

covdepGE: Covariate Dependent Graph Estimation

Installation

Run the following in R:

```
devtools::install_github("JacobHelwig/covdepGE")
```

Overview

Suppose $\mathbf{X} \in \mathbb{R}^{n \times p}$ is a data matrix of independent observations $\mathbf{X} = (\mathbf{x}_1, \dots, \mathbf{x}_p)$, where, for $j \in 1, \dots, p$:

$$\mathbf{x}_j \sim \mathcal{N}(\mu_j, \Sigma_{j,j}), \mathbf{x}_j \in \mathbb{R}^n \quad \mathbf{X} \sim \mathcal{N}(\mu, \Sigma) \quad (1)$$

The conditional dependence structure of $\mathbf{x}_1, \dots, \mathbf{x}_p$ can be modeled as an undirected graph \mathcal{G} such that:

$$\mathcal{G}_{i,j} = \begin{cases} 1 & \iff \text{Cov}(\mathbf{x}_i, \mathbf{x}_j) \neq 0 \\ 0 & \text{otherwise} \end{cases} \quad (2)$$

That is, there is an edge between the \mathbf{x}_i and \mathbf{x}_j nodes if, and only if, these variables are dependent on each other given all other variables.

Let \mathbf{Z} be an $n \times p'$ matrix of extraneous covariates. Further suppose that the conditional dependence structure of \mathbf{X} is not homogeneous across the individuals, and is instead a continuous function of the extraneous covariates $\mathbf{Z}(1)$. Then, this methodology aims to estimate a graph for each of the individuals, possibly unique to the individual, such that similar estimates are made for those who are similar to one another in terms of the extraneous covariates.

For an example application, see (1), wherein the sample was composed of healthy and cancerous individuals, $\mathbf{x}_1, \dots, \mathbf{x}_8$ were protein expression levels of 8 genes, and \mathbf{Z} was the copy number variation of a gene \mathbf{z} associated with cancer, $\mathbf{z} \notin \{\mathbf{x}_1, \dots, \mathbf{x}_8\}$.

Functionality

The main function, `covdepGE::covdepGE($\pmb{X}, \pmb{Z})`, estimates the posterior distribution of the graphical structure \mathcal{G}_l for each of the n individuals using a variational mean-field approximation. The function will output $n \times p \times p$ symmetric matrices \mathcal{A}_l , where $\mathcal{A}_{i,j}^{(l)}$ is the posterior inclusion probability of an edge between the node representing the i -th variable and the node representing the j -th variable.

There are also two plotting functions, `gg_adjMat` and `gg_inclusionCurve`, for visualizing the result of `covdepGE`. The primary functionality of `gg_adjMat` is to visualize the estimate to a specified individuals conditional dependence structure, however, it can also be used to visualize any numeric matrix. `gg_inclusionCurve` plots the posterior inclusion probability of an edge between two specified variables across all n individuals.

Demo

```
# install the package if necessary
if (!("covdepGE" %in% installed.packages())){
  devtools::install_github("JacobHelwig/covdepGE")
}
```

```
library(covdepGE)
```

```
set.seed(1)
```

```
n <- 100
```

```
p <- 4
```

```
# generate the extraneous covariate
```

```
Z_neg <- sort(runif(n / 2) * -1)
```

```
Z_pos <- sort(runif(n / 2))
```

```
Z <- c(Z_neg, Z_pos)
```

```
summary(Z)
```

```
##      Min.   1st Qu.   Median     Mean   3rd Qu.     Max.
```

```
## -0.99191 -0.55799  0.02277 -0.01475  0.45622  0.96062
```

```
# create true covariance structure for 2 groups: positive Z and negative Z
```

```
true_graph_pos <- true_graph_neg <- matrix(0, p + 1, p + 1)
```

```
true_graph_pos[1, 2] <- true_graph_pos[2, 1] <- 1
```

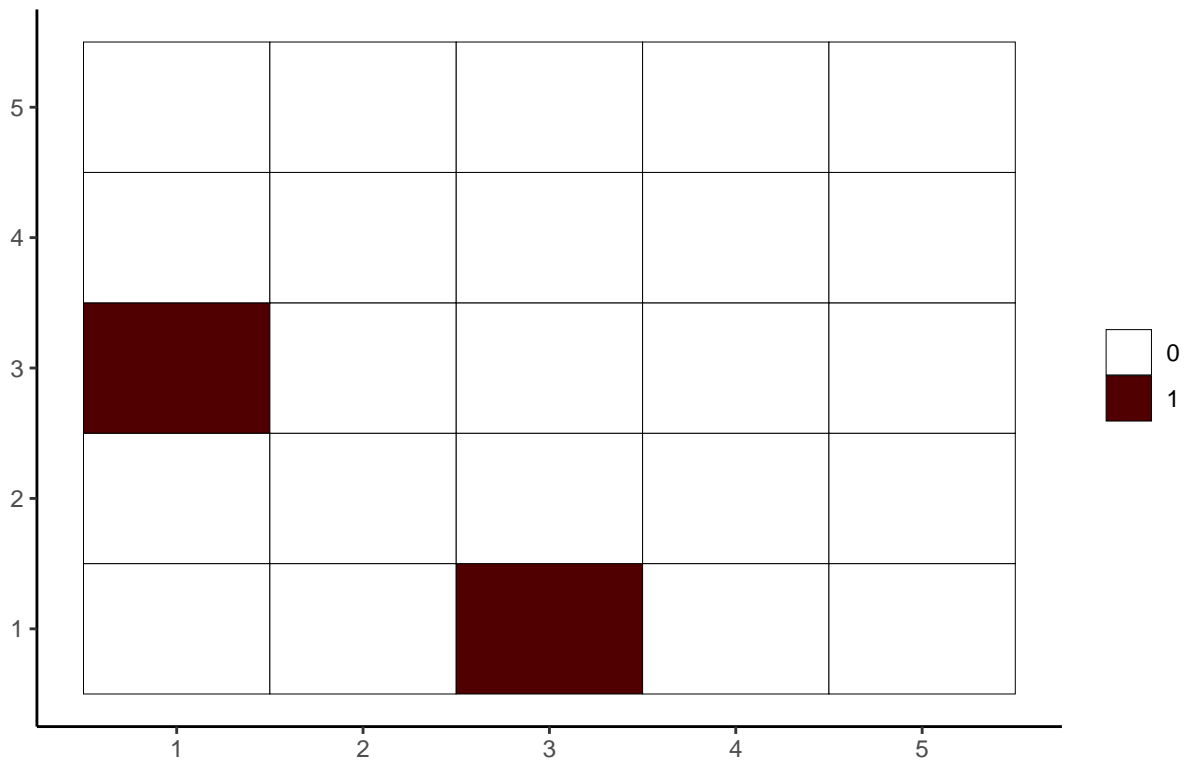
```
true_graph_neg[1, 3] <- true_graph_neg[3, 1] <- 1
```

```
# visualize the true covariance structures
```

```
(gg_adjMat(true_graph_neg) +
```

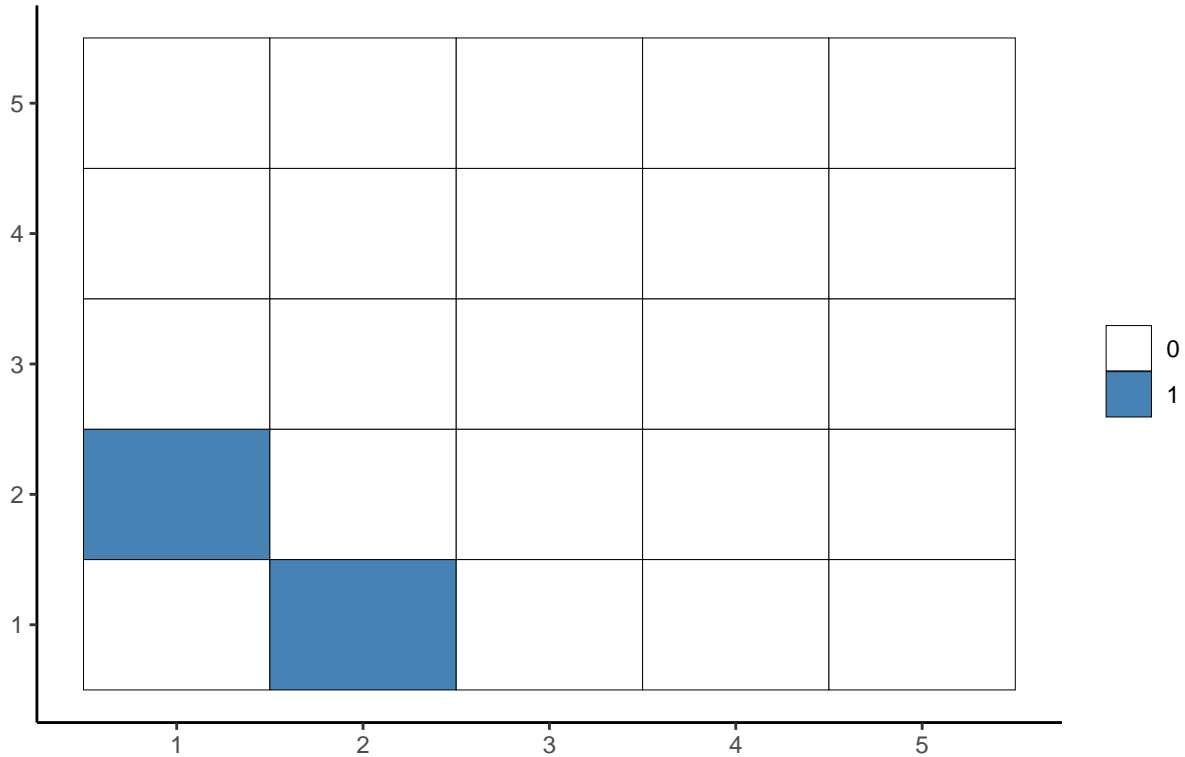
```
  ggplot2::ggtitle("True graph for individuals with negative Z (1,...,50)"))
```

True graph for individuals with negative Z (1,...,50)



```
(gg_adjMat(true_graph_pos, color1 = "steelblue") +
  ggplot2::ggtitle("True graph for individuals with positive Z (51,...,100)"))
```

True graph for individuals with positive Z (51,...,100)

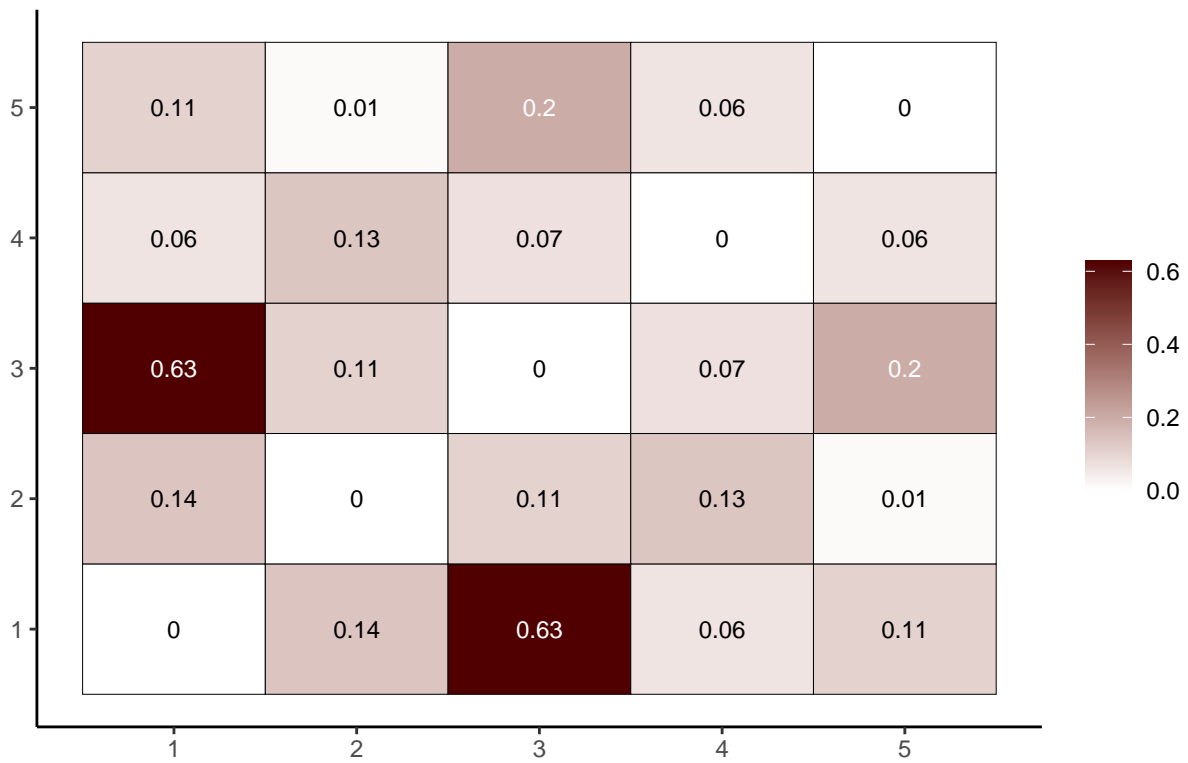


```
# generate the covariance matrices as a function of Z
sigma_mats_neg <- lapply(Z_neg, function(z) z * true_graph_neg + diag(p + 1))
sigma_mats_pos <- lapply(Z_pos, function(z) z * true_graph_pos + diag(p + 1))
sigma_mats <- c(sigma_mats_neg, sigma_mats_pos)

# generate the data using the covariance matrices
data_mat <- t(sapply(sigma_mats, MASS::mvrnorm, n = 1, mu = rep(0, p + 1)))

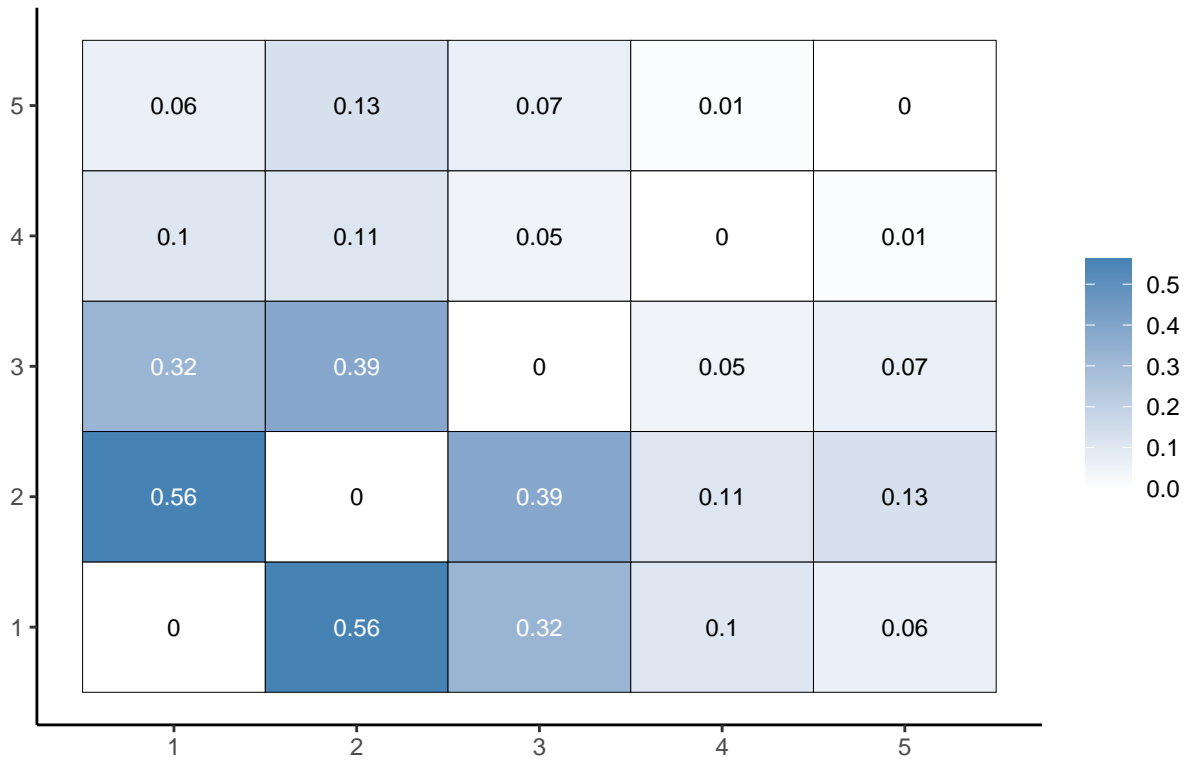
# visualize the sample correlation within each group
(gg_adjMat(abs(cor(data_mat[1:(n / 2), ])) - diag(p + 1)) +
  ggplot2::ggtitle("Correlation Matrix for Negative Z (1,...,50)"))
```

Correlation Matrix for Negative Z (1,...,50)



```
(gg_adjMat(abs(cor(data_mat[(n / 2 + 1):n, ])) - diag(p + 1),
  color1 = "steelblue") +
  ggplot2::ggtitle("Correlation Matrix for Positive Z (51,...,100)"))
```

Correlation Matrix for Positive Z (51,...,100)



```
# use varbvs to get the hyperparameter sigma
sigmasq <- rep(NA, p + 1)
for (j in 1:(p + 1)){
  sigmasq[j] <- mean(varbvs::varbvs(data_mat[ , -j], Z, data_mat[ , j], verbose = F)$sigma)
}
#sigmasq
#mean(sigmasq)

# estimate the covariance structure
#?covdepGE
out <- covdepGE(
  data_mat,
  Z, # extraneous covariates
  kde = T, # whether KDE should be used to calculate bandwidths
  sigmasq = mean(sigmasq), # hyperparameter residual variance
  var_min = 1e-4, # smallest sigmabeta_sq grid value
  var_max = 1, # largest sigmabeta_sq grid value
  n_sigma = 10, # length of the sigmabeta_sq grid
  pi_vec = seq(0.1, 0.3, 0.05), # prior inclusion probability grid
  norm = Inf, # norm to calculate the weights
  scale = T, # whether the extraneous covariates should be scaled
  tolerance = 1e-15, # variational parameter exit condition 1
  max_iter = 200, # variational parameter exit condition 2
  edge_threshold = 0.75, # minimum inclusion probability
  sym_method = "min", # how to symmetrize the alpha matrices
  print_time = T,
```

```

        warnings = T # whether warnings should be displayed
    )

## Warning in covdepGE(data_mat, Z, kde = T, sigmasq = mean(sigmasq), var_min =
## 1e-04, : Response 1: 3/50 candidate models did not converge in 200 iterations

## Warning in covdepGE(data_mat, Z, kde = T, sigmasq = mean(sigmasq), var_min =
## 1e-04, : Response 2: 17/50 candidate models did not converge in 200 iterations

## Warning in covdepGE(data_mat, Z, kde = T, sigmasq = mean(sigmasq), var_min =
## 1e-04, : Response 2: final model did not converge in 200 iterations

## Warning in covdepGE(data_mat, Z, kde = T, sigmasq = mean(sigmasq), var_min =
## 1e-04, : Response 3: 17/50 candidate models did not converge in 200 iterations

## Warning in covdepGE(data_mat, Z, kde = T, sigmasq = mean(sigmasq), var_min =
## 1e-04, : Response 3: final model did not converge in 200 iterations

## Warning in covdepGE(data_mat, Z, kde = T, sigmasq = mean(sigmasq), var_min =
## 1e-04, : For 1/5 responses, the selected value of sigmabeta_sq was on the grid
## boundary. See return value ELBO for details

## Warning in covdepGE(data_mat, Z, kde = T, sigmasq = mean(sigmasq), var_min =
## 1e-04, : For 5/5 responses, the selected value of pi was on the grid boundary.
## See return value ELBO for details

## Time difference of 3.737961 secs

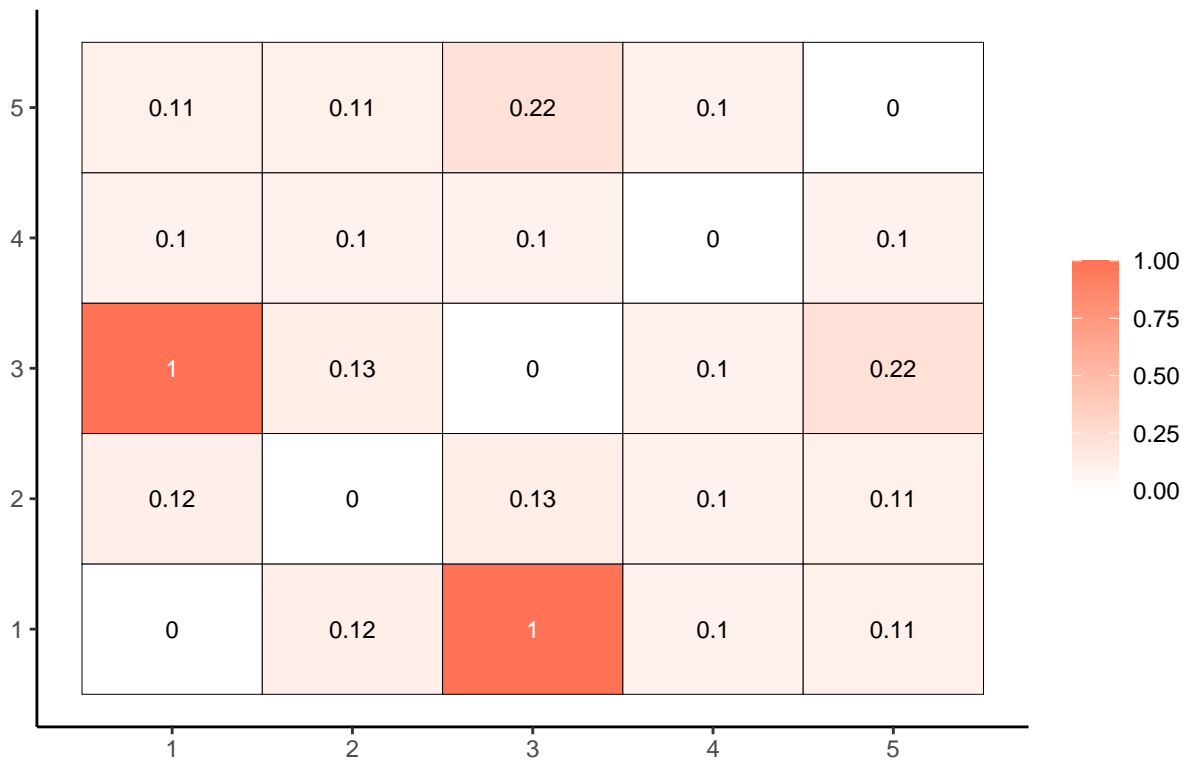
# grid search results
#out$ELBO

# individual-specific bandwidths calculated using KDE
#out$bandwidths

# analyze results
#?gg_adjMat
gg_adjMat(out, 1, color1 = "coral1")

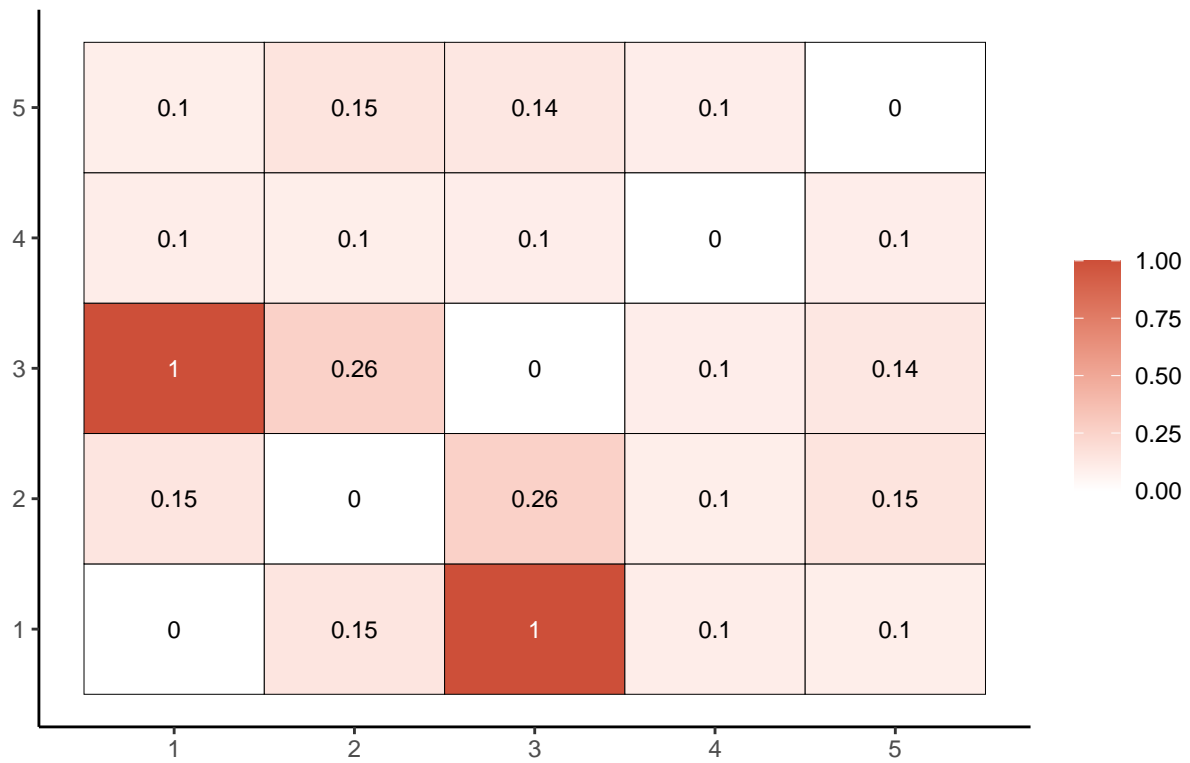
```

Posterior inclusion probabilities for individual 1



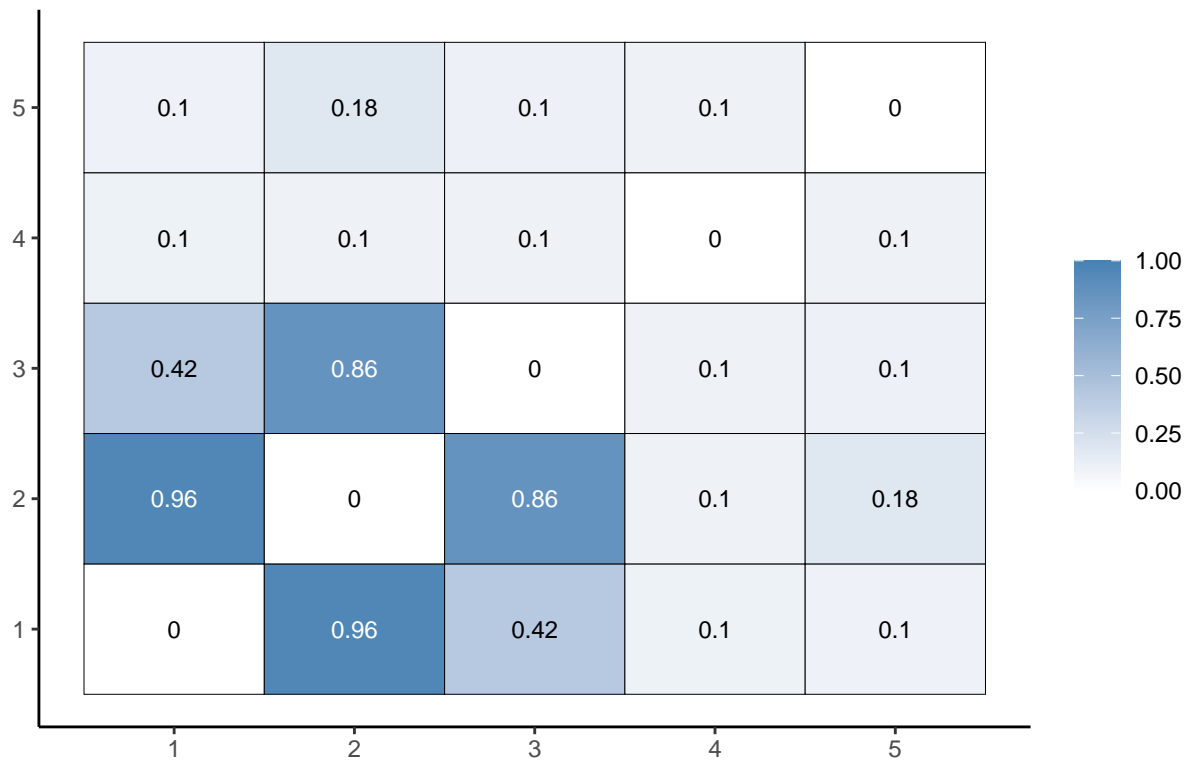
```
gg_adjMat(out, 50, color1 = "tomato3")
```

Posterior inclusion probabilities for individual 50



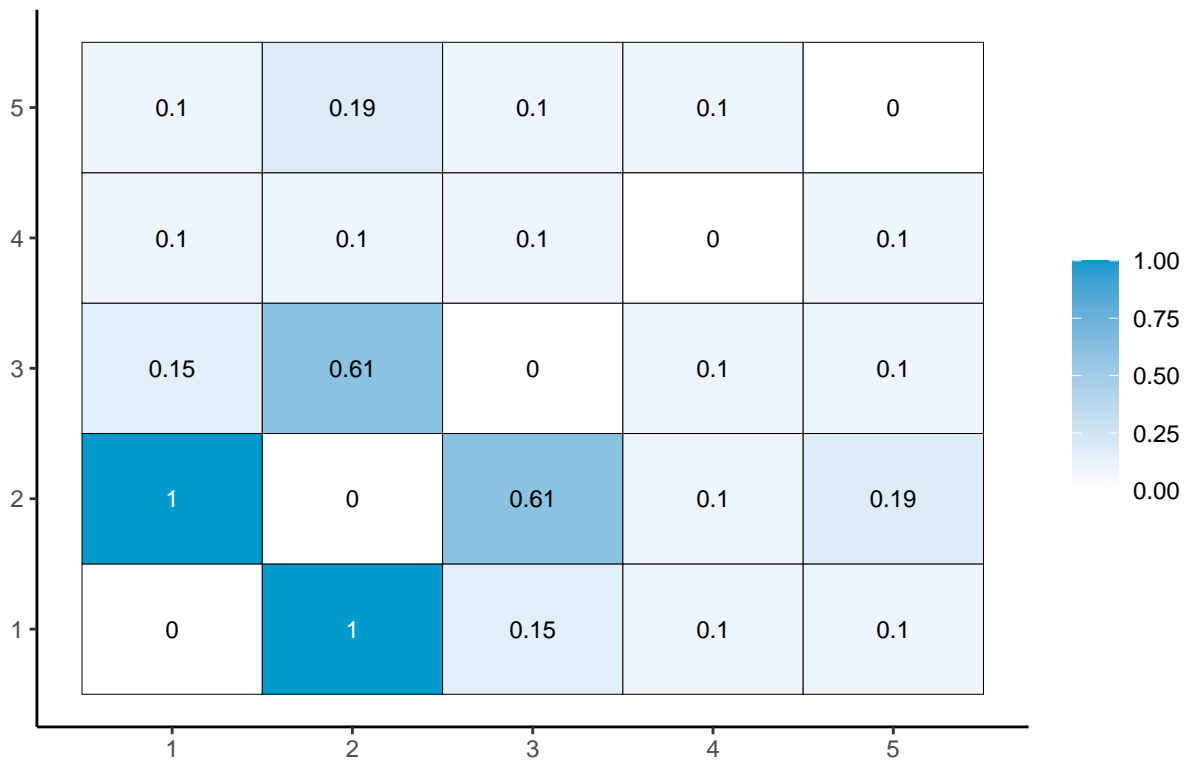
```
gg_adjMat(out, 60, color1 = "steelblue")
```


Posterior inclusion probabilities for individual 60

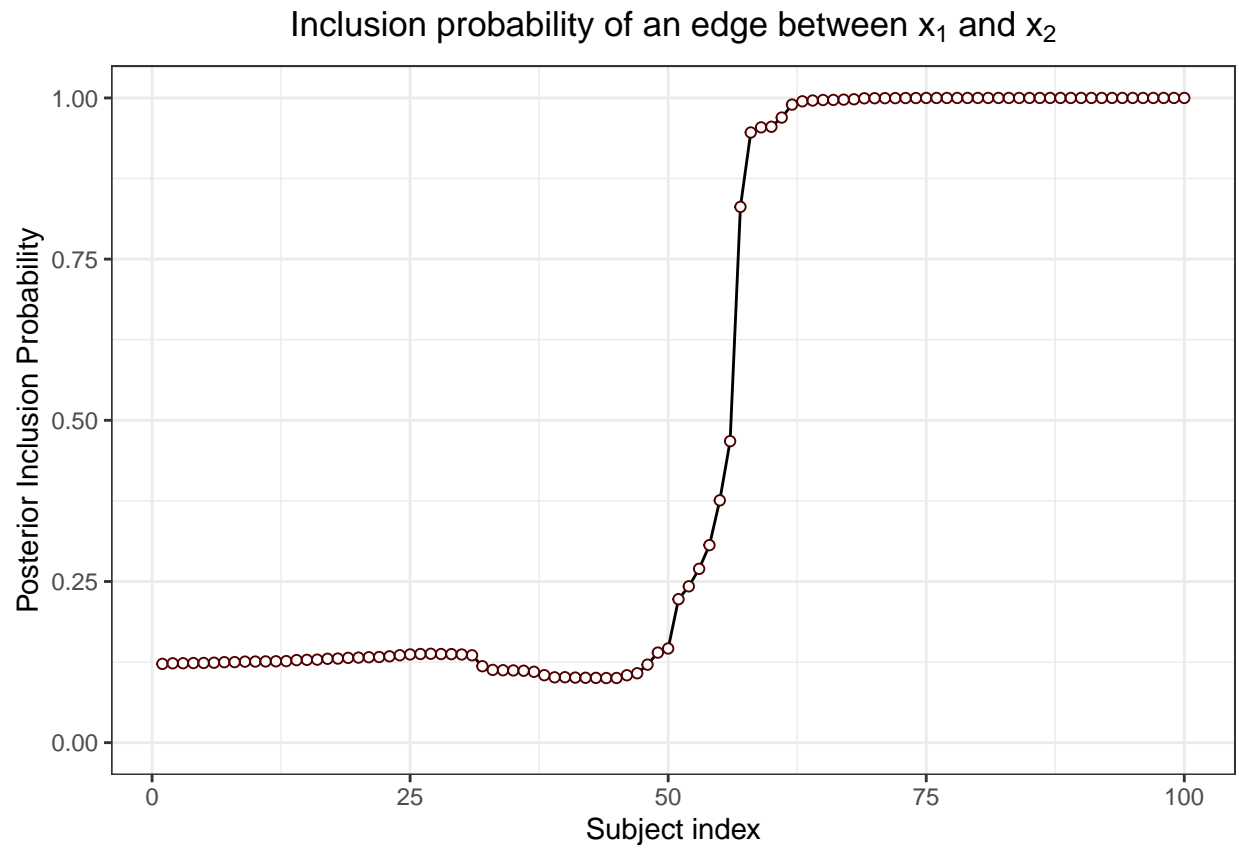


```
gg_adjMat(out, 100, color1 = "deepskyblue3")
```

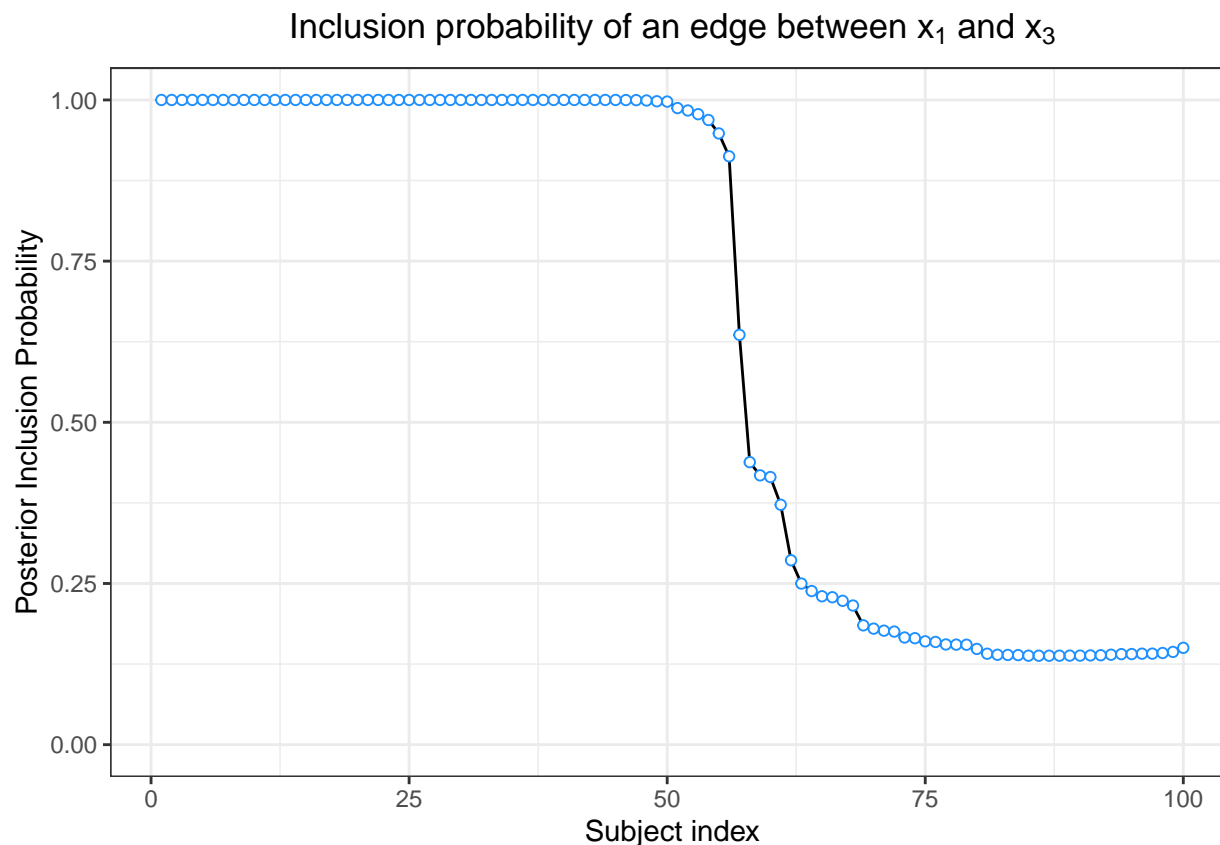
Posterior inclusion probabilities for individual 100



```
##gg_inclusionCurve  
gg_inclusionCurve(out, 1, 2)
```



```
gg_inclusionCurve(out, 1, 3, point_color = "dodgerblue")
```



```
# find sensitivity, specificity, and accuracy

# true positives
TP_neg <- sum(sapply(out$graphs[1:(n / 2)],
                    function(graph) sum(graph == 1 & true_graph_neg == 1)))
TP_pos <- sum(sapply(out$graphs[(n / 2 + 1):n],
                    function(graph) sum(graph == 1 & true_graph_pos == 1)))
TP <- TP_neg + TP_pos

# total positives
num_pos <- sum(true_graph_pos) * n / 2 + sum(true_graph_neg) * n / 2

# true negatives
TN_neg <- sum(sapply(out$graphs[1:(n / 2)],
                    function(graph) sum(graph == 0 & true_graph_neg == 0)))
TN_pos <- sum(sapply(out$graphs[(n / 2 + 1):n],
                    function(graph) sum(graph == 0 & true_graph_pos == 0)))
TN <- TN_neg + TN_pos

# total negatives
num_neg <- length(true_graph_pos) * n - num_pos

(sensitivity <- TP / num_pos)
```

```
## [1] 0.94
```

```
(specificity <- TN / num_neg)
```

```
## [1] 0.9686957
```

```
(accuracy <- (TN + TP) / (num_pos + num_neg))
```

```
## [1] 0.9664
```

To-do

- Topic 16 slides ideas
- 2 ideas from Mclust: logo and loading bar

```
# -----  
# /-----/ / / / / / /-----/ /-----/  
# / / / / / / / / / / /-----/ / / / / / /  
# / /-----/ / / \ / \ / \ /-----/ / / / / / /  
# \-----/ \-----/ \ / \ / \ /-----/ \-----/
```

- Change the weight calculation loop so that the inner loop range is constant, no matter `kde`
- Add Carbonetto-Stephens reference in documentation
- Check how to refer to σ^2 (regression coefficient variance) and σ_β^2 (slab variance)
- Create a vignette demonstrating usage on a simple simulated dataset
- Model details in return
- Change the `idmod` probs to logbase 10
- Remove CS argument
- Parallelization of the “main loop” over the predictors in `covdepGE_main.R`. This is complicated by the C++ code, however, two potential solutions are:
 - StackOverflow suggestion
 - RcppParallel

Bibliography

- (1) Dasgupta S., Ghosh P., Pati D., Mallick B. “An approximate Bayesian approach to covariate dependent graphical modeling.” 2021