How to use Fast Step Graph

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To install the last version of this package directly from GitHub uncomment and run:

```r
# library(devtools)
# use "quiet = FALSE" if you want to see the outputs of this command
# devtools::install_github("juancolonna/FastStepGraph", quiet = TRUE, force = TRUE)

# Then, load it:
library(FastStepGraph)
```

Simulate Gaussian Data with an Autoregressive (AR) Model:

```r
set.seed(1234567)
phi <- 0.4
p <- 50  # number of variables (dimension)
n <- 30  # number of samples

# Generate Data from a Gaussian distribution
data <- FastStepGraph::SigmaAR(n, p, phi)
X <− scale(data$X)  # standardizing variables
```

To fit the Omega matrix with `FastStepGraph()` function you have to know the optimal values of \( \alpha_f \) and \( \alpha_b \). If you don’t know these values, try to find them using cross-validation as follows:

```r
t0 <- Sys.time()  # INITIAL TIME
res <- FastStepGraph::cv.FastStepGraph(X, data_shuffle = TRUE)
difftime(Sys.time(), t0, units = "secs")
#> Time difference of 3.542877 secs
# print(res$alpha_f_opt)
# print(res$alpha_b_opt)
```

If your input variables are non-standardized (with zero mean and unit variance), we recommend that you set `data_scale=TRUE`. Subsequently, calculate the Omega matrix by calling the `FastStepGraph()` function passing the optimal parameters \( \alpha_f \) and \( \alpha_b \) found by cross-validation to fit the final model:

```r
t0 <- Sys.time()  # INITIAL TIME
G <- FastStepGraph::FastStepGraph(X, alpha_f = res$alpha_f_opt, alpha_b = res$alpha_b_opt)
difftime(Sys.time(), t0, units = "secs")
#> Time difference of 0.002274036 secs
# print(G$Omega)
```

You can also perform these two steps, the cross-validation to obtain the ideal parameters and return the fitted model, in a single step by setting the `return_model=TRUE` option as follows:

```r
t0 <- Sys.time()  # INITIAL TIME
res <- FastStepGraph::cv.FastStepGraph(X, return_model=TRUE, data_shuffle = TRUE)
```

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The arguments `n_folds = 5`, `alpha_f_min = 0.1`, `alpha_f_max = 0.9`, `n_alpha = 32` (size of the grid search) and `nei.max = 5`, have defaults values and can be omitted. Note that, `cv.FastStepGraph(X)` is not an exhaustive grid search over $\alpha_f$ and $\alpha_b$. This is a heuristic that tests only a few $\alpha_b$ values starting with the rule $\alpha_b = \alpha_f^2$. It is recommended to shuffle the rows of $X$ before running cross-validation. The default value is `data_shuffle = TRUE`, but if you want to disable row shuffle, set it to `data_shuffle = FALSE`.

To increase time performance, you can run `cv.FastStepGraph(X, parallel = TRUE)` in parallel. Before, you'll need to install and register a parallel backend. To run on a Linux system the `doParallel` dependency must be installed `install.packages("doParallel")`. These parallel packages will also require the following dependencies: `foreach`, `iterators` and `parallel`. Make sure you satisfy them. Then, call the method setting the parameter `parallel = TRUE`, as follows:

```r
t0 <- Sys.time()  # INITIAL TIME
# use 'n_cores = NULL' to set the maximum number of cores minus one on your machine
res <- FastStepGraph::cv.FastStepGraph(X, return_model=TRUE, parallel = TRUE, n_cores = 2)
difftime(Sys.time(), t0, units = "secs")
# print(res$alpha_f_opt)
# print(res$alpha_b_opt)
# print(res$Omega)
```

Remember, you can set the `n_cores` parameter to a value equal to the number of cores you have, but be careful as this may overload your system. Setting it to 1 disables parallel processing, and setting it to a number greater than the number of available cores does not improve efficiency.