# Package 'texteffect'

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Title Discovering Latent Treatments in Text Corpora and Estimating Their Causal Effects

Author Christian Fong <christianfong@stanford.edu>

Maintainer Christian Fong <christianfong@stanford.edu>

**Depends** R (>= 3.3), MASS, boot, ggplot2

Description Implements the approach described in Fong and Grimmer (2016) <https: //aclweb.org/anthology/P/P16/P16-1151.pdf> for automatically discovering latent treatments from a corpus and estimating the average marginal component effect (AMCE) of each treatment. The data is divided into a training and test set. The supervised Indian Buffet Process (sibp) is used to discover latent treatments in the training set. The fitted model is then applied to the test set to infer the values of the latent treatments in the test set. Finally, Y is regressed on the latent treatments in the test set to estimate the causal effect of each treatment.

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BioSample

Sample from the Fong and Grimmer Wikipedia Biography Data

# Description

This data set gives a small sample of the data used in "Discovery of Treatments from Text Corpora" by Christian Fong and Justin Grimmer. This sample is intended as a toy data set for use in the examples of this package's documentation. A real data set should include far more observations.

# Usage

BioSample

# Format

A data frame consisting of 51 columns (including an outcome measure and counts for each word in a 50 word vocabulary) and 250 observations.

# Source

Data collected using the Wikipedia API and an original survey experiment by Fong and Grimmer.

#### References

Fong, Christian and Justin Grimmer. (2016). Discovery of Treatments from Text Corpora. Proceedings of the 54th Annual Meeting of the Association for Computational Linguistics, 1600-1609.

infer\_Z

Infer Treatments on the Test Set

#### Description

infer\_Z uses an sibp object fitted on a training set to infer the treatments in a test set.

# Usage

infer\_Z(sibp.fit, X, newX = FALSE)

# infer\_Z

# Arguments

sibp.fit	A sibp object.
Х	The covariates for the data set where Z is to be inferred. Usually, the user should Use the same X used to call the sibp function.
newX	Set to TRUE if the X supplied is not the training and test set. Used primarily for followup validation experiments. Defaults to FALSE.

# Details

This function applies the mapping from words to treatments discovered in the training set to infer which observations have which treatments in the test set. Usually, users will be better served by calling sibp\_amce, which calls this function internally before returning estimates and confidence intervals for the average marginal component effects.

#### Value

nu

Informally, the probability that the row document has the column treatment. Formally, the parameter for the variatioanl approximation of z\_i,k, which is a Bernoulli distribution.

# Author(s)

Christian Fong

# References

Fong, Christian and Justin Grimmer. 2016. "Discovery of Treatments from Text Corpora" Proceedings of the 54th Annual Meeting of the Association for Computational Linguistics. https: //aclweb.org/anthology/P/P16/P16-1151.pdf

#### See Also

sibp, sibp\_amce

# Examples

```
##Load the Wikipedia biography data
data(BioSample)
```

```
# Divide into training and test sets
Y <- BioSample[,1]
X <- BioSample[,-1]
set.seed(1)
train.ind <- sample(1:nrow(X), size = 0.5*nrow(X), replace = FALSE)
# Fit an sIBP on the training data
sibp.fit <- sibp(X, Y, K = 2, alpha = 4, sigmasq.n = 0.8,
train.ind = train.ind)
# Infer the latent treatments in the test set</pre>
```

infer\_Z(sibp.fit, X)

sibp

# Description

sibp discovers latent binary treatments within a corpus, as described by Fong and Grimmer (2016).

#### Usage

```
sibp(X, Y, K, alpha, sigmasq.n,
a = 0.1, b = 0.1, sigmasq.A = 5,
train.ind, G = NULL, silent = FALSE )
```

#### Arguments

Х	The covariates for all observations in the training set, where each row is a document and each column is the count of a word.
Υ	The outcome for all observations in the training set.
К	The number of treatments to be discovered.
alpha	A parameter that influences how common the treatments are. When alpha is large, the treatments are common.
sigmasq.n	A parameter determining the variance of the word counts conditional on the treatments. When sigmasq.n is large, the treatments must explain most of the variation in X.
а	A parameter that, together with b, influences the variance of the treatment effects and the outcomes. $a = 0.1$ is a reasonably diffuse choice.
b	A parameter that, together with a, influences the variance of the treatment effects and the outcomes. $b = 0.1$ is a reasonably diffuse choice.
sigmasq.A	A parameter determining the variance of the effect of the treatments on word counts. A diffuse choice, such as 5, is usually appropriate.
train.ind	The indices of the observations in the training set, usually obtained from get_training_set().
G	An optional group membership matrix. The AMCE for a given treatment is permitted to vary as a function of the individual's group.
silent	If TRUE, prints how much the parameters have moved every 10 iterations of sIBP.

# Details

Fits a supervised Indian Buffet Process using variational inference. Before running this function, the data should be divided into a training set and a test set. This function should be run on the training set to discover latent treatments in the data that seem to be correlated with the outcome.

It is recommended to use linksibp\_param\_search instead of this function to search over multiple configurations of the most important parameters. So long as only the training data is used, the analyst can freely experimient with as many parameter configurations as he likes without corrupting his causal inferences. Once a parameter configuration is chosen, the user can then use sibp\_amce on the test set to estimate the average marginal component effect (AMCE) for each treatment.

# Value

nu	Informally, the probability that the row document has the column treatment. Formally, the parameter for the variational approximation of z_i,k, which is a Bernoulli distribution.
m	Informally, the effect of having each treatment on the outcome. Formally, the mean parameter for the variational approximation of the posterior distribution of beta, which is a normal distribution. Note that this is in the training sample, and it is inappropriate to use this posterior as the basis for causal inference. It is instead necessary to estimate effects using the test set, see sibp_amce.
S	The variance parameter for the posterior distribution of beta, which is a normal distribution.
lambda	A matrix where the kth row contains the shape parameters for the variational approximation of the posterior distribution of pi_k, which is a beta distribution.
phi	Informally, the effect of the row treatment on the column word. Formally, the mean parameter for the variational approximation of the posterior distribution of A, which is a normal distribution.
big.Phi	The variance parameter for the variational approximation of the posterior distribution of A, which is a normal distribution. The kth element of the list corresponds to a treatment k.
с	The shape parameter for the variational approximation of the posterior distribu- tion of tau, which is a gamma distribution.
d	The rate parameter for the variational approximation of the posterior distribution of tau, which is a gamma distribution.
К	The number of treatments.
D	The number of words in the vocabulary.
alpha	The alpha used to call this function.
а	The a used to call this function.
b	The b used to call this function.
sigmasq.A	The sigmasq.A used to call this function.
sigmasq.n	The sigmasq.n used to call this function.
train.ind	The indices of the observations in the training set.
test.ind	The indices of the observations in the test set.

# Author(s)

Christian Fong

# References

Fong, Christian and Justin Grimmer. 2016. "Discovery of Treatments from Text Corpora" Proceedings of the 54th Annual Meeting of the Association for Computational Linguistics. https://aclweb.org/anthology/P/P16/P16-1151.pdf

# See Also

sibp\_param\_search, sibp\_top\_words, sibp\_amce

# Examples

```
##Load the Wikipedia biography data
data(BioSample)
# Divide into training and test sets
Y <- BioSample[,1]</pre>
X <- BioSample[,-1]</pre>
set.seed(1)
train.ind <- sample(1:nrow(X), size = 0.5*nrow(X), replace = FALSE)</pre>
# Search sIBP for several parameter configurations; fit each to the training set
sibp.search <- sibp_param_search(X, Y, K = 2, alphas = c(2,4), sigmasq.ns = c(0.8, 1),
 iters = 1, train.ind = train.ind)
## Not run:
# Get metric for evaluating most promising parameter configurations
sibp_rank_runs(sibp.search, X, 10)
# Qualitatively look at the top candidates
sibp_top_words(sibp.search[["4"]][["0.8"]][[1]], colnames(X), 10, verbose = TRUE)
sibp_top_words(sibp.search[["4"]][["1"]][[1]], colnames(X), 10, verbose = TRUE)
# Select the most interest treatments to investigate
sibp.fit <- sibp.search[["4"]][["0.8"]][[1]]</pre>
# Estimate the AMCE using the test set
amce<-sibp_amce(sibp.fit, X, Y)</pre>
# Plot 95% confidence intervals for the AMCE of each treatment
sibp_amce_plot(amce)
## End(Not run)
```

sibp\_amce

Infer Treatments on the Test Set

# Description

sibp\_amce uses an sibp object fitted on a training set to estimate the AMCE with the test set.

# Usage

```
sibp_amce(sibp.fit, X, Y, G = NULL, seed = 0, level = 0.05, thresh = 0.5)
sibp_amce_plot(sibp.amce, xlab = "Feature", ylab = "Outcome", subs = NULL)
```

# sibp\_amce

# Arguments

sibp.fit	A sibp object.
Х	The covariates for the full data set. The division between the training and test set is handled inside the function.
Y	The outcomes for the full data set. The division between the training and test set is handled inside the function.
G	A group membership matrix. The AMCE for a given treatment is permitted to vary as a function of the individual's group.
seed	The seed
level	The level of the confidence intervals to be obtained.
thresh	The treatment will = 1 when $nu \ge thresh and 0$ otherwise. This avoids problems due to misclassification error.
sibp.amce	The table returned by codesibp_amce.
xlab	The label for the x-axis of the plot.
ylab	The label for the y-axis of the plot.
subs	The susbet of the coefficients to plot. By default, plots all coefficients.

# Details

Nothing

# Value

sibp.amce A table where the first column is the index of the treatment, the second column ("effect") is the estimated AMCE, the third column ("L") is the lower bound of the confidence interval, and the fourth column ("U") is the upper bound of the confidence interval.

# Author(s)

Christian Fong

# References

Fong, Christian and Justin Grimmer. 2016. "Discovery of Treatments from Text Corpora" Proceedings of the 54th Annual Meeting of the Association for Computational Linguistics. https://aclweb.org/anthology/P/P16/P16-1151.pdf

# See Also

sibp

# Examples

```
##Load the sample of Wikipedia biography data
data(BioSample)
# Divide into training and test sets
Y <- BioSample[,1]
X <- BioSample[,-1]
set.seed(1)
train.ind <- sample(1:nrow(X), size = 0.5*nrow(X), replace = FALSE)
# Fit an sIBP on the training data
sibp.fit <- sibp(X, Y, K = 2, alpha = 4, sigmasq.n = 0.8,
train.ind = train.ind)
sibp.amce <- sibp_amce(sibp.fit, X, Y)
sibp_amce_plot(sibp.amce)</pre>
```

sibp\_exclusivity Calculate Exclusivity Metric

#### Description

sibp\_exculsivity calculates the coherence metric for an sibp object fit on a training set. sibp\_rank\_runs runs sibp\_exclusivity on each element in the list returned by sibp\_param\_search, and ranks the parameter configurations from most to least promising.

# Usage

```
sibp_exclusivity(sibp.fit, X, num.words = 10)
sibp_rank_runs(sibp.search, X, num.words = 10)
```

# Arguments

sibp.fit	A sibp object.
sibp.search	A list of sibp object fit using the training set, obtained using sibp_param_search.
Х	The covariates for the full data set. The division between the training and test set is handled inside the function.
num.words	The top words whose coherence will be evaluated.

# Details

The metric is formally described at the top of page 1605 of https://aclweb.org/anthology/P/ P16/P16-1151.pdf. The purpose of this metric is merely to suggest which parameter configurations might contain the most interesting treatments to test if there are too many configurations to investigate manually. The choice of the parameter configuration should always be made on the basis of which treatments are substantively the most interesting, see sibp\_top\_words.

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# Value

exclusivity An exclusivity matrix which quantifies the degree to which the top words in a treatment appear in documents that have that treatment but not in documents that lack that treatment.

#### exclusivity\_rank

A table that ranks the treatments discovered by the various runs from sibp.search from most exclusive to least exclusive.

# Author(s)

Christian Fong

#### References

Fong, Christian and Justin Grimmer. 2016. "Discovery of Treatments from Text Corpora" Proceedings of the 54th Annual Meeting of the Association for Computational Linguistics. https://aclweb.org/anthology/P/P16/P16-1151.pdf

#### See Also

sibp\_param\_search, sibp\_top\_words

#### Examples

sibp\_param\_search Search Parameter Configurations for Supervised Indian Buffet Process (sibp)

#### Description

sibp\_param\_search runs sibp for a variety of parameter configurations, so that the user can then test the effects fo the most interesting treatments.

# Usage

```
sibp_param_search(X, Y, K, alphas, sigmasq.ns, iters,
a = 0.1, b = 0.1, sigmasq.A = 5, train.ind = train.ind,
G = NULL, seed = 0)
```

#### Arguments

Х	The covariates for the full data set. The division between the training and test set is handled inside the function.
Y	The outcomes for the full data set. The division between the training and test set is handled inside the function.
К	The number of treatments to be discovered.
alphas	A vector of values of alpha to try.
sigmasq.ns	A vector of values of sigmasq.n to try.
iters	The number of starting values to attempt for each combination of alpha and sigmasq.n.
a	A parameter.
b	A parameter.
sigmasq.A	A parameter.
train.ind	The indices of the observations in the training set, usually obtained from get_training_set().
G	An optional group membership matrix. The AMCE for a given treatment is permitted to vary as a function of the individual's group.
seed	The seed to be used, so the result can be replicated.

# Details

Fits a supervised Indian Buffet Process using variational inference for combinations of alpha and sigmasq.n. alpha influences how common the treatments are (where larger alphas imply more common treatments) and sigmasq.n influences how much of the variation of the outcome must be explained by the treatments. These parameters are the most important for determining the quality of the treatments discovered, so it is usually a good idea to experiment with many combinations. Because the treatments discovered can be sensitive to starting values, it is also usually a good idea to try each combination of alpha and sigmasq.n several times by setting iters > 1.

Because this function uses only the training data, the user can experiment with many parameter configurations without corrupting the inferences made with the test set. The choice of parameters is equivalent to the choice of hypotheses to test, so the analyst should choose the parameter configuration that leads to the most substantively interesting treatments. sibp\_top\_words can be applied to each element of the list returned by this function to determine which parameter configurations lead to interesting treatments. Often, it will be impractical to manually investigate every parameter configuration. In such cases, sibp\_rank\_runs can be used to automatically identify some of the most promising candidates.

#### Value

paramslist

sibp\_top\_words

# Author(s)

Christian Fong

#### References

Fong, Christian and Justin Grimmer. 2016. "Discovery of Treatments from Text Corpora" Proceedings of the 54th Annual Meeting of the Association for Computational Linguistics. https://aclweb.org/anthology/P/P16/P16-1151.pdf

# See Also

sibp\_rank\_runs, sibp\_top\_words, sibp\_amce

#### Examples

```
##Load the sample of Wikipedia biography data
data(BioSample)
# Divide into training and test sets
Y <- BioSample[,1]</pre>
X <- BioSample[,-1]</pre>
set.seed(1)
train.ind <- sample(1:nrow(X), size = 0.5*nrow(X), replace = FALSE)</pre>
# Search sIBP for several parameter configurations; fit each to the training set
sibp.search <- sibp_param_search(X, Y, K = 2, alphas = c(2,4),</pre>
                                  sigmasq.ns = c(0.8, 1), iters = 1,
     train.ind = train.ind)
## Not run:
# Get metric for evaluating most promising parameter configurations
sibp_rank_runs(sibp.search, X, 10)
# Qualitatively look at the top candidates
sibp_top_words(sibp.search[["4"]][["0.8"]][[1]], colnames(X), 10, verbose = TRUE)
sibp_top_words(sibp.search[["4"]][["1"]][[1]], colnames(X), 10, verbose = TRUE)
## End(Not run)
```

sibp\_top\_words Report Words Most Associated with each Treatment

# Description

sibp\_top\_words returns a data frame of the words most associated with each treatment.

#### Usage

```
sibp_top_words(sibp.fit, words, num.words = 10, verbose = FALSE)
```

# Arguments

sibp.fit	A sibp object.
words	The actual words, usually obtained through colnames(X).
num.words	The number of top words to report.
verbose	If set to true, reports how common each treatment is (so that the analyst can focus on the common treatments) and how closely associated each word is with each treatment.

#### Details

The choice of the parameter configuration should always be made on the basis of which treatments are substantively the most interesting. This function provides one natural way of discovering which words are most associated with each treatment (the mean parameter for the posterior distribution of phi, where phi is the effect of the treatment on the count of word w) and therefore helps to determine which treatments are most interesting.

# Value

top.words	A data frame where each column consists of the top ten words (in order) associ
	ated with a given treatment.

#### Author(s)

Christian Fong

# References

Fong, Christian and Justin Grimmer. 2016. "Discovery of Treatments from Text Corpora" Proceedings of the 54th Annual Meeting of the Association for Computational Linguistics. https://aclweb.org/anthology/P/P16/P16-1151.pdf

# See Also

sibp

# Examples

##Load the Wikipedia biography data
data(BioSample)

```
# Divide into training and test sets
Y <- BioSample[,1]
X <- BioSample[,-1]
set.seed(1)
train.ind <- sample(1:nrow(X), size = 0.5*nrow(X), replace = FALSE)
# Fit an sIBP on the training data
sibp.fit <- sibp(X, Y, K = 2, alpha = 4, sigmasq.n = 0.8,
train.ind = train.ind)</pre>
```

sibp\_top\_words

sibp\_top\_words(sibp.fit, colnames(X))

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