

Package: sanba (via r-universe)

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Type Package

Title Fitting Shared Atoms Nested Models via MCMC or Variational Bayes

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Description An efficient tool for fitting nested mixture models based on a shared set of atoms via Markov Chain Monte Carlo and variational inference algorithms. Specifically, the package implements the common atoms model (Denti et al., 2023), its finite version (similar to D'Angelo et al., 2023), and a hybrid finite-infinite model (D'Angelo and Denti, 2026). All models implement univariate nested mixtures with Gaussian kernels equipped with a normal-inverse gamma prior distribution on the parameters. Additional functions are provided to help analyze the results of the fitting procedure. References: Denti, Camerlenghi, Guindani, Mira (2023) <[doi:10.1080/01621459.2021.1933499](https://doi.org/10.1080/01621459.2021.1933499)>, D'Angelo, Canale, Yu, Guindani (2023) <[doi:10.1111/biom.13626](https://doi.org/10.1111/biom.13626)>, D'Angelo, Denti (2026) <[doi:10.1214/24-BA1458](https://doi.org/10.1214/24-BA1458)>.

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Contents

compute_postdens	2
compute_psm	4
estimate_G	5
estimate_partition	6
fit_CAM	8
fit_fiSAN	12
fit_fSAN	16
get_model	19
number_clusters	21
plot.SANmcmc	22
plot.SANmcmc_postdens	24
plot.SANvi	25
plot_vi_allocation_prob	25
print.SANmcmc	26
print.SANvi	27
summary.SANmcmc	27
summary.SANvi	28

Index 29

compute_postdens	<i>Compute Posterior Predictive Density Samples for a Group</i>
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Description

Computes posterior predictive density estimates for a selected group from a fitted SAN model estimated via MCMC. For each retained MCMC iteration, the function evaluates the corresponding mixture density on a user-defined grid and stores the resulting density curves.

The returned object can be visualized with `plot.SANmcmc_postdens`, which displays the collection of posterior density curves together with pointwise posterior summaries.

Usage

```
compute_postdens(
  object,
  group_ind = 1,
  mcmc_considered = 500,
  lim = 1,
  length_yseq = 500
)
```

Arguments

object	An object of class SANmcmc, typically returned by fit_CAM(), fit_fiSAN(), or fit_fSAN() when est_method = "MCMC".
group_ind	Integer specifying the group for which posterior densities should be computed.
mcmc_considered	Number of MCMC iterations (counting backward from the last saved iteration) used to compute posterior density samples.
lim	Non-negative numeric value controlling the extension of the evaluation grid beyond the observed data range. The density is evaluated on seq(min(y) - lim, max(y) + lim, length.out = length_yseq).
length_yseq	Number of grid points used to evaluate the density.

Value

An object of class SANmcmc_postdens, containing:

mcmcdens A matrix of dimension length_yseq x mcmc_considered, where each column contains the posterior predictive density evaluated on the grid for one MCMC iteration.

x Numeric vector containing the evaluation grid.

YG Two-column matrix containing the observations and corresponding group labels for the selected group.

Examples

```
# Generate example data
set.seed(123)
y <- c(rnorm(100), rnorm(100, 5))
g <- rep(1:2, each = 100)

# Fit fiSAN via MCMC
est <- fit_fiSAN(y, g, est_method = "MCMC")

# Compute posterior density samples for group 1
postdens <- compute_postdens(
  est,
  group_ind = 1,
  mcmc_considered = 50
)

# Plot posterior density summaries
plot(postdens)
```

 compute_psm

Compute and Plot the Posterior Similarity Matrix

Description

The function computes and plots the posterior similarity matrix (PSM), either for the whole dataset, or separately for each group. The function takes as input an object from `fit_CAM`, `fit_fiSAN`, or `fit_fSAN`, used with the `est_method = "MCMC"` argument.

Usage

```
compute_psm(
  object,
  distributional = FALSE,
  group_specific = FALSE,
  plot = TRUE,
  ncores = 0
)
```

Arguments

<code>object</code>	An object of class <code>SANmcmc</code> .
<code>distributional</code>	Logical (default <code>FALSE</code>). If <code>FALSE</code> , the function computes the posterior similarity matrix (PSM) for the observational partition (i.e., between individual observations). If <code>TRUE</code> , it computes the PSM at the distributional level, that is, between groups.
<code>group_specific</code>	Logical (default <code>FALSE</code>). If <code>FALSE</code> , the function considers the overall PSM. If <code>TRUE</code> , the function considers the group-specific PSMs. This argument only affects the observational partition, i.e., when <code>distributional</code> is <code>FALSE</code> .
<code>plot</code>	Logical (default <code>TRUE</code>). Whether to plot the PSM.
<code>ncores</code>	A parameter to pass to the <code>salso::salso()</code> function. The number of CPU cores to use for parallel computing; a value of zero indicates the use of all cores of the system.

Value

The function `compute_psm` returns and plots the posterior similarity matrix. When `distributional = FALSE`, if `group_specific = FALSE`, the output is a matrix of dimension $N \times N$; if `group_specific = TRUE`, the output is a list of length J (the number of groups), where each entry contains a matrix of dimension $N_j \times N_j$. If `distributional = TRUE`, the output is a matrix of dimension $J \times J$.

Examples

```
# Generate example data
set.seed(123)
y <- c(rnorm(100), rnorm(100, 5))
```

```

g <- rep(1:2,rep(100,2))
plot(y,col=g)
# Fitting fiSAN via MCMC
est <- fit_fiSAN(y, g, est_method = "MCMC")
est
# Estimate PSM
psm_overall <- compute_psm(est)
# Estimate distributional PSM
psm_distrib <- compute_psm(est, distributional = TRUE)

```

estimate_G

Estimate the Atoms and Weights of the Discrete Mixing Distributions

Description

The function computes the posterior means of the atoms and weights characterizing the discrete mixing distributions. The function takes as input an object from `fit_CAM`, `fit_fiSAN`, or `fit_fSAN`, used with the `est_method = "VI"` argument, and returns an object of class `SANvi_G`.

Usage

```

estimate_G(object)

## S3 method for class 'SANvi_G'
plot(x, DC_num = NULL, lim = 2, ...)

## S3 method for class 'SANvi_G'
summary(object, thr = 0.01, ...)

## S3 method for class 'SANvi_G'
print(x, thr = 0.01, ...)

```

Arguments

<code>object</code>	an object of class <code>SANvi_G</code> (usually, the result of a call to <code>estimate_G</code>).
<code>x</code>	an object of class <code>SANvi_G</code> (usually, the result of a call to <code>estimate_G</code>).
<code>DC_num</code>	an integer or a vector of integers indicating which distributional clusters to plot.
<code>lim</code>	optional value for the <code>plot</code> method to adjust the limits of the x-axis (the default is 2). The atoms are plotted on a range given by $\min(\text{posterior means}) - \text{lim}$, $\max(\text{posterior means}) + \text{lim}$.
<code>...</code>	ignored.
<code>thr</code>	argument for the <code>print</code> method. It should be a small positive number, representing a threshold. If the posterior weight of a specific shared atom is below the threshold, the atom is not reported.

Value

The function `estimate_G` returns an object of class `SANvi_G`, which is a matrix comprising the posterior means, variances, and weights of each estimated DC (one mixture component for each row).

Examples

```
# Generate example data
set.seed(123)
y <- c(rnorm(100),rnorm(100,5))
g <- rep(1:2,rep(100,2))
plot(y,col=g)
# Fitting fiSAN via variational inference
est <- fit_fiSAN(y,g,vi_param= list(n_runs = 10))
est
summary(est)
# Estimate posterior atoms and weights
G <- estimate_G(est)
summary(G)
```

estimate_partition *Estimate the Observational and Distributional Partition*

Description

Given the output of a `sanba` model-fitting function, this method estimates both the observational and distributional partitions. For MCMC objects, it computes a point estimate using `salso::salso()`; for Variational Inference (VI) objects, the cluster allocation is determined by the label with the highest estimated variational probability.

Usage

```
estimate_partition(object, ...)

## S3 method for class 'SANvi'
estimate_partition(object, ordered = TRUE, ...)

## S3 method for class 'SANmcmc'
estimate_partition(object, ordered = TRUE, add_burnin = 0, ncores = 0, ...)

## S3 method for class 'partition_mcmc'
summary(object, ...)

## S3 method for class 'partition_vi'
summary(object, ...)

## S3 method for class 'partition_mcmc'
print(x, ...)
```

```

## S3 method for class 'partition_vi'
print(x, ...)

## S3 method for class 'partition_mcmc'
plot(
  x,
  DC_num = NULL,
  type = c("ecdf", "boxplot", "scatter"),
  alt_palette = FALSE,
  ...
)

## S3 method for class 'partition_vi'
plot(
  x,
  DC_num = NULL,
  type = c("ecdf", "boxplot", "scatter"),
  alt_palette = FALSE,
  ...
)

```

Arguments

object	Object of class SANmcmc (usually, the result of a call to fit_fiSAN , fit_fSAN , or fit_CAM with <code>est_method = "MCMC"</code>) or SANvi (the result of a call to fit_fiSAN , fit_fSAN , or fit_CAM with <code>est_method = "VI"</code>).
...	Additional graphical parameters to be passed to the plot function.
ordered	Logical, if TRUE (default), the function sorts the distributional cluster labels reflecting the increasing values of medians of the data assigned to each DC. If FALSE, no ordering is applied.
add_burnin	Integer (default = 0). Number of observations to discard as additional burn-in (only for SANmcmc objects).
ncores	A parameter to pass to the <code>salso::salso()</code> function (only for SANmcmc objects). The number of CPU cores to use for parallel computing; a value of zero indicates the use of all cores on the system.
x	The result of a call to estimate_partition .
DC_num	An integer or a vector of integers indicating which distributional clusters to plot.
type	What type of plot should be drawn. Available types are "boxplot", "ecdf", and "scatter".
alt_palette	Logical, the color palette to be used. Default is R base colors (<code>alt_palette = FALSE</code>).

Value

A list of class `partition_vi` or `partition_mcmc` containing

- `obs_level`: a data frame containing the data values, their group indexes, and the observational and distributional clustering assignments for each observation.
- `dis_level`: a vector with the distributional clustering assignment for each unit.

See Also

[salso::salso\(\)](#)

Examples

```
set.seed(123)
y <- c(rnorm(40,0,0.3), rnorm(20,5,0.3))
g <- c(rep(1:6, each = 10))
out <- fit_fSAN(y = y, group = g, "VI", vi_param = list(n_runs = 10))
plot(out)
clust <- estimate_partition(out)
summary(clust)
plot(clust, lwd = 2, alt_palette = TRUE)
plot(clust, type = "scatter", alt_palette = FALSE, cex = 2)

set.seed(123)
y <- c(rnorm(40,0,0.3), rnorm(20,5,0.3))
g <- c(rep(1:6, each = 10))
out <- fit_fSAN(y = y, group = g, "MCMC", mcmc_param=list(nrep=500,burn=200))
plot(out)
clust <- estimate_partition(out)
summary(clust)
plot(clust, lwd = 2)
plot(clust, type = "boxplot", alt_palette = TRUE)
plot(clust, type = "scatter", alt_palette = TRUE, cex = 2, pch = 4)
```

fit_CAM

Fit the Common Atoms Mixture Model

Description

`fit_CAM` fits the common atoms mixture model (CAM) of Denti et al. (2023) with Gaussian kernels and normal-inverse gamma priors on the unknown means and variances. The function returns an object of class `SANmcmc` or `SANvi` depending on the chosen computational approach (MCMC or VI).

Usage

```
fit_CAM(y, group, est_method = c("VI", "MCMC"),
        prior_param = list(),
        vi_param = list(),
        mcmc_param = list())
```

Arguments

y	Numerical vector of observations (required).
group	Numerical vector of the same length of y, indicating the group membership (required).
est_method	Character, specifying the preferred estimation method. It can be either "VI" or "MCMC".
prior_param	<p>A list containing</p> <p>$m_0, \tau_0, \lambda_0, \gamma_0$ Hyperparameters on $(\mu, \sigma^2) \sim NIG(m_0, \tau_0, \lambda_0, \gamma_0)$. The default is (0, 0.01, 3, 2).</p> <p>hyp_alpha1, hyp_alpha2 If a random α is used, (hyp_alpha1, hyp_alpha2) specify the hyperparameters. The default is (1,1). The prior is $\alpha \sim \text{Gamma}(\text{hyp_alpha1}, \text{hyp_alpha2})$.</p> <p>alpha Distributional DP parameter if fixed (optional). The distribution is $\pi \sim GEM(\alpha)$.</p> <p>hyp_beta1, hyp_beta2 If a random β is used, (hyp_beta1, hyp_beta2) specify the hyperparameters. The default is (1,1). The prior is $\beta \sim \text{Gamma}(\text{hyp_beta1}, \text{hyp_beta2})$.</p> <p>beta Observational DP parameter if fixed (optional). The distribution is $\omega_k \sim GEM(\beta)$.</p>
vi_param	<p>A list of variational inference-specific settings containing</p> <p>maxL, maxK Integers, the upper bounds for the observational and distributional clusters to fit, respectively. The default is (50, 20).</p> <p>epsilon The threshold controlling the convergence criterion.</p> <p>n_runs Number of starting points considered for the estimation.</p> <p>seed Random seed to control the initialization.</p> <p>maxSIM The maximum number of CAVI iterations to perform.</p> <p>warmstart Logical, if TRUE, the observational means of the cluster atoms are initialized with a k-means algorithm.</p> <p>verbose Logical, if TRUE the iterations are printed.</p>
mcmc_param	<p>A list of MCMC inference-specific settings containing</p> <p>nrep, burn Integers, the number of total MCMC iterations, and the number of discarded iterations, respectively.</p> <p>maxL, maxK Integers, the upper bounds for the observational and distributional clusters to fit, respectively. The default is (50, 20).</p> <p>seed Random seed to control the initialization.</p> <p>warmstart Logical, if TRUE, the observational means of the cluster atoms are initialized with a k-means algorithm. If FALSE, the starting points can be passed through the parameters nclus_start, mu_start, sigma2_start, M_start, S_start, alpha_start, beta_start</p> <p>verbose Logical, if TRUE the iterations are printed.</p>

Details

The common atoms mixture model is used to perform inference in nested settings, where the data are organized into J groups. The data should be continuous observations (Y_1, \dots, Y_J) , where each $Y_j = (y_{1,j}, \dots, y_{n_j,j})$ contains the n_j observations from group j , for $j = 1, \dots, J$. The function takes as input the data as a numeric vector y in this concatenated form. Hence, y should be a vector of length $n_1 + \dots + n_J$. The group parameter is a numeric vector of the same size as y , indicating the group membership for each individual observation. Notice that with this specification, the observations in the same group need not be contiguous as long as the correspondence between the variables y and group is maintained.

Model

The data are modeled using a Gaussian likelihood, where both the mean and the variance are observational cluster-specific, i.e.,

$$y_{i,j} \mid M_{i,j} = l \sim N(\mu_l, \sigma_l^2)$$

where $M_{i,j} \in \{1, 2, \dots\}$ is the observational cluster indicator of observation i in group j . The prior on the model parameters is a normal-inverse gamma distribution $(\mu_l, \sigma_l^2) \sim NIG(m_0, \tau_0, \lambda_0, \gamma_0)$, i.e., $\mu_l \mid \sigma_l^2 \sim N(m_0, \sigma_l^2/\tau_0)$, $1/\sigma_l^2 \sim \text{Gamma}(\lambda_0, \gamma_0)$ (shape, rate).

Clustering

The model clusters both observations and groups. The clustering of groups (distributional clustering) is provided by the allocation variables $S_j \in \{1, 2, \dots\}$, with

$$Pr(S_j = k \mid \dots) = \pi_k \quad \text{for } k = 1, 2, \dots$$

The distribution of the probabilities is $\{\pi_k\}_{k=1}^\infty \sim GEM(\alpha)$, where GEM is the Griffiths-Engen-McCloskey distribution of parameter α , which characterizes the stick-breaking construction of the DP (Sethuraman, 1994).

The clustering of observations (observational clustering) is provided by the allocation variables $M_{i,j} \in \{1, 2, \dots\}$, with

$$Pr(M_{i,j} = l \mid S_j = k, \dots) = \omega_{l,k} \quad \text{for } k = 1, 2, \dots; l = 1, 2, \dots$$

The distribution of the probabilities is $\{\omega_{l,k}\}_{l=1}^\infty \sim GEM(\beta)$ for all $k = 1, 2, \dots$

Value

fit_CAM returns a list of class SANvi, if est_method = "VI", or SANmcmc, if est_method = "MCMC". The list contains the following elements:

model Name of the fitted model.

params List containing the data and the parameters used in the simulation. Details below.

sim List containing the optimized variational parameters or the simulated values. Details below.

time Total computation time.

Data and parameters: params is a list with the following components:

- y, group, Nj, J: Data, group labels, group frequencies, and number of groups.
- K, L: Number of distributional and observational mixture components.

- $m\theta$, $\tau\theta$, $\lambda\theta$, $\gamma\theta$: Model hyperparameters.
- (hyp_alpha1, hyp_alpha2) or alpha: hyperparameters on α (if α random); or provided value for α (if fixed).
- (hyp_beta1, hyp_beta2) or beta: hyperparameters on β (if β random); or provided value for β (if fixed).
- seed: The random seed adopted to replicate the run.
- epsilon, n_runs: The threshold controlling the convergence criterion and the number of starting points considered.
- nrep, burnin: If est_method = "MCMC", the number of total MCMC iterations, and the number of discarded ones.

Simulated values: depending on the algorithm, it returns a list with the optimized variational parameters or a list with the chains of the simulated values.

Variational inference: sim is a list with the following components:

- theta_l: Matrix of size (maxL, 4). Each row is a posterior variational estimate of the four normal-inverse gamma hyperparameters.
- XI: A list of length J. Each element is a matrix of size (Nj, maxL) containing the posterior variational probability of assignment of the i-th observation in the j-th group to the l-th OC, i.e., $\hat{\xi}_{i,j,l} = \hat{Q}(M_{i,j} = l)$.
- RHO: Matrix of size (J, maxK). Each row is a posterior variational probability of assignment of the j-th group to the k-th DC, i.e., $\hat{\rho}_{j,k} = \hat{Q}(S_j = k)$.
- a_tilde_k, b_tilde_k: Vector of updated variational parameters of the beta distributions governing the distributional stick-breaking process.
- a_bar_lk, b_bar_lk: Matrix of updated variational parameters of the beta distributions governing the observational stick-breaking process (arranged by column).
- conc_hyper: If the concentration parameters are chosen to be random, a vector with the four updated hyperparameters.
- Elbo_val: Vector containing the values of the ELBO.

MCMC inference: sim is a list with the following components:

- mu: Matrix of size (nrep, maxL). Each row is a posterior sample of the mean parameter for each observational cluster (μ_1, \dots, μ_L).
- sigma2: Matrix of size (nrep, maxL). Each row is a posterior sample of the variance parameter for each observational cluster ($\sigma_1^2, \dots, \sigma_L^2$).
- obs_cluster: Matrix of size (nrep, n), with $n = \text{length}(y)$. Each row is a posterior sample of the observational cluster allocation variables ($M_{1,1}, \dots, M_{n,J}$).
- distr_cluster: Matrix of size (nrep, J), with $J = \text{length}(\text{unique}(\text{group}))$. Each row is a posterior sample of the distributional cluster allocation variables (S_1, \dots, S_J).
- pi: Matrix of size (nrep, maxK). Each row is a posterior sample of the distributional cluster probabilities ($\pi_1, \dots, \pi_{\text{maxK}}$).
- omega: 3-d array of size (maxL, maxK, nrep). Each slice is a posterior sample of the observational cluster probabilities. In each slice, each column k is a vector (of length maxL) observational cluster probabilities ($\omega_{1,k}, \dots, \omega_{\text{maxL},k}$) for distributional cluster k .

- alpha: Vector of length nrep of posterior samples of the parameter α .
- beta: Vector of length nrep of posterior samples of the parameter β .
- maxK: Vector of length nrep of the number of distributional DP components used by the slice sampler.
- maxL: Vector of length nrep of the number of observational DP components used by the slice sampler.

References

- Denti, F., Camerlenghi, F., Guindani, M., and Mira, A. (2023). A Common Atoms Model for the Bayesian Nonparametric Analysis of Nested Data. *Journal of the American Statistical Association*, 118(541), 405-416. DOI: 10.1080/01621459.2021.1933499
- Sethuraman, A.J. (1994). A Constructive Definition of Dirichlet Priors, *Statistica Sinica*, 4, 639–650.

Examples

```
set.seed(123)
y <- c(rnorm(60), rnorm(40, 5))
g <- rep(1:2, rep(50, 2))

out_vi <- fit_CAM(y, group = g, est_method = "VI", vi_param = list(n_runs = 1,
  epsilon = .01 ))
out_vi

out_mcmc <- fit_CAM(y = y, group = g, est_method = "MCMC",
  mcmc_param = list(nrep = 50, burn = 20))
out_mcmc
```

 fit_fiSAN

Fit the Finite-Infinite Shared Atoms Mixture Model

Description

fit_fiSAN fits the finite-infinite shared atoms nested (fiSAN) mixture model with Gaussian kernels and normal-inverse gamma priors on the unknown means and variances. The function returns an object of class SANmcmc or SANvi depending on the chosen computational approach (MCMC or VI).

Usage

```
fit_fiSAN(y, group, est_method = c("VI", "MCMC"),
  prior_param = list(),
  vi_param = list(),
  mcmc_param = list())
```

Arguments

<code>y</code>	Numerical vector of observations (required).
<code>group</code>	Numerical vector of the same length of <code>y</code> , indicating the group membership (required).
<code>est_method</code>	Character, specifying the preferred estimation method. It can be either "VI" or "MCMC".
<code>prior_param</code>	A list containing: <code>m0, tau0, lambda0, gamma0</code> Hyperparameters on $(\mu, \sigma^2) \sim NIG(m_0, \tau_0, \lambda_0, \gamma_0)$. The default is (0, 0.01, 3, 2). <code>hyp_alpha1, hyp_alpha2</code> If a random α is used, (<code>hyp_alpha1, hyp_alpha2</code>) specify the hyperparameters. The default is (1,1). The prior is $\alpha \sim \text{Gamma}(\text{hyp_alpha1}, \text{hyp_alpha2})$. <code>alpha</code> Distributional DP parameter if fixed (optional). The distribution is $\pi \sim \text{GEM}(\alpha)$. <code>b_dirichlet</code> The hyperparameter of the symmetric observational Dirichlet distribution. The default is $1/\text{maxL}$.
<code>vi_param</code>	A list of variational inference-specific settings containing: <code>maxL, maxK</code> Integers, the upper bounds for the observational and distributional clusters to fit, respectively. The default is (50, 20). <code>epsilon</code> The threshold controlling the convergence criterion. <code>n_runs</code> Number of starting points considered for the estimation. <code>seed</code> Random seed to control the initialization. <code>maxSIM</code> The maximum number of CAVI iterations to perform. <code>warmstart</code> Logical, if TRUE, the observational means of the cluster atoms are initialized with a k-means algorithm. <code>verbose</code> Logical, if TRUE the iterations are printed.
<code>mcmc_param</code>	A list of MCMC inference-specific settings containing: <code>nrep, burn</code> Integers, the number of total MCMC iterations, and the number of discarded iterations, respectively. <code>maxL, maxK</code> Integers, the upper bounds for the observational and distributional clusters to fit, respectively. The default is (50, 20). <code>seed</code> Random seed to control the initialization. <code>warmstart</code> Logical, if TRUE, the observational means of the cluster atoms are initialized with a k-means algorithm. If FALSE, the starting points can be passed through the parameters <code>nclus_start, mu_start, sigma2_start, M_start, S_start, alpha_start</code> . <code>verbose</code> Logical, if TRUE the iterations are printed.

Details**Data structure**

The finite-infinite common atoms mixture model is used to perform inference in nested settings, where the data are organized into J groups. The data should be continuous observations (Y_1, \dots, Y_J) ,

where each $Y_j = (y_{1,j}, \dots, y_{n_j,j})$ contains the n_j observations from group j , for $j = 1, \dots, J$. The function takes as input the data as a numeric vector y in this concatenated form. Hence, y should be a vector of length $n_1 + \dots + n_J$. The group parameter is a numeric vector of the same size as y , indicating the group membership for each individual observation. Notice that with this specification, the observations in the same group need not be contiguous as long as the correspondence between the variables y and group is maintained.

Model

The data are modeled using a Gaussian likelihood, where both the mean and the variance are observational-cluster-specific:

$$y_{i,j} \mid M_{i,j} = l \sim N(\mu_l, \sigma_l^2)$$

where $M_{i,j} \in \{1, \dots, L\}$ is the observational cluster indicator of observation i in group j . The prior on the model parameters is a normal-inverse gamma distribution $(\mu_l, \sigma_l^2) \sim NIG(m_0, \tau_0, \lambda_0, \gamma_0)$, i.e., $\mu_l \mid \sigma_l^2 \sim N(m_0, \sigma_l^2/\tau_0)$, $1/\sigma_l^2 \sim \text{Gamma}(\lambda_0, \gamma_0)$ (shape, rate).

Clustering

The model clusters both observations and groups. The clustering of groups (distributional clustering) is provided by the allocation variables $S_j \in \{1, 2, \dots\}$, with:

$$Pr(S_j = k \mid \dots) = \pi_k \quad \text{for } k = 1, 2, \dots$$

The distribution of the probabilities is $\{\pi_k\}_{k=1}^{\infty} \sim GEM(\alpha)$, where GEM is the Griffiths-Engen-McCloskey distribution of parameter α , which characterizes the stick-breaking construction of the DP (Sethuraman, 1994).

The clustering of observations (observational clustering) is provided by the allocation variables $M_{i,j} \in \{1, \dots, L\}$, with:

$$Pr(M_{i,j} = l \mid S_j = k, \dots) = \omega_{l,k} \quad \text{for } k = 1, 2, \dots; l = 1, \dots, L.$$

The distribution of the probabilities is $(\omega_{1,k}, \dots, \omega_{L,k}) \sim \text{Dirichlet}_L(b, \dots, b)$ for all $k = 1, 2, \dots$. Here, the dimension L is fixed.

Value

`fit_fiSAN` returns a list of class `SANvi`, if `est_method = "VI"`, or `SANmcmc`, if `est_method = "MCMC"`. The list contains the following elements:

`model` Name of the fitted model.

`params` List containing the data and the parameters used in the simulation. Details below.

`sim` List containing the optimized variational parameters or the simulated values. Details below.

`time` Total computation time.

Data and parameters: `params` is a list with the following components:

- `y`, `group`, `Nj`, `J`: Data, group labels, group frequencies, and number of groups.
- `K`, `L`: Number of distributional and observational mixture components.
- `m0`, `tau0`, `lambda0`, `gamma0`: Model hyperparameters.
- `(hyp_alpha1, hyp_alpha2)` or `alpha`: Hyperparameters on α (if α random); or provided value for α (if fixed).

- `b_dirichlet`: Provided value for b .
- `seed`: The random seed adopted to replicate the run.
- `epsilon`, `n_runs`: If `est_method = "VI"`, the threshold controlling the convergence criterion and the number of starting points considered.
- `nrep`, `burnin`: If `est_method = "MCMC"`, the number of total MCMC iterations, and the number of discarded ones.

Simulated values: Depending on the algorithm, it returns a list with the optimized variational parameters or a list with the chains of the simulated values.

Variational inference: `sim` is a list with the following components:

- `theta_l`: Matrix of size $(\text{maxL}, 4)$. Each row is a posterior variational estimate of the four normal-inverse gamma hyperparameters.
- `XI`: A list of length J . Each element is a matrix of size (N_j, maxL) , the posterior variational assignment probabilities $\hat{Q}(M_{i,j} = l)$.
- `RHO`: Matrix of size (J, maxK) , with the posterior variational assignment probabilities $\hat{Q}(S_j = k)$.
- `a_tilde_k`, `b_tilde_k`: Vector of updated variational parameters of the beta distributions governing the distributional stick-breaking process.
- `conc_hyper`: If the concentration parameter is random, this contains its updated hyperparameters.
- `b_dirichlet_lk`: Matrix of updated variational parameters of the Dirichlet distributions governing observational clustering.
- `Elbo_val`: Vector containing the values of the ELBO.

MCMC inference: `sim` is a list with the following components:

- `mu`: Matrix of size $(\text{nrep}, \text{maxL})$ with samples of the observational cluster means.
- `sigma2`: Matrix of size $(\text{nrep}, \text{maxL})$ with samples of the observational cluster variances.
- `obs_cluster`: Matrix of size (nrep, n) with posterior samples of the observational cluster allocations.
- `distr_cluster`: Matrix of size (nrep, J) with posterior samples of the distributional cluster allocations.
- `pi`: Matrix of size $(\text{nrep}, \text{maxK})$ with posterior samples of the distributional cluster weights.
- `omega`: Array of size $(\text{maxL}, \text{maxK}, \text{nrep})$ with observational cluster weights.
- `alpha`: Vector of length `nrep` with posterior samples of α .
- `maxK`: Vector of length `nrep` with number of active distributional components.

Examples

```
set.seed(123)
y <- c(rnorm(60), rnorm(40, 5))
g <- rep(1:2, rep(50, 2))
plot(density(y[g==1]), xlim = c(-5,10), main = "Group-specific density")
lines(density(y[g==2]), col = 2)
```

```

out_vi <- fit_fiSAN(y, group = g, est_method = "VI",
                  vi_param = list(n_runs = 1))
out_vi

out_mcmc <- fit_fiSAN(y = y, group = g, est_method = "MCMC")
out_mcmc

```

fit_fSAN

Fit the Finite Shared Atoms Mixture Model

Description

fit_fSAN fits the finite shared atoms nested (fSAN) mixture model with Gaussian kernels and normal-inverse gamma priors on the unknown means and variances. The function returns an object of class SANmcmc or SANvi depending on the chosen computational approach (MCMC or VI).

Usage

```

fit_fSAN(y, group, est_method = c("VI", "MCMC"),
        prior_param = list(),
        vi_param = list(),
        mcmc_param = list())

```

Arguments

y	Numerical vector of observations (required).
group	Numerical vector of the same length of y, indicating the group membership (required).
est_method	Character, specifying the preferred estimation method. It can be either "VI" or "MCMC".
prior_param	A list containing m_0 , τ_0 , λ_0 , γ_0 Hyperparameters on $(\mu, \sigma^2) \sim NIG(m_0, \tau_0, \lambda_0, \gamma_0)$. The default is (0, 0.01, 3, 2). a_dirichlet The hyperparameter of the symmetric distributional Dirichlet distribution. The default is $1/\max K$. b_dirichlet The hyperparameter of the symmetric observational Dirichlet distribution. The default is $1/\max L$.
vi_param	A list of variational inference-specific settings, containing maxL, maxK Integers, the upper bounds for the observational and distributional clusters to fit, respectively. The default is (50, 20). epsilon The threshold controlling the convergence criterion. n_runs Number of starting points considered for the estimation. seed Random seed to control the initialization. maxSIM The maximum number of CAVI iteration to perform.

	warmstart	Logical, if TRUE, the observational means of the cluster atoms are initialized with a k-means algorithm.
	verbose	Logical, if TRUE the iterations are printed.
mcmc_param		A list of MCMC inference-specific settings, containing
	nrep, burn	Integers, the number of total MCMC iterations, and the number of discarded iterations, respectively.
	maxL, maxK	Integers, the upper bounds for the observational and distributional clusters to fit, respectively. The default is (50, 20).
	seed	Random seed to control the initialization.
	warmstart	Logical, if TRUE, the observational means of the cluster atoms are initialized with a k-means algorithm. If FALSE, the starting points can be passed through the parameters nclus_start, mu_start, sigma2_start, M_start, S_start
	verbose	Logical, if TRUE the iterations are printed.

Details

Data structure

The finite common atoms mixture model is used to perform inference in nested settings, where the data are organized into J groups. The data should be continuous observations (Y_1, \dots, Y_J) , where each $Y_j = (y_{1,j}, \dots, y_{n_j,j})$ contains the n_j observations from group j , for $j = 1, \dots, J$. The function takes as input the data as a numeric vector \mathbf{y} in this concatenated form. Hence \mathbf{y} should be a vector of length $n_1 + \dots + n_J$. The group parameter is a numeric vector of the same size as \mathbf{y} indicating the group membership for each individual observation. Notice that with this specification the observations in the same group need not be contiguous as long as the correspondence between the variables \mathbf{y} and group is maintained.

Model

The data are modeled using a Gaussian likelihood, where both the mean and the variance are observational-cluster-specific, i.e.,

$$y_{i,j} \mid M_{i,j} = l \sim N(\mu_l, \sigma_l^2)$$

where $M_{i,j} \in \{1, \dots, L\}$ is the observational cluster indicator of observation i in group j . The prior on the model parameters is a normal-inverse gamma distribution $(\mu_l, \sigma_l^2) \sim NIG(m_0, \tau_0, \lambda_0, \gamma_0)$, i.e., $\mu_l \mid \sigma_l^2 \sim N(m_0, \sigma_l^2/\tau_0)$, $1/\sigma_l^2 \sim \text{Gamma}(\lambda_0, \gamma_0)$ (shape, rate).

Clustering

The model performs a clustering of both observations and groups. The clustering of groups (distributional clustering) is provided by the allocation variables $S_j \in \{1, \dots, K\}$, with

$$Pr(S_j = k \mid \dots) = \pi_k \quad \text{for } k = 1, \dots, K.$$

The distribution of the probabilities is $(\pi_1, \dots, \pi_K) \sim \text{Dirichlet}_K(a, \dots, a)$. Here, the dimension K is fixed.

The clustering of observations (observational clustering) is provided by the allocation variables $M_{i,j} \in \{1, \dots, L\}$, with

$$Pr(M_{i,j} = l \mid S_j = k, \dots) = \omega_{l,k} \quad \text{for } k = 1, \dots, K; l = 1, \dots, L.$$

The distribution of the probabilities is $(\omega_{1,k}, \dots, \omega_{L,k}) \sim \text{Dirichlet}_L(b, \dots, b)$ for all $k = 1, \dots, K$. Here, the dimension L is fixed.

Value

fit_fSAN returns a list of class SANvi, if est_method = "VI", or SANmcmc, if est_method = "MCMC". The list contains the following elements:

model Name of the fitted model.
 params List containing the data and the parameters used in the simulation. Details below.
 sim List containing the optimized variational parameters or the simulated values. Details below.
 time Total computation time.

Data and parameters: params is a list with the following components:

- y, group, Nj, J: Data, group labels, group frequencies, and number of groups.
- K, L: Number of distributional and observational mixture components.
- m0, tau0, lambda0, gamma0: Model hyperparameters.
- a_dirichlet: Provided value for a .
- b_dirichlet: Provided value for b .
- seed: The random seed adopted to replicate the run.
- epsilon, n_runs: The threshold controlling the convergence criterion and the number of starting points considered
- nrep, burnin: If est_method = "MCMC", the number of total MCMC iterations, and the number of discarded ones.

Simulated values: depending on the algorithm, it returns a list with the optimized variational parameters or a list with the chains of the simulated values.

Variational inference: sim is a list with the following components:

- theta_l: Matrix of size (maxL, 4). Each row is a posterior variational estimate of the four normal-inverse gamma hyperparameters.
- XI : A list of length J. Each element is a matrix of size (Nj, maxL) posterior variational probability of assignment of assignment of the i-th observation in the j-th group to the l-th OC, i.e., $\hat{\xi}_{i,j,l} = \hat{Q}(M_{i,j} = l)$.
- RHO: Matrix of size (J, maxK). Each row is a posterior variational probability of assignment of the j-th group to the k-th DC, i.e., $\hat{\rho}_{j,k} = \hat{Q}(S_j = k)$.
- a_dirichlet_k: Vector of updated variational parameters of the Dirichlet distribution governing the distributional clustering.
- b_dirichlet_lk: Matrix of updated variational parameters of the Dirichlet distributions governing the observational clustering (arranged by column).
- Elbo_val: Vector containing the values of the ELBO.

MCMC inference: sim is a list with the following components:

- mu: Matrix of size (nrep, maxL). Each row is a posterior sample of the mean parameter of each observational cluster (μ_1, \dots, μ_L) .
- sigma2: Matrix of size (nrep, maxL). Each row is a posterior sample of the variance parameter of each observational cluster $(\sigma_1^2, \dots, \sigma_L^2)$.

- `obs_cluster`: Matrix of size $(nrep, n)$, with $n = \text{length}(y)$. Each row is a posterior sample of the observational cluster allocation variables $(M_{1,1}, \dots, M_{n,J})$.
- `distr_cluster`: Matrix of size $(nrep, J)$, with $J = \text{length}(\text{unique}(\text{group}))$. Each row is a posterior sample of the distributional cluster allocation variables (S_1, \dots, S_J) .
- `pi`: Matrix of size $(nrep, \text{maxK})$. Each row is a posterior sample of the distributional cluster probabilities $(\pi_1, \dots, \pi_{\text{maxK}})$.
- `omega`: 3-d array of size $(\text{maxL}, \text{maxK}, nrep)$. Each slice is a posterior sample of the observational cluster probabilities. In each slice, each column k is a vector (of length maxL) observational cluster probabilities $(\omega_{1,k}, \dots, \omega_{\text{maxL},k})$ for distributional cluster k .

Examples

```
set.seed(123)
y <- c(rnorm(60), rnorm(40, 5))
g <- rep(1:2, rep(50, 2))
plot(density(y[g==1]), xlim = c(-5,10), main = "Group-specific density")
lines(density(y[g==2]), col = 2)

out_vi <- fit_fSAN(y, group = g, est_method = "VI", vi_param = list(n_runs = 1))
out_vi

out_mcmc <- fit_fSAN(y = y, group = g, est_method = "MCMC",
                    mcmc_param = list(nrep = 100, burn= 50))
out_mcmc
```

get_model

Accessor Functions for SAN Model Outputs

Description

The functions `get_model`, `get_time`, `get_params`, `get_sim`, and `get_seed_best_run` provide convenient access to specific components of model output objects of class `SANmcmc` (fitted via MCMC) or `SANvi` (fitted via variational inference).

Specifically:

- `get_model`: returns a character string specifying the model type used;
- `get_time`: returns the total runtime of the estimation procedure;
- `get_params`: returns a list containing data and prior hyperparameters;
- `get_sim`: returns the fitted quantities, such as posterior draws (MCMC) or variational estimates (VI);
- `get_seed_best_run`: (only for `SANvi`) returns the random seed associated with the run that achieved the highest ELBO.

Usage

```
get_model(object, ...)

## S3 method for class 'SANvi'
get_model(object, ...)

## S3 method for class 'SANmcmc'
get_model(object, ...)

get_time(object, ...)

## S3 method for class 'SANvi'
get_time(object, ...)

## S3 method for class 'SANmcmc'
get_time(object, ...)

get_params(object, ...)

## S3 method for class 'SANvi'
get_params(object, ...)

## S3 method for class 'SANmcmc'
get_params(object, ...)

get_sim(object, ...)

## S3 method for class 'SANvi'
get_sim(object, ...)

## S3 method for class 'SANmcmc'
get_sim(object, ...)

get_seed_best_run(object, ...)

## S3 method for class 'SANvi'
get_seed_best_run(object, ...)
```

Arguments

object	An object of class SANmcmc or SANvi, as returned by fit_fSAN , fit_fiSAN , or fit_CAM .
...	ignored.

Value

The requested component from the fitted model object. See the function descriptions above for details.

Examples

```
set.seed(123)
y <- c(rnorm(40, 0, 0.3), rnorm(20, 5, 0.3))
g <- c(rep(1:6, each = 10))

out <- fit_fSAN(y = y, group = g, est_method = "MCMC",
               mcmc_param = list(nrep = 500, burn = 200))

get_model(out)
get_time(out)
hp <- get_params(out)
sims <- get_sim(out)
```

number_clusters

Estimate the Number of Observational and Distributional Clusters

Description

Computes the estimated number of observational clusters (OC) and distributional clusters (DC) from a fitted SAN model object.

For variational inference (SANvi objects), the function returns point estimates based on posterior mode assignments. For MCMC-based inference (SANmcmc objects), it returns the mean, median, and variance of the number of clusters across posterior samples.

Usage

```
number_clusters(object, ...)
```

Arguments

object	An object of class SANvi or SANmcmc, typically, the output of a call to <code>fit_fiSAN</code> , <code>fit_fSAN</code> , or <code>fit_CAM</code> .
...	ignored.

Value

A data frame reporting the estimated number of observational (OC) and distributional (DC) clusters.

- For SANvi: a single-row data frame with the point estimates.
- For SANmcmc: a three-row data frame with the mean, median, and variance across MCMC samples.

Examples

```
# Generate example data
set.seed(123)
y <- c(rnorm(60), rnorm(40, 5))
g <- rep(1:2, each = 50)

plot(density(y[g == 1]), xlim = c(-5, 10), main = "Group-specific density")
lines(density(y[g == 2]), col = 2)

# Fit fiSAN via MCMC
est_mcmc <- fit_fiSAN(y, g, est_method = "MCMC")
number_clusters(est_mcmc)

# Fit fiSAN via Variational Inference
est_vi <- fit_fiSAN(y, g, est_method = "VI")
number_clusters(est_vi)
```

plot.SANmcmc

Visual Check of the Convergence of the MCMC Output

Description

Plot method for objects of class `SANmcmc`. Check the convergence of the MCMC through visual inspection of the chains.

Usage

```
## S3 method for class 'SANmcmc'
plot(
  x,
  param = c("mu", "sigma2", "pi", "num_clust", "alpha", "beta"),
  show_density = TRUE,
  add_burnin = 0,
  show_convergence = TRUE,
  trunc_plot = 2,
  ...
)
```

Arguments

<code>x</code>	Object of class <code>SANmcmc</code> (usually, the result of a call to <code>fit_CAM</code> , <code>fit_fiSAN</code> , or <code>fit_fSAN</code> , used with the <code>est_method = "MCMC"</code> argument).
<code>param</code>	String with the names of the parameters to check. It can be one of <code>"mu"</code> , <code>"sigma2"</code> , <code>"pi"</code> , <code>"num_clust"</code> , <code>"alpha"</code> , <code>"beta"</code> .
<code>show_density</code>	Logical (default <code>TRUE</code>). Should a kernel estimate of the density be plotted?

add_burnin	Integer (default = 0). Additional number of observations to discard in the burn-in.
show_convergence	Logical (default TRUE). Should a superimposed red line of the cumulative mean be plotted?
trunc_plot	Integer (default = 10). For multidimensional parameters, the maximum number of components to be plotted.
...	Ignored.

Value

The function displays the traceplots and posterior density estimates of the parameters sampled in the MCMC algorithm.

Note

The function is not available for the observational weights ω .

Examples

```
set.seed(123)
y <- c(rnorm(40,0,0.3), rnorm(20,5,0.3))
g <- c(rep(1,30), rep(2, 30))
out <- fit_fiSAN(y = y, group = g, "MCMC", mcmc_param = list(nrep = 500, burn = 200))
plot(out, param = "mu", trunc_plot = 2)
plot(out, param = "sigma2", trunc_plot = 2)
plot(out, param = "alpha", trunc_plot = 1)
plot(out, param = "alpha", add_burnin = 100)
plot(out, param = "pi", trunc_plot = 4, show_density = FALSE)

out <- fit_CAM(y = y, group = g, "MCMC",
mcmc_param = list(nrep = 500, burn = 200, seed= 1234))
plot(out, param = "mu", trunc_plot = 2)
plot(out, param = "sigma2", trunc_plot = 2)
plot(out, param = "alpha")
plot(out, param = "pi", trunc_plot = 2)
plot(out, param = "pi", trunc_plot = 5)
plot(out, param = "num_clust", trunc_plot = 5)
plot(out, param = "beta", trunc_plot = 2)

out <- fit_fSAN(y = y, group = g, "MCMC", mcmc_param = list(nrep = 500, burn = 200))
plot(out, param = "mu", trunc_plot = 2)
plot(out, param = "sigma2", trunc_plot = 2)
plot(out, param = "pi", trunc_plot = 4,
      show_convergence = FALSE, show_density = FALSE)
```

plot.SANmcmc_postdens *Plot Posterior Density Samples*

Description

Visualizes the output of [compute_postdens](#). The plot displays all posterior density curves obtained from the selected MCMC iterations, together with pointwise posterior summaries and the observed data values.

Specifically, the plot includes:

- all sampled density curves (gray lines),
- the posterior median density (black line),
- the posterior mean density (red line),
- pointwise credible interval bounds (blue lines),
- observed data values shown as rug-like points on the horizontal axis.

Usage

```
## S3 method for class 'SANmcmc_postdens'  
plot(x, alpha = 0.025, ...)
```

Arguments

x	An object of class SANmcmc_postdens returned by compute_postdens .
alpha	Significance level used to construct pointwise credible intervals. The default alpha = 0.025 corresponds to a 95% pointwise credible interval.
...	Additional graphical arguments passed to plotting methods.

Value

Invisibly returns NULL. The function is called for its side effect of producing a plot.

See Also

[compute_postdens](#)

plot.SANvi

Visual Check of the Convergence of the VI Output

Description

Plot method for objects of class `SANvi`. The function displays two graphs. The left plot shows the progression of all the ELBO values as a function of the iterations. The right plots shows the ELBO increments between successive iterations of the best run on a log scale (note: increments should always be positive).

Usage

```
## S3 method for class 'SANvi'
plot(x, ...)
```

Arguments

`x` Object of class `SANvi` (usually, the result of a call to `fit_CAM`, `fit_fiSAN`, or `fit_fSAN`, used with the `est_method = "VI"` argument).

`...` Ignored.

Value

The function plots the path followed by the ELBO and its subsequent differences.

Examples

```
set.seed(123)
y <- c(rnorm(200,0,0.3), rnorm(100,5,0.3))
g <- c(rep(1,150), rep(2, 150))
out <- fit_fSAN(y = y, group = g, "VI", vi_param = list(n_runs = 2))
plot(out)
```

plot_vi_allocation_prob

Plot Variational Cluster Allocation Probabilities

Description

Produces visualizations of the posterior cluster allocation probabilities from a SAN model fitted via variational inference. The function supports plotting either the observation-level (OC) or distribution-level (DC) allocation probabilities, depending on the argument `distributional`.

This function applies to objects returned by `fit_CAM`, `fit_fiSAN`, or `fit_fSAN` when used with `est_method = "VI"`.

Usage

```
plot_vi_allocation_prob(object, distributional = FALSE, ...)
```

Arguments

object An object of class `SANvi`, representing a model fitted via variational inference.

distributional Logical (default `FALSE`). If `FALSE`, plots the allocation probabilities of individual observations to observational clusters (OC). If `TRUE`, plots the allocation probabilities of groups to distributional clusters (DC).

... Additional graphical parameters passed to the underlying `image()` function (or equivalent), for customizing the plot (e.g., `col`, `main`, `xlab`, `ylab`).

Value

The function plots the variational cluster allocation probabilities.

Examples

```
# Generate example data
set.seed(123)
y <- c(rnorm(60), rnorm(40, 5))
g <- rep(1:2, each = 50)

# Fit fiSAN via VI
est <- fit_fiSAN(y, g, est_method = "VI")

# Plot observational cluster probabilities
plot_vi_allocation_prob(est)

# Plot distributional cluster probabilities
plot_vi_allocation_prob(est, distributional = TRUE)
```

```
print.SANmcmc          Print the MCMC Output
```

Description

Print method for objects of class `SANmcmc`.

Usage

```
## S3 method for class 'SANmcmc'
print(x, ...)
```

Arguments

x Object of class `SANmcmc`.

... Ignored.

Value

The function prints a summary of the fitted model.

print.SANvi	<i>Print the Variational Inference Output</i>
-------------	---

Description

Print method for objects of class SANvi.

Usage

```
## S3 method for class 'SANvi'
print(x, ...)
```

Arguments

x	Object of class SANvi.
...	Further arguments passed to or from other methods.

Value

The function prints a summary of the fitted model.

summary.SANmcmc	<i>Summarize the MCMC Output</i>
-----------------	----------------------------------

Description

Summary method for objects of class SANmcmc.

Usage

```
## S3 method for class 'SANmcmc'
summary(object, ...)
```

Arguments

object	of class SANmcmc.
...	Ignored.

Value

The function prints a summary of the fitted model.

summary.SANvi	<i>Summarize the Variational Inference Output</i>
---------------	---

Description

Summary method for objects of class SANvi.

Usage

```
## S3 method for class 'SANvi'  
summary(object, ...)
```

Arguments

object	Object of class SANvi.
...	Further arguments passed to or from other methods.

Value

The function prints a summary of the fitted model.

Index

compute_postdens, [2](#), [24](#)
compute_psm, [4](#)

estimate_G, [5](#)
estimate_partition, [6](#), [7](#)

fit_CAM, [7](#), [8](#), [20](#), [21](#), [25](#)
fit_fiSAN, [7](#), [12](#), [20](#), [21](#), [25](#)
fit_fSAN, [7](#), [16](#), [20](#), [21](#), [25](#)

get_model, [19](#)
get_params (get_model), [19](#)
get_seed_best_run (get_model), [19](#)
get_sim (get_model), [19](#)
get_time (get_model), [19](#)

number_clusters, [21](#)

plot.partition_mcmc
 (estimate_partition), [6](#)
plot.partition_vi (estimate_partition),
 [6](#)
plot.SANmcmc, [22](#)
plot.SANmcmc_postdens, [2](#), [24](#)
plot.SANvi, [25](#)
plot.SANvi_G (estimate_G), [5](#)
plot_vi_allocation_prob, [25](#)
print.partition_mcmc
 (estimate_partition), [6](#)
print.partition_vi
 (estimate_partition), [6](#)
print.SANmcmc, [26](#)
print.SANvi, [27](#)
print.SANvi_G (estimate_G), [5](#)

salso::salso(), [6](#), [8](#)
summary.partition_mcmc
 (estimate_partition), [6](#)
summary.partition_vi
 (estimate_partition), [6](#)
summary.SANmcmc, [27](#)
summary.SANvi, [28](#)
summary.SANvi_G (estimate_G), [5](#)