

```
vignette("banocc-vignette") for more examples.
```

# **Examples**

# **Index**

```
* datasets
    banocc_model, 2
    {\tt compositions\_hard\_null, 3}
    compositions_neg_spike, 4
    compositions_null, 4
    compositions_pos_spike, 5
    counts_hard_null, 5
    counts_neg_spike, 6
    counts_null, 6
    counts_pos_spike, 7
banocc, 2
banocc-package (banocc), 2
banocc\_model, 2
compositions_hard_null, 3
compositions_neg_spike, 4
{\tt compositions\_null, 4}
compositions_pos_spike, 5
counts_hard_null, 5
counts_neg_spike, 6
counts_null, 6
counts_pos_spike, 7
get_banocc_output, 7
run_banocc, 8
sampling, 9, 10
stan, 9
```