# Package 'COMIX'

## November 23, 2022

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## Description

The function computes (and by default plots) estimates of the autocovariance or autocorrelation function for the different parameters of the model. This is a wrapper for coda::acf.

## Usage

```
acfParams(
   res,
   params = c("w", "xi", "xi0", "psi", "G", "E", "eta"),
   only_non_trivial_clusters = TRUE,
   lag.max = NULL,
   type = c("correlation", "covariance", "partial"),
   plot = TRUE,
   ...
)
```

## Arguments

res	An object of class COMIX or tidyChainCOMIX.						
params	A character vector naming the parameters to compute and plot the autocorrel tion plots for.						
only_non_trivial_clusters							
	Logical, if TRUE only compute and/or plot the autocorrelation for the clusters that are estimated to be non-empty.						
lag.max	maximum lag at which to calculate the autocorrelation. See more details at ?acf.						
type	Character string giving the type of autocorrelation to be computed. See more details at ?acf.						
plot	Logical. If TRUE (the default) the autocorrelation is plotted.						
	Other arguments passed to acf.						

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#### Value

An acfParamsCOMIX object which is a named list, with a named element for each requested parameter. Each element is an object of class acf (from the coda package). #' @examples library(COMIX) # Number of observations for each sample (row) and cluster (column): njk <- matrix( c( 150, 300, 250, 200 ), nrow = 2, byrow = TRUE )

- # Dimension of data: p <- 3
- # Scale and skew parameters for first cluster: Sigma1 <- matrix(0.5, nrow = p, ncol = p) + diag(0.5, nrow = p) alpha1 <- rep(0, p) alpha1[1] <- -5 # location parameter for first cluster in first sample: xi11 <- rep(0, p) # location parameter for first cluster in second sample (aligned with first): xi21 <- rep(0, p)
- # Scale and skew parameters for second cluster: Sigma2 <- matrix(-1/3, nrow = p, ncol = p) + diag(1 + 1/3, nrow = p) alpha2 <- rep(0, p) alpha2[2] <- 5 # location parameter for second cluster in first sample: xi12 <- rep(3, p) # location parameter for second cluster in second sample (misaligned with first): xi22 <- rep(4, p)
- # Sample data: set.seed(1) Y <- rbind( sn::rmsn(njk[1, 1], xi = xi11, Omega = Sigma1, alpha = alpha1), sn::rmsn(njk[1, 2], xi = xi12, Omega = Sigma2, alpha = alpha2), sn::rmsn(njk[2, 1], xi = xi21, Omega = Sigma1, alpha = alpha1), sn::rmsn(njk[2, 2], xi = xi22, Omega = Sigma2, alpha = alpha2))
- $C \leftarrow c(rep(1, rowSums(njk)[1]), rep(2, rowSums(njk)[2]))$
- prior <- list(zeta = 1, K = 10) pmc <- list(naprt = 5, nburn = 200, nsave = 200) # Reasonable usage pmc <- list(naprt = 5, nburn = 2, nsave = 5) # Minimal usage for documentation # Fit the model: res <- comix(Y, C, pmc = pmc, prior = prior)
- # Relabel to resolve potential label switching issues: res\_relab <- relabelChain(res) effssz <- effectiveSampleSize(res\_relab, "w") # Or: tidy\_chain <- tidyChain(res\_relab, "w") acf\_w <- acf-Params(tidy\_chain, "w")
- # (see vignette for a more detailed example)

calibrate	This function aligns multiple samples so that their location parameters
	are equal.

#### **Description**

This function aligns multiple samples so that their location parameters are equal.

#### **Usage**

```
calibrate(x, reference.group = NULL)
```

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#### **Arguments**

x An object of class COMIX.

reference.group

An integer between 1 and the number of groups in the data (length(unique(C))). Defaults to NULL. If NULL, the samples are aligned so that their location parameters are set to be at the estimated group location parameter. If an integer, the samples are aligned so that their location parameters are the same as the location parameter of sample reference.group.

#### Value

A named list of 3:

- Y\_cal: a nrow(x\$data\$Y) × ncol(x\$data\$Y) matrix, a calibrated version of the original data.
- calibration\_distribution: an x\$pmc\$nsave × ncol(x\$data\$Y) × nrow(x\$data\$Y) array storing the difference between the estimated sample-specific location parameter and the group location parameter for each saved step of the chain.
- calibration\_median: a nrow(x\$data\$Y) × ncol(x\$data\$Y) matrix storing the median difference between the estimated sample-specific location parameter and the group location parameter for each saved step of the chain. This matrix is equal to the difference between the uncalibrated data (x\$data\$Y) and the calibrated data (Y\_cal).

```
library(COMIX)
# Number of observations for each sample (row) and cluster (column):
nik <-
  matrix(
    c(
      150, 300,
      250, 200
    ),
    nrow = 2,
    byrow = TRUE
# Dimension of data:
# Scale and skew parameters for first cluster:
Sigma1 <- matrix(0.5, nrow = p, ncol = p) + diag(0.5, nrow = p)
alpha1 <- rep(0, p)
alpha1[1] <- -5
# location parameter for first cluster in first sample:
xi11 < - rep(0, p)
# location parameter for first cluster in second sample (aligned with first):
xi21 \leftarrow rep(0, p)
# Scale and skew parameters for second cluster:
```

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```
Sigma2 <- matrix(-1/3, nrow = p, ncol = p) + diag(1 + 1/3, nrow = p)
alpha2 <- rep(0, p)
alpha2[2] <- 5
# location parameter for second cluster in first sample:
xi12 < - rep(3, p)
# location parameter for second cluster in second sample (misaligned with first):
xi22 < - rep(4, p)
# Sample data:
set.seed(1)
Y <-
  rbind(
    sn::rmsn(njk[1, 1], xi = xi11, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[1, 2], xi = xi12, Omega = Sigma2, alpha = alpha2),
    sn::rmsn(njk[2, 1], xi = xi21, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[2, 2], xi = xi22, Omega = Sigma2, alpha = alpha2)
C \leftarrow c(rep(1, rowSums(njk)[1]), rep(2, rowSums(njk)[2]))
prior <- list(zeta = 1, K = 10)</pre>
pmc <- list(naprt = 5, nburn = 200, nsave = 200) # Reasonable usage</pre>
pmc <- list(naprt = 5, nburn = 2, nsave = 5) # Minimal usage for documentation</pre>
# Fit the model:
res <- comix(Y, C, pmc = pmc, prior = prior)</pre>
# Relabel to resolve potential label switching issues:
res_relab <- relabelChain(res)</pre>
# Generate calibrated data:
cal <- calibrateNoDist(res_relab)</pre>
# Compare raw and calibrated data: (see plot in vignette)
# par(mfrow=c(1, 2))
\# plot(Y, col = C, xlim = range(Y[,1]), ylim = range(Y[,2]))
# Get posterior estimates for the model parameters:
res_summary <- summarizeChain(res_relab)</pre>
# Check for instance, the cluster assignment labels:
table(res_summary$t)
# Indeed the same as
colSums(njk)
# Or examine the skewness parameter for the non-trivial clusters:
res_summary$alpha[ , unique(res_summary$t)]
# And compare those to
cbind(alpha1, alpha2)
# (see vignette for a more detailed example)
```

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calibrateNoDist

This function aligns multiple samples so that their location parameters are equal.

#### **Description**

This function aligns multiple samples so that their location parameters are equal.

#### Usage

```
calibrateNoDist(x, reference.group = NULL)
```

## **Arguments**

x An object of class COMIX.

reference.group

An integer between 1 and the number of groups in the data (length(unique(C))). Defaults to NULL. If NULL, the samples are aligned so that their location parameters are set to be at the estimated group location parameter. If an integer, the samples are aligned so that their location parameters are the same as the location parameter of sample reference.group.

#### Value

A named list of 2:

- Y\_cal: a nrow(x\$data\$Y) × ncol(x\$data\$Y) matrix, a calibrated version of the original data.
- calibration\_median: a nrow(x\$data\$Y) × ncol(x\$data\$Y) matrix storing the median difference between the estimated sample-specific location parameter and the group location parameter for each saved step of the chain. This matrix is equal to the difference between the uncalibrated data (x\$data\$Y) and the calibrated data (Y\_cal).

```
library(COMIX)
# Number of observations for each sample (row) and cluster (column):
njk <-
    matrix(
        c(
            150, 300,
            250, 200
        ),
        nrow = 2,
        byrow = TRUE
    )
# Dimension of data:
p <- 3</pre>
```

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```
# Scale and skew parameters for first cluster:
Sigma1 <- matrix(0.5, nrow = p, ncol = p) + diag(0.5, nrow = p)
alpha1 <- rep(0, p)
alpha1[1] <- -5
# location parameter for first cluster in first sample:
xi11 < - rep(0, p)
# location parameter for first cluster in second sample (aligned with first):
xi21 < - rep(0, p)
# Scale and skew parameters for second cluster:
Sigma2 \leftarrow matrix(-1/3, nrow = p, ncol = p) + diag(1 + 1/3, nrow = p)
alpha2 <- rep(0, p)
alpha2[2] <- 5
# location parameter for second cluster in first sample:
xi12 < - rep(3, p)
# location parameter for second cluster in second sample (misaligned with first):
xi22 < - rep(4, p)
# Sample data:
set.seed(1)
Y <-
  rbind(
    sn::rmsn(njk[1, 1], xi = xi11, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[1, 2], xi = xi12, Omega = Sigma2, alpha = alpha2),
    sn::rmsn(njk[2, 1], xi = xi21, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[2, 2], xi = xi22, Omega = Sigma2, alpha = alpha2)
C \leftarrow c(rep(1, rowSums(njk)[1]), rep(2, rowSums(njk)[2]))
prior \leftarrow list(zeta = 1, K = 10)
pmc <- list(naprt = 5, nburn = 200, nsave = 200) # Reasonable usage</pre>
pmc <- list(naprt = 5, nburn = 2, nsave = 5) # Minimal usage for documentation
# Fit the model:
res <- comix(Y, C, pmc = pmc, prior = prior)</pre>
# Relabel to resolve potential label switching issues:
res_relab <- relabelChain(res)</pre>
# Generate calibrated data:
cal <- calibrateNoDist(res_relab)</pre>
# Compare raw and calibrated data: (see plot in vignette)
# par(mfrow=c(1, 2))
\# plot(Y, col = C, xlim = range(Y[,1]), ylim = range(Y[,2]) )
# Get posterior estimates for the model parameters:
res_summary <- summarizeChain(res_relab)</pre>
# Check for instance, the cluster assignment labels:
table(res_summary$t)
# Indeed the same as
colSums(njk)
```

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```
# Or examine the skewness parameter for the non-trivial clusters:
res_summary$alpha[ , unique(res_summary$t)]
# And compare those to
cbind(alpha1, alpha2)
# (see vignette for a more detailed example)
```

comix

This function generates a sample from the posterior of COMIX.

#### Description

This function generates a sample from the posterior of COMIX.

#### Usage

```
comix(Y, C, prior = NULL, pmc = NULL, state = NULL, ncores = 2)
```

## **Arguments**

Y Matrix of the data. Each row represents an observation.

C Vector of the group label of each observation. Labels must be integers starting from 1.

prior

A list giving the prior information. If unspecified, a default prior is used. The list includes the following parameters:

- zeta: Coarsening parameter. A number between 0 and 1. zeta = 1: sample from standard posterior; zeta < 1: sample from power posterior. The lower zeta is, the more flexible the kernels become.
- K: Maximal number of mixture components.
- eta\_prior Parameters for gamma prior for concentration parameter of the stick breaking process prior for the weights.
- m0: Number of degrees of freedom for the inverse Wishart prior for Sigma, the covariance matrix of the kernels.
- Lambda: Mean parameter for the inverse Wishart prior for Sigma, the covariance matrix of the kernels.
- b0: Mean parameter for the multivariate normal distribution that is the prior for the group mean parameter xi0.
- B0: Covariance parameter for the multivariate normal distribution that is the prior for the group mean parameter xi0.
- e0: Number of degrees of freedom for the inverse Wishart prior for  $E_k$ , the covariance matrix of the multivariate normal from which  $\xi_{j,k}$  are drawn.
- E0: Mean parameter for the inverse Wishart prior for  $E_k$ , the covariance matrix of the multivariate normal from which  $\xi_{j,k}$  are drawn.
- merge\_step: Introduce step to merge mixture components with small KL divergence. Default is merge\_step = TRUE.

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merge\_par: Parameter controlling merging radius. Default is merge\_par = 0.1.

pmc

A list giving the Population Monte Carlo (PMC) parameters:

- npart: Number of PMC particles.
- nburn: Number of burn-in steps
- nsave: Number of steps in the chain after burn-in.
- nskip: Thinning parameter, number of steps to skip between saving steps after burn-in.
- ndisplay: Display status of chain after every ndisplay steps.

state

A list giving the initial cluster labels:

 t: An integer vector, same length as the number of rows of Y, with cluster labels between 1 and K.

ncores

The number of CPU cores to utilize in parallel. Defaults to 2.

#### Value

An object of class COMIX, a list of 4:

chain, a named list:

- t: an nsave × nrow(Y) matrix with estimated cluster labels for each saved step of the chain and each observation in the data Y.
- z: a nsave  $\times$  nrow(Y) matrix with estimated values of the latent  $z_{i,j}$  variable for the parameterization of the multivariate skew normal distribution used in the sampler for each saved step of the chain and each observation in the data Y.
- W: an length(unique(C))  $\times$  K  $\times$
- nsave: array storing the estimated sample- and cluster-specific weights for each saved step of the chain.
- xi: an length(unique(C)) × (ncol(Y) x K) × nsave array storing the estimated sampleand cluster-specific multivariate skew normal location parameters of the kernel for each saved step of the chain.
- xi0: an  $ncol(Y) \times K \times$
- nsave: array storing the estimated cluster-specific group location parameters for each saved step of the chain.
- psi: an ncol(Y) × K × nsave array storing the estimated cluster-specific skew parameters of the kernels in the parameterization of the multivariate skew normal distribution used in the sampler for each saved step of the chain.
- G: an ncol(Y) × (ncol(Y) x K) × nsave array storing the estimated cluster-specific multivariate skew normal scale matrix (in row format) of the kernel used in the sampler for each saved step of the chain.
- E: an ncol(Y) × (ncol(Y) x K) × nsave array storing the estimated covariance matrix (in row format) of the multivariate normal distribution from which the sample- and cluster-specific location parameters are drawn for each saved step of the chain.
- eta: a nsave × 1 matrix storing the estimated Dirichlet Process Mixture concentration parameter for each saved step of the chain.

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Sigma: an ncol(Y) × (ncol(Y) x K) × nsave array storing the estimated cluster-specific
multivariate skew normal scale matrix (in row format) of the kernel for each saved step
of the chain.

- alpha: an ncol(Y) × K × nsave array storing the estimated cluster-specific skew parameters of the kernel's multivariate skew normal distribution for each saved step of the chain.
- data, a named list that includes the matrix of the data Y and C the vector of the group label of each observation.
- prior and pmc, the lists, as above, that were provided as inputs to the function.

```
library(COMIX)
# Number of observations for each sample (row) and cluster (column):
  matrix(
   c(
      150, 300,
      250, 200
   ),
   nrow = 2,
    byrow = TRUE
# Dimension of data:
p < -3
# Scale and skew parameters for first cluster:
Sigma1 <- matrix(0.5, nrow = p, ncol = p) + diag(0.5, nrow = p)
alpha1 <- rep(0, p)
alpha1[1] <- -5
# location parameter for first cluster in first sample:
xi11 < - rep(0, p)
# location parameter for first cluster in second sample (aligned with first):
xi21 < - rep(0, p)
# Scale and skew parameters for second cluster:
Sigma2 \leftarrow matrix(-1/3, nrow = p, ncol = p) + diag(1 + 1/3, nrow = p)
alpha2 < - rep(0, p)
alpha2[2] <- 5
# location parameter for second cluster in first sample:
xi12 < - rep(3, p)
# location parameter for second cluster in second sample (misaligned with first):
xi22 < - rep(4, p)
# Sample data:
set.seed(1)
Y <-
  rbind(
    sn::rmsn(njk[1, 1], xi = xi11, Omega = Sigma1, alpha = alpha1),
```

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```
sn::rmsn(njk[1, 2], xi = xi12, Omega = Sigma2, alpha = alpha2),
    sn::rmsn(njk[2, 1], xi = xi21, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[2, 2], xi = xi22, Omega = Sigma2, alpha = alpha2)
  )
C \leftarrow c(rep(1, rowSums(njk)[1]), rep(2, rowSums(njk)[2]))
prior <- list(zeta = 1, K = 10)</pre>
pmc <- list(naprt = 5, nburn = 200, nsave = 200) # Reasonable usage
pmc <- list(naprt = 5, nburn = 2, nsave = 5) # Minimal usage for documentation</pre>
# Fit the model:
res <- comix(Y, C, pmc = pmc, prior = prior)</pre>
# Relabel to resolve potential label switching issues:
res_relab <- relabelChain(res)</pre>
# Generate calibrated data:
cal <- calibrateNoDist(res_relab)</pre>
# Compare raw and calibrated data: (see plot in vignette)
# par(mfrow=c(1, 2))
\# plot(Y, col = C, xlim = range(Y[,1]), ylim = range(Y[,2]) )
# Get posterior estimates for the model parameters:
res_summary <- summarizeChain(res_relab)</pre>
# Check for instance, the cluster assignment labels:
table(res_summary$t)
# Indeed the same as
colSums(njk)
# Or examine the skewness parameter for the non-trivial clusters:
res_summary$alpha[ , unique(res_summary$t)]
# And compare those to
cbind(alpha1, alpha2)
# (see vignette for a more detailed example)
```

effectiveSampleSize

This function creates an object that summarizes the effective sample size for the parameters of the model.

#### **Description**

This function creates an object that summarizes the effective sample size for the parameters of the model.

## Usage

```
effectiveSampleSize(res, params = c("w", "xi", "xi0", "psi", "G", "E", "eta"))
```

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#### Arguments

res An object of class COMIX or tidyChainCOMIX.

params A character vector naming the parameters to compute the effective sample size

for.

#### Value

An effectiveSampleSizeCOMIX object which is a named list, with a named element for each requested parameter. Each element is a data frame that includes the effective sample size for the parameter.

```
library(COMIX)
# Number of observations for each sample (row) and cluster (column):
njk <-
  matrix(
    c(
      150, 300,
      250, 200
    ),
    nrow = 2,
    byrow = TRUE
  )
# Dimension of data:
p <- 3
# Scale and skew parameters for first cluster:
Sigma1 <- matrix(0.5, nrow = p, ncol = p) + diag(0.5, nrow = p)
alpha1 < - rep(0, p)
alpha1[1] <- -5
# location parameter for first cluster in first sample:
xi11 < - rep(0, p)
# location parameter for first cluster in second sample (aligned with first):
xi21 \leftarrow rep(0, p)
# Scale and skew parameters for second cluster:
Sigma2 \leftarrow matrix(-1/3, nrow = p, ncol = p) + diag(1 + 1/3, nrow = p)
alpha2 < - rep(0, p)
alpha2[2] <- 5
# location parameter for second cluster in first sample:
xi12 < - rep(3, p)
# location parameter for second cluster in second sample (misaligned with first):
xi22 < - rep(4, p)
# Sample data:
set.seed(1)
Y <-
  rbind(
    sn::rmsn(njk[1, 1], xi = xi11, Omega = Sigma1, alpha = alpha1),
```

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```
sn::rmsn(njk[1, 2], xi = xi12, Omega = Sigma2, alpha = alpha2),
    sn::rmsn(njk[2, 1], xi = xi21, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[2, 2], xi = xi22, Omega = Sigma2, alpha = alpha2)
  )
C \leftarrow c(rep(1, rowSums(njk)[1]), rep(2, rowSums(njk)[2]))
prior <- list(zeta = 1, K = 10)</pre>
pmc <- list(naprt = 5, nburn = 200, nsave = 200) # Reasonable usage</pre>
pmc <- list(naprt = 5, nburn = 2, nsave = 5) # Minimal usage for documentation</pre>
# Fit the model:
res <- comix(Y, C, pmc = pmc, prior = prior)</pre>
# Relabel to resolve potential label switching issues:
res_relab <- relabelChain(res)</pre>
effssz <- effectiveSampleSize(res_relab, "w")</pre>
# Or:
tidy_chain <- tidyChain(res_relab, "w")</pre>
effssz <- effectiveSampleSize(tidy_chain, "w")</pre>
# (see vignette for a more detailed example)
```

gewekeParams

This function creates an object that summarizes the Geweke convergence diagnostic.

## Description

This function creates an object that summarizes the Geweke convergence diagnostic.

#### Usage

```
gewekeParams(
   res,
   params = c("w", "xi", "xi0", "psi", "G", "E", "eta"),
   frac1 = 0.1,
   frac2 = 0.5,
   probs = c(0.025, 0.975)
)
```

## **Arguments**

res	An object of class COMIX or tidyChainCOMIX.
params	A character vector naming the parameters to compute the Geweke diagnostic for.
frac1	Double, fraction to use from beginning of chain.
frac2	Double, fraction to use from end of chain.
probs	A vector of 2 doubles, probabilities denoting the limits of a confidence interval for the convergence diagnostic.

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#### Value

An gewekeParamsCOMIX object which is a named list, with a named element for each requested parameter. Each element is a data frame that includes the Geweke diagnostic and result of a stationarity test for the parameter.

```
library(COMIX)
# Number of observations for each sample (row) and cluster (column):
nik <-
  matrix(
    c(
      150, 300,
      250, 200
    ),
    nrow = 2.
    byrow = TRUE
# Dimension of data:
p < -3
# Scale and skew parameters for first cluster:
Sigma1 <- matrix(0.5, nrow = p, ncol = p) + diag(0.5, nrow = p)
alpha1 <- rep(0, p)
alpha1[1] <- -5
# location parameter for first cluster in first sample:
xi11 < - rep(0, p)
# location parameter for first cluster in second sample (aligned with first):
xi21 \leftarrow rep(0, p)
# Scale and skew parameters for second cluster:
Sigma2 \leftarrow matrix(-1/3, nrow = p, ncol = p) + diag(1 + 1/3, nrow = p)
alpha2 < - rep(0, p)
alpha2[2] <- 5
# location parameter for second cluster in first sample:
xi12 < - rep(3, p)
# location parameter for second cluster in second sample (misaligned with first):
xi22 < - rep(4, p)
# Sample data:
set.seed(1)
Y <-
  rbind(
    sn::rmsn(njk[1, 1], xi = xi11, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[1, 2], xi = xi12, Omega = Sigma2, alpha = alpha2),
    sn::rmsn(njk[2, 1], xi = xi21, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[2, 2], xi = xi22, Omega = Sigma2, alpha = alpha2)
C \leftarrow c(rep(1, rowSums(njk)[1]), rep(2, rowSums(njk)[2]))
```

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```
prior <- list(zeta = 1, K = 10)
pmc <- list(naprt = 5, nburn = 200, nsave = 200) # Reasonable usage
pmc <- list(naprt = 5, nburn = 2, nsave = 5) # Minimal usage for documentation
# Fit the model:
res <- comix(Y, C, pmc = pmc, prior = prior)

# Relabel to resolve potential label switching issues:
res_relab <- relabelChain(res)
effssz <- effectiveSampleSize(res_relab, "w")
# Or:
tidy_chain <- tidyChain(res_relab, "w")
gwk <- gewekeParams(tidy_chain, "w")
# (see vignette for a more detailed example)</pre>
```

heidelParams

This function creates an object that summarizes the Heidelberg-Welch convergence diagnostic.

## Description

This function creates an object that summarizes the Heidelberg-Welch convergence diagnostic.

## Usage

```
heidelParams(
    res,
    params = c("w", "xi", "xi0", "psi", "G", "E", "eta"),
    eps = 0.1,
    pvalue = 0.05
)
```

#### **Arguments**

res An object of class COMIX or tidyChainCOMIX.

params A character vector naming the parameters to compute the Heidelberg-Welch

diagnostic for.

eps Target value for ratio of halfwidth to sample mean.

pvalue Significance level to use.

#### Value

An heidelParamsCOMIX object which is a named list, with a named element for each requested parameter. Each element is a data frame that includes the Heidelberg-Welch diagnostic and results of a stationarity test for the parameter.

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```
library(COMIX)
# Number of observations for each sample (row) and cluster (column):
nik <-
  matrix(
    c(
      150, 300,
      250, 200
    ),
    nrow = 2,
    byrow = TRUE
  )
# Dimension of data:
p < -3
# Scale and skew parameters for first cluster:
Sigma1 <- matrix(0.5, nrow = p, ncol = p) + diag(0.5, nrow = p)
alpha1 < - rep(0, p)
alpha1[1] <- -5
# location parameter for first cluster in first sample:
xi11 < - rep(0, p)
# location parameter for first cluster in second sample (aligned with first):
xi21 \leftarrow rep(0, p)
# Scale and skew parameters for second cluster:
Sigma2 \leftarrow matrix(-1/3, nrow = p, ncol = p) + diag(1 + 1/3, nrow = p)
alpha2 <- rep(0, p)
alpha2Γ21 <- 5
# location parameter for second cluster in first sample:
xi12 < - rep(3, p)
# location parameter for second cluster in second sample (misaligned with first):
xi22 < - rep(4, p)
# Sample data:
set.seed(1)
Y <-
  rbind(
    sn::rmsn(njk[1, 1], xi = xi11, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[1, 2], xi = xi12, Omega = Sigma2, alpha = alpha2),
    sn::rmsn(njk[2, 1], xi = xi21, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[2, 2], xi = xi22, Omega = Sigma2, alpha = alpha2)
C <- c(rep(1, rowSums(njk)[1]), rep(2, rowSums(njk)[2]))</pre>
prior \leftarrow list(zeta = 1, K = 10)
pmc <- list(naprt = 5, nburn = 200, nsave = 200) # Reasonable usage
pmc <- list(naprt = 5, nburn = 2, nsave = 5) # Minimal usage for documentation
# Fit the model:
res <- comix(Y, C, pmc = pmc, prior = prior)</pre>
```

plotEffective Sample Size

```
# Relabel to resolve potential label switching issues:
res_relab <- relabelChain(res)
effssz <- effectiveSampleSize(res_relab, "w")
# Or:
tidy_chain <- tidyChain(res_relab, "w")
hd <- heidelParams(tidy_chain, "w")
# (see vignette for a more detailed example)</pre>
```

plotEffectiveSampleSize

This function creates plots for the effective sample size for the parameters of the model.

## Description

This function creates plots for the effective sample size for the parameters of the model.

#### Usage

```
plotEffectiveSampleSize(effssz, param)
```

## **Arguments**

effssz An object of class effectiveSampleSizeCOMIX as created by the function effectiveSampleSize.

param Character, naming the parameter to create a plot of effective sample sizes.

#### Value

A ggplot2 plot containing the effective sample size plot.

```
library(COMIX)
# Number of observations for each sample (row) and cluster (column):
njk <-
    matrix(
        c(
            150, 300,
            250, 200
        ),
        nrow = 2,
        byrow = TRUE
    )

# Dimension of data:
p <- 3
# Scale and skew parameters for first cluster:
Sigma1 <- matrix(0.5, nrow = p, ncol = p) + diag(0.5, nrow = p)</pre>
```

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```
alpha1 < - rep(0, p)
alpha1[1] <- -5
# location parameter for first cluster in first sample:
xi11 < - rep(0, p)
# location parameter for first cluster in second sample (aligned with first):
xi21 \leftarrow rep(0, p)
# Scale and skew parameters for second cluster:
Sigma2 \leftarrow matrix(-1/3, nrow = p, ncol = p) + diag(1 + 1/3, nrow = p)
alpha2 <- rep(0, p)
alpha2[2] <- 5
# location parameter for second cluster in first sample:
xi12 < - rep(3, p)
# location parameter for second cluster in second sample (misaligned with first):
xi22 < - rep(4, p)
# Sample data:
set.seed(1)
Υ <-
  rbind(
    sn::rmsn(njk[1, 1], xi = xi11, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[1, 2], xi = xi12, Omega = Sigma2, alpha = alpha2),
    sn::rmsn(njk[2, 1], xi = xi21, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[2, 2], xi = xi22, Omega = Sigma2, alpha = alpha2)
C \leftarrow c(rep(1, rowSums(njk)[1]), rep(2, rowSums(njk)[2]))
prior <- list(zeta = 1, K = 10)</pre>
pmc <- list(naprt = 5, nburn = 200, nsave = 200) # Reasonable usage</pre>
pmc <- list(naprt = 5, nburn = 2, nsave = 5) # Minimal usage for documentation</pre>
# Fit the model:
res <- comix(Y, C, pmc = pmc, prior = prior)</pre>
# Relabel to resolve potential label switching issues:
res_relab <- relabelChain(res)</pre>
effssz <- effectiveSampleSize(res_relab, "w")</pre>
# Or:
tidy_chain <- tidyChain(res_relab, "w")</pre>
effssz <- effectiveSampleSize(tidy_chain, "w")</pre>
plotEffectiveSampleSize(effssz, "w")
# (see vignette for a more detailed example)
```

plotGewekeParams

This function creates plots for the Geweke diagnostic and results of test of stationarity for the parameters of the model.

#### **Description**

This function creates plots for the Geweke diagnostic and results of test of stationarity for the parameters of the model.

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#### Usage

```
plotGewekeParams(gwk, param)
```

#### **Arguments**

gwk An object of class gewekeParamsCOMIX as created by the function gewekeParams.

Param Character, naming the parameter to create a plot of the Geweke diagnostic for.

#### Value

A ggplot2 plot containing the Geweke diagnostic plot.

```
library(COMIX)
# Number of observations for each sample (row) and cluster (column):
  matrix(
    c(
      150, 300,
      250, 200
    ),
    nrow = 2,
    byrow = TRUE
# Dimension of data:
p < -3
# Scale and skew parameters for first cluster:
Sigma1 <- matrix(0.5, nrow = p, ncol = p) + diag(0.5, nrow = p)
alpha1 < - rep(0, p)
alpha1[1] <- -5
# location parameter for first cluster in first sample:
xi11 < - rep(0, p)
# location parameter for first cluster in second sample (aligned with first):
xi21 \leftarrow rep(0, p)
# Scale and skew parameters for second cluster:
Sigma2 <- matrix(-1/3, nrow = p, ncol = p) + diag(1 + 1/3, nrow = p)
alpha2 < - rep(0, p)
alpha2[2] <- 5
# location parameter for second cluster in first sample:
xi12 < - rep(3, p)
# location parameter for second cluster in second sample (misaligned with first):
xi22 \leftarrow rep(4, p)
# Sample data:
set.seed(1)
Y <-
  rbind(
```

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```
sn::rmsn(njk[1, 1], xi = xi11, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[1, 2], xi = xi12, Omega = Sigma2, alpha = alpha2),
    sn::rmsn(njk[2, 1], xi = xi21, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[2, 2], xi = xi22, Omega = Sigma2, alpha = alpha2)
 )
C \leftarrow c(rep(1, rowSums(njk)[1]), rep(2, rowSums(njk)[2]))
prior <- list(zeta = 1, K = 10)</pre>
pmc <- list(naprt = 5, nburn = 200, nsave = 200) # Reasonable usage</pre>
pmc <- list(naprt = 5, nburn = 2, nsave = 5) # Minimal usage for documentation</pre>
# Fit the model:
res <- comix(Y, C, pmc = pmc, prior = prior)
# Relabel to resolve potential label switching issues:
res_relab <- relabelChain(res)</pre>
effssz <- effectiveSampleSize(res_relab, "w")</pre>
tidy_chain <- tidyChain(res_relab, "w")</pre>
gwk <- gewekeParams(tidy_chain, "w")</pre>
plotGewekeParams(gwk, "w")
# (see vignette for a more detailed example)
```

plotHeidelParams

This function creates plots for the Heidelberg-Welch diagnostic and results of test of stationarity for the parameters of the model.

#### Description

This function creates plots for the Heidelberg-Welch diagnostic and results of test of stationarity for the parameters of the model.

#### Usage

```
plotHeidelParams(hd, param)
```

#### **Arguments**

hd An object of class heidelParamsCOMIX as created by the function heidelParams.

param Character, naming the parameter to create a plot of the Heidelberg-Welch diag-

nostic for.

#### Value

A ggplot2 plot containing the Heidelberg-Welch diagnostic plot.

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```
library(COMIX)
# Number of observations for each sample (row) and cluster (column):
nik <-
  matrix(
    c(
      150, 300,
      250, 200
    ),
    nrow = 2,
    byrow = TRUE
  )
# Dimension of data:
p < -3
# Scale and skew parameters for first cluster:
Sigma1 <- matrix(0.5, nrow = p, ncol = p) + diag(0.5, nrow = p)
alpha1 < - rep(0, p)
alpha1[1] <- -5
# location parameter for first cluster in first sample:
xi11 < - rep(0, p)
# location parameter for first cluster in second sample (aligned with first):
xi21 \leftarrow rep(0, p)
# Scale and skew parameters for second cluster:
Sigma2 \leftarrow matrix(-1/3, nrow = p, ncol = p) + diag(1 + 1/3, nrow = p)
alpha2 <- rep(0, p)
alpha2Γ21 <- 5
# location parameter for second cluster in first sample:
xi12 < - rep(3, p)
# location parameter for second cluster in second sample (misaligned with first):
xi22 < - rep(4, p)
# Sample data:
set.seed(1)
Y <-
  rbind(
    sn::rmsn(njk[1, 1], xi = xi11, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[1, 2], xi = xi12, Omega = Sigma2, alpha = alpha2),
    sn::rmsn(njk[2, 1], xi = xi21, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[2, 2], xi = xi22, Omega = Sigma2, alpha = alpha2)
C <- c(rep(1, rowSums(njk)[1]), rep(2, rowSums(njk)[2]))</pre>
prior \leftarrow list(zeta = 1, K = 10)
pmc <- list(naprt = 5, nburn = 200, nsave = 200) # Reasonable usage
pmc <- list(naprt = 5, nburn = 2, nsave = 5) # Minimal usage for documentation
# Fit the model:
res <- comix(Y, C, pmc = pmc, prior = prior)</pre>
```

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```
# Relabel to resolve potential label switching issues:
res_relab <- relabelChain(res)
effssz <- effectiveSampleSize(res_relab, "w")
# Or:
tidy_chain <- tidyChain(res_relab, "w")
hd <- heidelParams(tidy_chain, "w")
plotHeidelParams(hd, "w")
# (see vignette for a more detailed example)</pre>
```

plotTracePlots

This function creates trace plots for different parameters of the MCMC chain.

## Description

This function creates trace plots for different parameters of the MCMC chain.

#### Usage

```
plotTracePlots(res, param)
```

## **Arguments**

res An object of class COMIX or tidyChainCOMIX.

param Character, naming the parameter to create a trace plot for.

#### Value

A ggplot2 plot containing the trace plot.

```
library(COMIX)
# Number of observations for each sample (row) and cluster (column):
njk <-
matrix(
    c(
       150, 300,
       250, 200
    ),
    nrow = 2,
    byrow = TRUE
    )

# Dimension of data:
p <- 3
# Scale and skew parameters for first cluster:
Sigma1 <- matrix(0.5, nrow = p, ncol = p) + diag(0.5, nrow = p)</pre>
```

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```
alpha1 < - rep(0, p)
alpha1[1] < - -5
# location parameter for first cluster in first sample:
xi11 < - rep(0, p)
# location parameter for first cluster in second sample (aligned with first):
xi21 \leftarrow rep(0, p)
# Scale and skew parameters for second cluster:
Sigma2 \leftarrow matrix(-1/3, nrow = p, ncol = p) + diag(1 + 1/3, nrow = p)
alpha2 <- rep(0, p)
alpha2[2] <- 5
# location parameter for second cluster in first sample:
xi12 < - rep(3, p)
# location parameter for second cluster in second sample (misaligned with first):
xi22 < - rep(4, p)
# Sample data:
set.seed(1)
Y <-
  rbind(
    sn::rmsn(njk[1, 1], xi = xi11, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[1, 2], xi = xi12, Omega = Sigma2, alpha = alpha2),
    sn::rmsn(njk[2, 1], xi = xi21, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[2, 2], xi = xi22, Omega = Sigma2, alpha = alpha2)
C \leftarrow c(rep(1, rowSums(njk)[1]), rep(2, rowSums(njk)[2]))
prior <- list(zeta = 1, K = 10)</pre>
pmc <- list(naprt = 5, nburn = 200, nsave = 200) # Reasonable usage</pre>
pmc <- list(naprt = 5, nburn = 2, nsave = 5) # Minimal usage for documentation</pre>
# Fit the model:
res <- comix(Y, C, pmc = pmc, prior = prior)</pre>
# Relabel to resolve potential label switching issues:
res_relab <- relabelChain(res)</pre>
plotTracePlots(res_relab, "w")
# Or:
tidy_chain <- tidyChain(res_relab, "w")</pre>
plotTracePlots(tidy_chain, "w")
# (see vignette for a more detailed example)
```

relabelChain

This function relabels the chain to avoid label switching issues.

#### **Description**

This function relabels the chain to avoid label switching issues.

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#### Usage

```
relabelChain(res)
```

#### **Arguments**

res

An object of class COMIX.

## Value

An object of class COMIX where res\$chain\$t is replaced with the new labels.

```
library(COMIX)
# Number of observations for each sample (row) and cluster (column):
nik <-
 matrix(
   c(
      150, 300,
      250, 200
   ),
   nrow = 2,
    byrow = TRUE
# Dimension of data:
p <- 3
# Scale and skew parameters for first cluster:
Sigma1 <- matrix(0.5, nrow = p, ncol = p) + diag(0.5, nrow = p)
alpha1 <- rep(0, p)
alpha1[1] <- -5
# location parameter for first cluster in first sample:
xi11 < - rep(0, p)
# location parameter for first cluster in second sample (aligned with first):
xi21 \leftarrow rep(0, p)
# Scale and skew parameters for second cluster:
Sigma2 \leftarrow matrix(-1/3, nrow = p, ncol = p) + diag(1 + 1/3, nrow = p)
alpha2 < - rep(0, p)
alpha2[2] <- 5
# location parameter for second cluster in first sample:
xi12 < - rep(3, p)
# location parameter for second cluster in second sample (misaligned with first):
xi22 < - rep(4, p)
# Sample data:
set.seed(1)
Y <-
  rbind(
    sn::rmsn(njk[1, 1], xi = xi11, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[1, 2], xi = xi12, Omega = Sigma2, alpha = alpha2),
```

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```
sn::rmsn(njk[2, 1], xi = xi21, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[2, 2], xi = xi22, Omega = Sigma2, alpha = alpha2)
C \leftarrow c(rep(1, rowSums(njk)[1]), rep(2, rowSums(njk)[2]))
prior \leftarrow list(zeta = 1, K = 10)
pmc <- list(naprt = 5, nburn = 200, nsave = 200) # Reasonable usage
pmc <- list(naprt = 5, nburn = 2, nsave = 5) # Minimal usage for documentation</pre>
# Fit the model:
res <- comix(Y, C, pmc = pmc, prior = prior)</pre>
# Relabel to resolve potential label switching issues:
res_relab <- relabelChain(res)</pre>
# Generate calibrated data:
cal <- calibrateNoDist(res_relab)</pre>
# Compare raw and calibrated data: (see plot in vignette)
# par(mfrow=c(1, 2))
\# plot(Y, col = C, xlim = range(Y[,1]), ylim = range(Y[,2]) )
# Get posterior estimates for the model parameters:
res_summary <- summarizeChain(res_relab)</pre>
# Check for instance, the cluster assignment labels:
table(res_summary$t)
# Indeed the same as
colSums(njk)
# Or examine the skewness parameter for the non-trivial clusters:
res_summary$alpha[ , unique(res_summary$t)]
# And compare those to
cbind(alpha1, alpha2)
# (see vignette for a more detailed example)
```

summarizeChain

This function provides post-hoc estimates of the model parameters.

#### Description

This function provides post-hoc estimates of the model parameters.

#### Usage

```
summarizeChain(res)
```

#### **Arguments**

res

An object of class COMIX.

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#### Value

#### A named list:

 xi0: a ncol(res\$data\$Y) × res\$prior\$K matrix storing the posterior mean of the group location parameter.

- psi: a ncol(res\$data\$Y) × res\$prior\$K matrix storing the posterior mean of the multivariate skew normal kernels skewness parameter (in the parameterization used in the sampler).
- alpha: a ncol(res\$data\$Y) × res\$prior\$K matrix storing the posterior mean of the multivariate skew normal kernels skewness parameter.
- W: a length(unique(res\$data\$C)) × res\$prior\$K matrix storing the posterior mean of the mixture weights for each sample and cluster.
- xi: an length(unique(res\$data\$C)) × ncol(res\$data\$Y) × res\$prior\$K array storing
  the the posterior mean of the multivariate skew normal kernels location parameter for each
  sample and cluster.
- Sigma: an ncol(res\$data\$Y) × ncol(res\$data\$Y) × res\$prior\$K array storing the the posterior mean of the scaling matrix of the multivariate skew normal kernels for each cluster.
- G: an ncol(res\$data\$Y) × ncol(res\$data\$Y) × res\$prior\$K array storing the posterior mean of the scaling matrix of the multivariate skew normal kernels for each cluster (in the parameterization used in the sampler).
- E: an ncol(res\$data\$Y) × ncol(res\$data\$Y) × res\$prior\$K array storing the posterior mean of the covariance matrix of the multivariate normal distributions for each cluster form which the sample specific location parameters are drawn.
- meanvec: an length(unique(resdataC))  $\times$  ncol(resdataY)  $\times$  respriorK array storing the posterior mean of the multivariate skew normal kernels mean parameter for each sample and cluster.
- meanvec0: a ncol(res\$data\$Y)  $\times$  res\$prior\$K matrix storing the posterior mean of the group mean parameter.
- t: Vector of length nrow(x\$data\$Y). Each element is the mode of the posterior distribution of cluster labels.
- eta: scalar, the mean of the posterior distribution of the estimated Dirichlet Process Mixture concentration parameter.

```
library(COMIX)
# Number of observations for each sample (row) and cluster (column):
njk <-
    matrix(
    c(
        150, 300,
        250, 200
    ),
    nrow = 2,
    byrow = TRUE
   )</pre>
```

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```
# Dimension of data:
p < -3
# Scale and skew parameters for first cluster:
Sigma1 <- matrix(0.5, nrow = p, ncol = p) + diag(0.5, nrow = p)
alpha1 <- rep(0, p)
alpha1[1] <- -5
# location parameter for first cluster in first sample:
xi11 < - rep(0, p)
# location parameter for first cluster in second sample (aligned with first):
xi21 \leftarrow rep(0, p)
# Scale and skew parameters for second cluster:
Sigma2 \leftarrow matrix(-1/3, nrow = p, ncol = p) + diag(1 + 1/3, nrow = p)
alpha2 <- rep(0, p)
alpha2[2] <- 5
# location parameter for second cluster in first sample:
xi12 < - rep(3, p)
# location parameter for second cluster in second sample (misaligned with first):
xi22 < - rep(4, p)
# Sample data:
set.seed(1)
Y <-
  rbind(
    sn::rmsn(njk[1, 1], xi = xi11, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[1, 2], xi = xi12, Omega = Sigma2, alpha = alpha2),
    sn::rmsn(njk[2, 1], xi = xi21, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[2, 2], xi = xi22, Omega = Sigma2, alpha = alpha2)
C \leftarrow c(rep(1, rowSums(njk)[1]), rep(2, rowSums(njk)[2]))
prior <- list(zeta = 1, K = 10)</pre>
pmc <- list(naprt = 5, nburn = 200, nsave = 200) # Reasonable usage
pmc <- list(naprt = 5, nburn = 2, nsave = 5) # Minimal usage for documentation</pre>
# Fit the model:
res <- comix(Y, C, pmc = pmc, prior = prior)</pre>
# Relabel to resolve potential label switching issues:
res_relab <- relabelChain(res)</pre>
# Generate calibrated data:
cal <- calibrateNoDist(res_relab)</pre>
# Compare raw and calibrated data: (see plot in vignette)
# par(mfrow=c(1, 2))
\# plot(Y, col = C, xlim = range(Y[,1]), ylim = range(Y[,2]))
# Get posterior estimates for the model parameters:
res_summary <- summarizeChain(res_relab)</pre>
# Check for instance, the cluster assignment labels:
table(res_summary$t)
```

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```
# Indeed the same as
colSums(njk)

# Or examine the skewness parameter for the non-trivial clusters:
res_summary$alpha[ , unique(res_summary$t)]
# And compare those to
cbind(alpha1, alpha2)

# (see vignette for a more detailed example)
```

tidyChain

This function creates tidy versions of the stored chain. This object can then be used as input for the other diagnostic functions in this package.

## **Description**

This function creates tidy versions of the stored chain. This object can then be used as input for the other diagnostic functions in this package.

#### Usage

```
tidyChain(
  res,
  params = c("t", "w", "xi", "xi0", "psi", "G", "E", "eta", "Sigma", "alpha")
)
```

### **Arguments**

res An object of class COMIX.

params A character vector naming the parameters to tidy.

#### Value

A tidyChainCOMIX object: a named list of class whose length is the length of params. Each element of the list contains a tibble with a tidy version of the samples from the MCMC chain.

```
library(COMIX)
# Number of observations for each sample (row) and cluster (column):
njk <-
    matrix(
        c(
            150, 300,
            250, 200
        ),
        nrow = 2,
        byrow = TRUE
        )</pre>
```

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```
# Dimension of data:
p <- 3
# Scale and skew parameters for first cluster:
Sigma1 <- matrix(0.5, nrow = p, ncol = p) + diag(0.5, nrow = p)
alpha1 <- rep(0, p)
alpha1[1] <- -5
# location parameter for first cluster in first sample:
xi11 < - rep(0, p)
# location parameter for first cluster in second sample (aligned with first):
xi21 \leftarrow rep(0, p)
# Scale and skew parameters for second cluster:
Sigma2 \leftarrow matrix(-1/3, nrow = p, ncol = p) + diag(1 + 1/3, nrow = p)
alpha2 <- rep(0, p)
alpha2[2] <- 5
# location parameter for second cluster in first sample:
xi12 < - rep(3, p)
# location parameter for second cluster in second sample (misaligned with first):
xi22 < - rep(4, p)
# Sample data:
set.seed(1)
Y <-
  rbind(
    sn::rmsn(njk[1, 1], xi = xi11, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[1, 2], xi = xi12, Omega = Sigma2, alpha = alpha2),
    sn::rmsn(njk[2, 1], xi = xi21, Omega = Sigma1, alpha = alpha1),
    sn::rmsn(njk[2, 2], xi = xi22, Omega = Sigma2, alpha = alpha2)
C \leftarrow c(rep(1, rowSums(njk)[1]), rep(2, rowSums(njk)[2]))
prior <- list(zeta = 1, K = 10)</pre>
pmc <- list(naprt = 5, nburn = 200, nsave = 200) # Reasonable usage
pmc <- list(naprt = 5, nburn = 2, nsave = 5) # Minimal usage for documentation</pre>
# Fit the model:
res <- comix(Y, C, pmc = pmc, prior = prior)</pre>
# Relabel to resolve potential label switching issues:
res_relab <- relabelChain(res)</pre>
tidy_chain <- tidyChain(res_relab)</pre>
# (see vignette for a more detailed example)
```

transform\_params

Convert between parameterizations of the multivariate skew normal distribution.

#### **Description**

Convert between parameterizations of the multivariate skew normal distribution.

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## Usage

```
transform_params(Sigma, alpha)
```

## **Arguments**

Sigma A scale matrix.

alpha A vector for the skew parameter.

#### Value

#### A list:

• delta: a reparameterized skewness vector, a transformed version of alpha.

- omega: a diagonal matrix of the same dimensions as Sigma, the diagonal elements are the square roots of the diagonal elements of Sigma.
- psi: another reparameterized skewness vector, utilized in the sampler.
- G: a reparameterized version of Sigma, utilized in the sampler.

```
library(COMIX)
# Scale and skew parameters:
Sigma <- matrix(0.5, nrow = 4, ncol = 4) + diag(0.5, nrow = 4)
alpha <- c(0, 0, 0, 5)
transformed_parameters <- transform_params(Sigma, alpha)</pre>
```

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