HTSClusterPlus package instructions

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HTSClusterPlus version 0.99.1

HTSClusterPlus is a package to perform co-expression analyses of RNA-seq data. The package implements both Gaussian and Poisson mixtures model, performs parameter estimation using an expectation-maximization algorithm, and performs model selection using either the Integrated Completed Likelihood (ICL) criterion or the slope heuristics approach.

The full User's Guide for HTSClusterPlus may be found by installing the HTSClusterPlus package and loading it into an R session via library(HTSClusterPlus). In R for Windows, the User's Guide is then available from the drop-down menu called "Vignettes". In other operating systems, type

- > library(HTSClusterPlus)
- > HTSClusterUsersGuide()

at the R prompt to open the User's Guide in a pdf viewer.