# Package 'mSigHdp'

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ActivateAndBurnin AnalyzeAndPlotretval ChainsDiagnosticPlot CleanChlist CombinePosteriorChains ExtendBurnin GenerateAverageCluster Generateppindex MultipleSetupAndPosterior ParallelPosteriorafterBurnin
PrepInit

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#### **Description**

Prepare an hdpState-class object and run the Gibbs sampling burnin.

# Usage

```
ActivateAndBurnin(
  input.catalog,
  seedNumber = 1,
  K.guess,
  multi.types = FALSE,
  verbose = TRUE,
  burnin = 4000,
  cpiter = 3,
  burnin.verbosity = 0,
  gamma.alpha = 1,
  gamma.beta = 1
)
```

# **Arguments**

```
input.catalog
                 Input spectra catalog as a matrix or in ICAMS format.
seedNumber
                 An integer that is used to generate separate random seeds for the call to dp_activate,
                 and before the call of hdp_burnin.
K.guess
                 Suggested initial value of the number of signatures, passed to dp_activate
                 as initcc.
                 A logical scalar or a character vector. If FALSE, The HDP analysis will regard
multi.types
                 all input spectra as one tumor type.
                 If TRUE, the HDP analysis will infer tumor types based on the string before "::"
                 in their names. e.g. tumor type for "SA.Syn.Ovary-AdenoCA::S.500" would be
                 "SA.Syn.Ovary-AdenoCA"
                 If multi.types is a character vector, then it should be of the same length as
                 the number of columns in input.catalog, and each value is the name of the
                 tumor type of the corresponding column in input.catalog.
                 e.g. c("SA.Syn.Ovary-AdenoCA", "SA.Syn.Kidney-RCC").
verbose
                 If TRUE then message progress information.
burnin
                 Pass to hdp_burnin burnin.
                 Pass to hdp_burnin cpiter.
cpiter
```

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```
burnin.verbosity
```

Pass to hdp\_burnin verbosity.

gamma.alpha

Shape parameter of the gamma distribution prior for the Dirichlet process concentration parameters; in this function the gamma distributions for all Dirichlet processes, except possibly the top level process, are the same.

gamma.beta

Inverse scale parameter (rate parameter) of the gamma distribution prior for the Dirichlet process concentration parameters; in this function the gamma distributions for all Dirichlet processes, except possibly the top level process, are the same.

#### Value

A list with 2 elements:

hdplist A list representation of an hdpState-class object.

likelihood A numeric vector with the likelihood at each iteration.

AnalyzeAndPlotretval

Evaluate and plot retval from CombinePosteriorChains

# **Description**

Evaluate and plot retval from CombinePosteriorChains

# Usage

```
AnalyzeAndPlotretval(
  retval,
  out.dir = NULL,
  ground.truth.sig = NULL,
  ground.truth.exp = NULL,
  verbose = TRUE,
  overwrite = TRUE,
  diagnostic.plot = TRUE)
```

## **Arguments**

retval the output from function CombinePosteriorChains

out.dir Directory that will be created for the output; if overwrite is FALSE then

abort if out .dir already exits.

ground.truth.sig

Optional. Either a string with the path to file with ground truth signatures or and ICAMS catalog with the ground truth signatures. These are the signatures used to construct the ground truth spectra.

ground.truth.exp

Optional. Ground truth exposure matrix or path to file with ground truth exposures. If NULL skip checks that need this information.

verbose If TRUE then message progress information.

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If TRUE plot diagnostic plot. This is optional because there are cases having error

ChainsDiagnosticPlot

Diagnostic plot for a hdpSampleMulti object

# **Description**

Diagnostic plot for a hdpSampleMulti object

# Usage

```
ChainsDiagnosticPlot(retval, out.dir, verbose)
```

# **Arguments**

retval ou	tput from (	CombinePosterior	Chains.A lis	st with the	following	elements:
-----------	-------------	------------------	--------------	-------------	-----------	-----------

**signature** The extracted signature profiles as a matrix; rows are mutation types, columns are samples (e.g. tumors).

**exposure** The inferred exposures as a matrix of mutation counts; rows are signatures, columns are samples (e.g. tumors).

multi.chains A hdpSampleMulti-class object. This object has the method chains which returns a list of hdpSampleChain-class objects. Each of these sample chains objects has a method final\_hdpState (actually the methods seems to be just hdp) that returns the hdpState from which it was generated.

out.dir

Directory that will be created for the output; if overwrite is FALSE then abort if out.dir already exits.

verbose

If TRUE then message progress information.

CleanChlist

If the job of Gibbs sampling from MultipleSetupAndPosterior has an error caught by R, the corresponding element of chlist has class try-error. If the job is stopped with, e.g. a segfault, the chlist element is NULL.

# Description

If the job of Gibbs sampling from MultipleSetupAndPosterior has an error caught by R, the corresponding element of chlist has class try-error. If the job is stopped with, e.g. a segfault, the chlist element is NULL.

```
CleanChlist(chlist, verbose = FALSE)
```

CombinePosteriorChains 5

#### **Arguments**

chlist A list of hdpSampleChain-class objects.

verbose If TRUE then message progress information.

#### Value

Invisibly, the clean, non-error chlist This is a list of hdpSampleChain-class objects.

CombinePosteriorChains

Extract components and exposures from multiple posterior sample chains

# **Description**

Extract components and exposures from multiple posterior sample chains

# Usage

```
CombinePosteriorChains(
  clean.chlist,
  input.catalog,
  multi.types,
  cluster.method = "kmedians",
  verbose = TRUE,
  cos.merge = 0.9,
  categ.CI = 0.95,
  exposure.CI = 0.95,
  min.sample = 1
)
```

# **Arguments**

clean.chlist A list of hdpSampleChain-class objects. Each element is the result of one posterior sample chain.

input.catalog

Input spectra catalog as a matrix or in ICAMS format.

multi.types

A logical scalar or a character vector. If FALSE, The HDP analysis will regard all input spectra as one tumor type.

If TRUE, the HDP analysis will infer tumor types based on the string before "::" in their names. e.g. tumor type for "SA.Syn.Ovary-AdenoCA::S.500" would be "SA.Syn.Ovary-AdenoCA"

If multi.types is a character vector, then it should be of the same length as the number of columns in input.catalog, and each value is the name of the tumor type of the corresponding column in input.catalog.

e.g. c("SA.Syn.Ovary-AdenoCA", "SA.Syn.Kidney-RCC").

cluster.method

A kccaFamily object. Deprecated. Future code will use "kmedians".

verbose If TRUE then message progress information.

6 ExtendBurnin

The cosine similarity threshold for merging raw clusters from the posterior samcos.merge pling chains into "components" i.e. signatures; passed to hdp\_extract\_components. A number the range [0, 1]. The level of the confidence interval used in step 4 of categ.CI hdp\_merge\_and\_extract\_components. This governs when "averaged raw cluster" get assigned to component 0, i.e. if the the confidence interval overlaps 0. Lower values make it less likely that an averaged raw cluster will be assigned to component 0. The CI in question is for the number of mutations in a given mutation class (e.g. ACA > AAA, internally called a "category"). If, for every mutation class, this CI overlaps 0, then the averaged raw cluster goes to component 0. exposure.CI A number in the range [0,1]. The level of the confidence interval used in step 5 of hdp\_merge\_and\_extract\_components. The CI in question here for the total number of mutations assigned to an averaged raw cluster. A "component" (i.e. signature) must have at least this many samples; passed to min.sample hdp\_merge\_and\_extract\_components.

#### Value

Invisibly, a list with the following elements:

- **signature** The extracted signature profiles as a matrix; rows are mutation types, columns are samples (e.g. tumors).
- **exposure** The inferred exposures as a matrix of mutation counts; rows are signatures, columns are samples (e.g. tumors).
- multi.chains A hdpSampleMulti-class object. This object has the method chains which returns a list of hdpSampleChain-class objects. Each of these sample chains objects has a method final\_hdpState (actually the methods seems to be just hdp) that returns the hdpState from which it was generated.
- **sum\_raw\_clusters\_after\_cos\_merge** A matrix containing aggregated spectra of raw clusters after cosine similarity merge step in hdp\_merge\_and\_extract\_components.
- **sum\_raw\_clusters\_after\_nonzero\_categ** A matrix containing aggregated spectra of raw clusters after non-zero category selecting step in hdp\_merge\_and\_extract\_components.
- clust\_hdp0\_ccc4 A matrix containing aggregated spectra of raw clusters moving to hdp.0 after non-zero category selection step in hdp\_merge\_and\_extract\_components.
- **clust\_hdp0\_ccc5** A matrix containing aggregated spectra of raw clusters moving to hdp.0 after non-zero observation selection step in hdp\_merge\_and\_extract\_components.

ExtendBurnin	Extend Burn in iteration for a list representation of an
	hdpState-class <i>object. This list is an output from</i> hdp_burnin <i>or</i> ActivateandBurnin.

# **Description**

Extend Burn in iteration for a list representation of an hdpState-class object. This list is an output from hdp\_burnin or ActivateandBurnin.

```
ExtendBurnin(hdplist, seedNumber = 1, burnin = 4000, cpiter = 3, verbosity = 0)
```

#### **Arguments**

hdplist A list representation of an hdpState-class object

seedNumber A random seed for setting the environment of hdp\_burnin.

burnin Pass to hdp\_posterior burnin.

cpiter Pass to hdp\_posterior cpiter.

verbosity Pass to hdp\_posterior verbosity.

#### Value

A list with hdp object after burn-in iteration and likelihood of iteration

GenerateAverageCluster

Generate average pattern of clusters of each posterior chain from combined list of multiple posterior sample chains

# **Description**

Generate average pattern of clusters of each posterior chain from combined list of multiple posterior sample chains

# Usage

```
GenerateAverageCluster(clean.chlist)
```

# **Arguments**

clean.chlist A list of multiple (or one) posterior sample chains.

#### Value

A list of matrices containing the average pattern of clusters within each posterior chain and a list of matrices containing the sum of each cluster in each posterior chain

Generate ppindex Generate index for a HDP structure and num.tumor.types for other functions

# **Description**

Generate index for a HDP structure and num.tumor.types for other functions

```
Generateppindex(multi.types, input.catalog)
```

#### **Arguments**

multi.types

A logical scalar or a character vector. If FALSE, The HDP analysis will regard all input spectra as one tumor type.

If TRUE, the HDP analysis will infer tumor types based on the string before "::" in their names. e.g. tumor type for "SA.Syn.Ovary-AdenoCA::S.500" would be "SA.Syn.Ovary-AdenoCA"

If multi.types is a character vector, then it should be of the same length as the number of columns in input.catalog, and each value is the name of the tumor type of the corresponding column in input.catalog.

```
e.g. c("SA.Syn.Ovary-AdenoCA", "SA.Syn.Kidney-RCC").
```

input.catalog

Input spectra catalog as a matrix or in ICAMS format.

MultipleSetupAndPosterior

Activate hierarchical Dirichlet processes and run posterior sampling in parallel.

# **Description**

Activate hierarchical Dirichlet processes and run posterior sampling in parallel.

#### Usage

```
MultipleSetupAndPosterior(
  input.catalog,
  seedNumber = 1,
  K.guess,
  multi.types = FALSE,
  verbose = TRUE,
  post.burnin = 4000,
  post.n = 50,
  post.space = 50,
  post.cpiter = 3,
  post.verbosity = 0,
  CPU.cores = 1,
  num.child.process = 4,
  gamma.alpha = 1,
  gamma.beta = 1,
  gamma0.alpha = gamma.alpha,
  gamma0.beta = gamma.beta,
  checkpoint.chlist = TRUE
)
```

# **Arguments**

```
input.catalog
```

Input spectra catalog as a matrix or in ICAMS format.

seedNumber A random seeds passed to dp\_activate.

K.guess Suggested initial value of the number of signatures, passed to dp\_activate as initco.

multi.types A logical scalar or a character vector. If FALSE, The HDP analysis will regard all input spectra as one tumor type.

If TRUE, the HDP analysis will infer tumor types based on the string before "::" in their names. e.g. tumor type for "SA.Syn.Ovary-AdenoCA::S.500" would be "SA.Syn.Ovary-AdenoCA"

If multi.types is a character vector, then it should be of the same length as the number of columns in input.catalog, and each value is the name of the tumor type of the corresponding column in input.catalog.

e.g. c("SA.Syn.Ovary-AdenoCA", "SA.Syn.Kidney-RCC").

verbose If TRUE then message progress information.

post.burnin Pass to hdp\_posterior burnin.

post.n Pass to hdp\_posterior n.

post.space Pass to hdp\_posterior space.
post.cpiter Pass to hdp\_posterior cpiter.

post.verbosity

Pass to hdp\_posterior verbosity.

CPU.cores Number of CPUs to use; there is no point in making this larger than num.child.process.

num.child.process

Number of posterior sampling chains; can set to 1 for testing.

Shape parameter of the gamma distribution prior for the Dirichlet process concentration parameters; in this function the gamma distributions for all Dirichlet processes, except possibly the top level process, are the same.

Inverse scale parameter (rate parameter) of the gamma distribution prior for the Dirichlet process concentration parameters; in this function the gamma distributions for all Dirichlet processes, except possibly the top level process, are the same.

gamma 0.alpha See figure B.1 from Nicola Robert's thesis. The shape parameter  $(\alpha_0)$  of the gamma distribution priors for the Dirichlet process concentration parameters  $(\gamma_0)$  for  $G_0$ .

gamma 0 . beta See figure B.1 from Nicola Robert's thesis. Inverse scale parameter (rate parameter,  $\beta_0$ ) of the gamma distribution priors for the Dirichlet process concentration parameters  $(\gamma_0)$  for  $G_0$ .

checkpoint.chlist

If TRUE, checkpoint the (unclean) chlist to "initial.chlist.Rdata" in the current working directory. and checkpoint the clean chlist to "clean.chlist.Rdata" in the current working directory.

#### Value

Invisibly, the clean chlist (output of CleanChlist). This is a list of hdpSampleChain-class objects.

ParallelPosteriorafterBurnin

Generate an HDP Gibbs sampling chain from a spectra catalog.

# **Description**

Generate an HDP Gibbs sampling chain from a spectra catalog.

#### Usage

```
ParallelPosteriorafterBurnin(
  retval,
  seedNumber = 1,
  verbose = TRUE,
  post.burnin = 4000,
  post.n = 50,
  post.space = 50,
  post.cpiter = 3,
  post.verbosity = 0,
  num.child.process = 2,
  CPU.cores = 2
)
```

#### **Arguments**

```
A list object containing hdplist after burn-in iteration and likelihood from BurninIteration.
retval
seedNumber
                Pass to hdp_posterior
verbose
                If TRUE then message progress information.
                Pass to hdp_posterior burnin. This can be set to a small number
post.burnin
                Pass to hdp_posterior n.
post.n
                Pass to hdp_posterior space.
post.space
post.cpiter Pass to hdp_posterior cpiter.
post.verbosity
                Pass to hdp_posterior verbosity.
num.child.process
                Number of posterior sampling chains; can set to 1 for testing.
                Number of CPUs to use; there is no point in making this larger than num.child.process.
CPU.cores
```

# Value

Invisibly, an hdpSampleChain-class object as returned from hdp\_posterior.

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PrepInit Initialize hdp object Allocate process index for hdp initialization. Pre pare for hdp_init	PrepInit
---	----------

#### **Description**

Initialize hdp object Allocate process index for hdp initialization. Prepare for hdp\_init

# Usage

```
PrepInit(
  multi.types,
  input.catalog,
  verbose = TRUE,
  K.guess,
  gamma.alpha = 1,
  gamma.beta = 1,
  gamma0.alpha = gamma.alpha,
  gamma0.beta = gamma.beta
)
```

#### **Arguments**

multi.types A logical scalar or a character vector. If FALSE, The HDP analysis will regard all input spectra as one tumor type.

If TRUE, the HDP analysis will infer tumor types based on the string before "::" in their names. e.g. tumor type for "SA.Syn.Ovary-AdenoCA::S.500" would be "SA.Syn.Ovary-AdenoCA"

If multi.types is a character vector, then it should be of the same length as the number of columns in input.catalog, and each value is the name of the tumor type of the corresponding column in input.catalog.

e.g. c("SA.Syn.Ovary-AdenoCA", "SA.Syn.Kidney-RCC").

input.catalog

Input spectra catalog as a matrix or in ICAMS format.

verbose If TRUE then message progress information.

 $\textbf{K.guess} \qquad \qquad \textbf{Suggested initial value of the number of signatures, passed to } \\ \textbf{dp\_activate}$ 

as initcc.

gamma.alpha Shape parameter of the gamma distribution prior for the Dirichlet process con-

centration parameters; in this function the gamma distributions for all Dirichlet

processes, except possibly the top level process, are the same.

gamma . beta Inverse scale parameter (rate parameter) of the gamma distribution prior for the

Dirichlet process concentration parameters; in this function the gamma distributions for all Dirichlet processes, except possibly the top level process, are the

same.

gamma0.alpha See figure B.1 from Nicola Robert's thesis. The shape parameter  $(\alpha_0)$  of the

gamma distribution priors for the Dirichlet process concentration parameters

 $(\gamma_0)$  for  $G_0$ .

gamma0.beta See figure B.1 from Nicola Robert's thesis. Inverse scale parameter (rate param-

eter,  $\beta_0$ ) of the gamma distribution priors for the Dirichlet process concentration

parameters  $(\gamma_0)$  for  $G_0$ .

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RunHdpParallel

Extract mutational signatures and optionally compare them to existing signatures and exposures.

#### **Description**

Extract mutational signatures and optionally compare them to existing signatures and exposures.

# Usage

```
RunHdpParallel(
  input.catalog,
  seedNumber = 1,
  K.guess,
  multi.types = FALSE,
  verbose = TRUE,
  post.burnin = 4000,
  post.n = 50,
  post.space = 50,
  post.cpiter = 3,
  post.verbosity = 0,
  CPU.cores = 1,
  num.child.process = 4,
  cos.merge = 0.9,
  min.sample = 1,
  categ.CI = 0.95,
  exposure.CI = 0.95,
  cluster.method = "kmedians",
  ground.truth.sig = NULL,
  ground.truth.exp = NULL,
  overwrite = TRUE,
  out.dir = NULL,
  gamma.alpha = 1,
  gamma.beta = 1,
  gamma0.alpha = gamma.alpha,
  gamma0.beta = gamma.beta,
  checkpoint.chlist = TRUE
)
```

#### **Arguments**

input.catalog

Input spectra catalog as a matrix or in ICAMS format.

seedNumber A random seeds passed to dp\_activate.

Suggested initial value of the number of signatures, passes

K.guess Suggested initial value of the number of signatures, passed to dp\_activate as initco.

multi.types A logical scalar or a character vector. If FALSE, The HDP analysis will regard all input spectra as one tumor type.

If TRUE, the HDP analysis will infer tumor types based on the string before "::" in their names. e.g. tumor type for "SA.Syn.Ovary-AdenoCA::S.500" would be "SA.Syn.Ovary-AdenoCA"

If multi.types is a character vector, then it should be of the same length as the number of columns in input.catalog, and each value is the name of the tumor type of the corresponding column in input.catalog.

e.g. c("SA.Syn.Ovary-AdenoCA", "SA.Syn.Kidney-RCC").

verbose If TRUE then message progress information.

post.burnin Pass to hdp\_posterior burnin.

post.n Pass to hdp\_posterior n.

post.space Pass to hdp\_posterior space.
post.cpiter Pass to hdp\_posterior cpiter.

post.verbosity

Pass to hdp\_posterior verbosity.

CPU.cores Number of CPUs to use; there is no point in making this larger than num.child.process. num.child.process

Number of posterior sampling chains; can set to 1 for testing.

cos.merge The cosine similarity threshold for merging raw clusters from the posterior sam-

 $pling\ chains\ into\ "components"\ i.e.\ signatures; passed\ to\ hdp\_\texttt{extract\_components}.$ 

min.sample A "component" (i.e. signature) must have at least this many samples; passed to

hdp\_merge\_and\_extract\_components.

categ.CI A number the range [0,1]. The level of the confidence interval used in step 4 of

hdp\_merge\_and\_extract\_components. This governs when "averaged raw cluster" get assigned to component 0, i.e. if the the confidence interval overlaps 0. Lower values make it less likely that an averaged raw cluster will be assigned to component 0. The CI in question is for the number of mutations in a given mutation class (e.g. ACA > AAA, internally called a "category"). If, for every mutation class, this CI overlaps 0, then the averaged raw cluster goes to

component 0.

exposure.CI A number in the range [0,1]. The level of the confidence interval used in step 5 of hdp\_merge\_and\_extract\_components. The CI in question here for the total

number of mutations assigned to an averaged raw cluster.

cluster.method

A kccaFamily object. Deprecated. Future code will use "kmedians".

ground.truth.sig

Optional. Either a string with the path to file with ground truth signatures or and ICAMS catalog with the ground truth signatures. These are the signatures used to construct the ground truth spectra.

ground.truth.exp

Optional. Ground truth exposure matrix or path to file with ground truth expo-

sures. If NULL skip checks that need this information.

overwrite If TRUE overwrite out.dir if it exists, otherwise raise an error.

out.dir Directory that will be created for the output; if overwrite is FALSE then

abort if out . dir already exits.

gamma .alpha Shape parameter of the gamma distribution prior for the Dirichlet process con-

centration parameters; in this function the gamma distributions for all Dirichlet

processes, except possibly the top level process, are the same.

gamma .beta Inverse scale parameter (rate parameter) of the gamma distribution prior for the

Dirichlet process concentration parameters; in this function the gamma distributions for all Dirichlet processes, except possibly the top level process, are the

same.

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gamma 0.alpha See figure B.1 from Nicola Robert's thesis. The shape parameter  $(\alpha_0)$  of the gamma distribution priors for the Dirichlet process concentration parameters  $(\gamma_0)$  for  $G_0$ .

gamma 0 . beta See figure B.1 from Nicola Robert's thesis. Inverse scale parameter (rate parameter,  $\beta_0$ ) of the gamma distribution priors for the Dirichlet process concentration parameters  $(\gamma_0)$  for  $G_0$ .

checkpoint.chlist

If TRUE, checkpoint the (unclean) chlist to "initial.chlist.Rdata" in the current working directory. and checkpoint the clean chlist to "clean.chlist.Rdata" in the current working directory.

#### Value

Invisibly, a list with the following elements:

- **signature** The extracted signature profiles as a matrix; rows are mutation types, columns are samples (e.g. tumors).
- **exposure** The inferred exposures as a matrix of mutation counts; rows are signatures, columns are samples (e.g. tumors).
- multi.chains A hdpSampleMulti-class object. This object has the method chains which returns a list of hdpSampleChain-class objects. Each of these sample chains objects has a method final\_hdpState (actually the methods seems to be just hdp) that returns the hdpState from which it was generated.
- **sum\_raw\_clusters\_after\_cos\_merge** A matrix containing aggregated spectra of raw clusters after cosine similarity merge step in hdp\_merge\_and\_extract\_components.
- **sum\_raw\_clusters\_after\_nonzero\_categ** A matrix containing aggregated spectra of raw clusters after non-zero category selecting step in hdp\_merge\_and\_extract\_components.
- clust\_hdp0\_ccc4 A matrix containing aggregated spectra of raw clusters moving to hdp.0 after non-zero category selection step in hdp\_merge\_and\_extract\_components.
- **clust\_hdp0\_ccc5** A matrix containing aggregated spectra of raw clusters moving to hdp.0 after non-zero observation selection step in hdp\_merge\_and\_extract\_components.

SetupAndActivate Generate an HDP Gibbs sampling chain from a spectra catalog.

# Description

Generate an HDP Gibbs sampling chain from a spectra catalog.

```
SetupAndActivate(
  input.catalog,
  seedNumber = 1,
  K.guess,
  multi.types = FALSE,
  verbose = TRUE,
  gamma.alpha = 1,
  gamma.beta = 1,
  gamma0.alpha = gamma.alpha,
  gamma0.beta = gamma.beta
)
```

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#### **Arguments**

input.catalog I

Input spectra catalog as a matrix or in ICAMS format.

seedNumber A random seeds passed to dp\_activate.

K.guess Suggested initial value of the number of signatures, passed to dp\_activate

as initcc.

multi.types A logical scalar or a character vector. If FALSE, The HDP analysis will regard

all input spectra as one tumor type.

If TRUE, the HDP analysis will infer tumor types based on the string before "::" in their names. e.g. tumor type for "SA.Syn.Ovary-AdenoCA::S.500" would be

"SA.Syn.Ovary-AdenoCA"

If multi.types is a character vector, then it should be of the same length as the number of columns in input.catalog, and each value is the name of the

tumor type of the corresponding column in input.catalog.

e.g. c("SA.Syn.Ovary-AdenoCA", "SA.Syn.Kidney-RCC").

verbose If TRUE then message progress information.

gamma.alpha Shape parameter of the gamma distribution prior for the Dirichlet process con-

centration parameters; in this function the gamma distributions for all Dirichlet

processes, except possibly the top level process, are the same.

gamma .beta Inverse scale parameter (rate parameter) of the gamma distribution prior for the

Dirichlet process concentration parameters; in this function the gamma distributions for all Dirichlet processes, except possibly the top level process, are the

same.

gamma0.alpha See figure B.1 from Nicola Robert's thesis. The shape parameter  $(\alpha_0)$  of the

gamma distribution priors for the Dirichlet process concentration parameters

 $(\gamma_0)$  for  $G_0$ .

gamma 0. beta See figure B.1 from Nicola Robert's thesis. Inverse scale parameter (rate param-

eter,  $\beta_0$ ) of the gamma distribution priors for the Dirichlet process concentration

parameters  $(\gamma_0)$  for  $G_0$ .

#### Value

Invisibly, an hdpState-class object as returned from dp\_activate.

SetupAndPosterior Generate an HDP Gibbs sampling chain from a spectra catalog.

# **Description**

Generate an HDP Gibbs sampling chain from a spectra catalog.

```
SetupAndPosterior(
  input.catalog,
  seedNumber = 1,
  K.guess,
  multi.types = FALSE,
```

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```
verbose = TRUE,
post.burnin = 4000,
post.n = 50,
post.space = 50,
post.cpiter = 3,
post.verbosity = 0,
gamma.alpha = 1,
gamma0.beta = 1,
gamma0.beta = gamma.alpha,
gamma0.beta = gamma.beta
```

# **Arguments**

input.catalog

Input spectra catalog as a matrix or in ICAMS format.

seedNumber A random seeds passed to dp\_activate.

 $\textbf{K.guess} \qquad \qquad \textbf{Suggested initial value of the number of signatures, passed to} \ \ \textbf{dp\_activate}$ 

as initcc.

 $\verb|multi.types| A logical scalar or a character vector. If \verb|FALSE|, The HDP| analysis will regard|$ 

all input spectra as one tumor type.

If TRUE, the HDP analysis will infer tumor types based on the string before "::" in their names. e.g. tumor type for "SA.Syn.Ovary-AdenoCA::S.500" would be

"SA.Syn.Ovary-AdenoCA"

If multi.types is a character vector, then it should be of the same length as the number of columns in input.catalog, and each value is the name of the

 $tumor\ type\ of\ the\ corresponding\ column\ in\ {\tt input.catalog}.$ 

 $e.g. \; \texttt{c("SA.Syn.Ovary-AdenoCA", "SA.Syn.Kidney-RCC")}.$ 

verbose If TRUE then message progress information.

post.burnin Pass to hdp\_posterior burnin.

post.n Pass to hdp\_posterior n.

post.space Pass to hdp\_posterior space.
post.cpiter Pass to hdp\_posterior cpiter.

post.verbosity

Pass to hdp\_posterior verbosity.

gamma.alpha Shape parameter of the gamma distribution prior for the Dirichlet process con-

centration parameters; in this function the gamma distributions for all Dirichlet

processes, except possibly the top level process, are the same.

gamma .beta Inverse scale parameter (rate parameter) of the gamma distribution prior for the

Dirichlet process concentration parameters; in this function the gamma distributions for all Dirichlet processes, except possibly the top level process, are the

same

gamma0.alpha See figure B.1 from Nicola Robert's thesis. The shape parameter  $(\alpha_0)$  of the

gamma distribution priors for the Dirichlet process concentration parameters  $(\gamma_0)$  for  $G_0$ .

(

gamma 0.beta See figure B.1 from Nicola Robert's thesis. Inverse scale parameter (rate param-

eter,  $\beta_0$ ) of the gamma distribution priors for the Dirichlet process concentration

parameters  $(\gamma_0)$  for  $G_0$ .

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# Value

Invisibly, an hdpSampleChain-class object as returned from hdp\_posterior.

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