NPBin

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We will analyze a subset of one of the sample dataset for illustration purposes.

```
library(npbin)
```

```
minimum_coverage <- 5 # minimum total coverage allowed
n_cores <- 47 # the number of cores to be used, can ONLY be 1 if run on Windows.

dt <- atac
colnames(dt)
#> [1] "chr" "location" "m"
#> [4] "xm" "winning.chip" "motif"
#> [7] "pval.mat.atSNP" "pval.pat.atSNP" "pval.rank.atSNP"
#> [10] "winnig.motif" "potential_TP" "potential_FP"

dt.ct <- data.table(dt)[m >= minimum_coverage, ]
```

for illustration purpose, keep only the 2000 points, remove this line will end up with analyzing the whole data set. It could be slow if only one core is used.

```
dt.ct <- dt.ct[1:2000,]
#> Error in eval(expr, envir, enclos): object 'dt.ct' not found
dt.ct[, p_hat:=xm / m]
#> Error in eval(expr, envir, enclos): object 'dt.ct' not found
n <- nrow(dt.ct)
#> Error in nrow(dt.ct): object 'dt.ct' not found

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n_breaks <- 11 # number of breaks
spline_order <- 4 # order of splines
breaks <- seq(0, 1, length.out = n_breaks)
pi_init <- hist(</pre>
```

```
dt.ct[, p_hat],
 breaks = seq(0, 1, length.out = n_breaks + spline_order - 3),
 plot = FALSE
)[["density"]] # initialized the weights using the histogram of p\_hat
\# Error in hist(dt.ct[, p_hat], breaks = seq(0, 1, length.out = n_breaks + : object 'dt.ct
estimate the overall model
overall_model_estimate <- emBinBspl(</pre>
  dt.ct[, xm],
 dt.ct[, m],
 breaks = breaks,
 k = spline_order,
 pi.init = pi_init,
 ncores = n_cores,
 err.max = 1e-3,
 iter.max = 200
#> Error in emBinBspl(dt.ct[, xm], dt.ct[, m], breaks = breaks, k = spline_order, : object
estimate the null model
null_model_estimate <- estNull(</pre>
 dt.ct[, xm],
 dt.ct[, m],
 overall_model_estimate,
 init = NULL,
 iter.max = 200,
 ncores = n_cores,
 ub = rep(log(1e4), 2),
  err.max = 1e-4
)
#> Error in estNull1(mod, pseq, ncores = ncores): object 'overall_model_estimate' not found
 fnp := null_model_estimate[["f"]]
][,
  fOnp := null_model_estimate[["f0"]]
][,
  locfdrnp := null_model_estimate[["locfdr"]]
][,
 fdrnp := locfdr2FDR(locfdrnp)
 ranknp := rank(locfdrnp, ties.method = "max")
#> Error in eval(expr, envir, enclos): object 'dt.ct' not found
```

```
names(null_model_estimate)
#> Error in eval(expr, envir, enclos): object 'null_model_estimate' not found
null model estimate$coef.null
#> Error in eval(expr, envir, enclos): object 'null_model_estimate' not found
Empirical Bayes test using p_hat
pct0 <- 0.45
empirical_bayes_beta_hat <- ebBeta(</pre>
 dt.ct[, xm],
 dt.ct[, m],
 dt.ct[, p hat],
 breaks = breaks,
 k = spline_order,
 pi.init = pi_init,
 pct0 = pct0,
 init = NULL,
 iter.max = 200,
 err.max = 1e-4,
 ncores = n_cores
)
\#> Error in emBspl(x, m, p, breaks = breaks, k = k, pi.init = pi.init, ncores = ncores, : of
 fhat := empirical_bayes_beta_hat[["f"]]
][,
 f0hat := empirical_bayes_beta_hat[["f0"]]
  locfdrhat := empirical_bayes_beta_hat[["locfdr"]]
][,
  fdrhat := locfdr2FDR(locfdrhat)
][,
 rankhat := rank(locfdrhat, ties.method = "max")
#> Error in eval(expr, envir, enclos): object 'dt.ct' not found
names(empirical_bayes_beta_hat)
#> Error in eval(expr, envir, enclos): object 'empirical_bayes_beta_hat' not found
null parameters of EBE
empirical_bayes_beta_hat[["coef.null"]]
#> Error in eval(expr, envir, enclos): object 'empirical_bayes_beta_hat' not found
```

```
p_binomial <- sapply(</pre>
  1:n,
 function(y) binom.test(dt.ct[y, xm], dt.ct[y, m])[["p.value"]]
)
\#> Error in lapply(X = X, FUN = FUN, ...): object 'n' not found
dt.ct[,
 pvbin := p_binomial
][,
  fdrbin := p.adjust(pvbin, method = "BH")
][,
 rankbin := rank(pvbin, ties.method = "max")
#> Error in eval(expr, envir, enclos): object 'dt.ct' not found
Evaluate the results using motifs
number of potential TP defined based on motif
dt.ct[, sum(potential TP)]
#> Error in eval(expr, envir, enclos): object 'dt.ct' not found
number of potential FP defined based on motif
dt.ct[, sum(potential_FP)]
#> Error in eval(expr, envir, enclos): object 'dt.ct' not found
find the number of TP and FP in top ranked SNPs
dt.ct[,
 tpnp := rank2nhit(ranknp ,potential_TP)
][,
  fpnp := rank2nhit(ranknp, potential_FP)
#> Error in eval(expr, envir, enclos): object 'dt.ct' not found
 tp_hat := rank2nhit(rankhat, potential_TP)
][,
  fp_hat := rank2nhit(rankhat, potential_FP)
#> Error in eval(expr, envir, enclos): object 'dt.ct' not found
dt.ct[,
 tpbin := rank2nhit(rankbin, potential_TP)
][,
 fpbin := rank2nhit(rankbin, potential_FP)
#> Error in eval(expr, envir, enclos): object 'dt.ct' not found
```

plot the accuracy measure as in the main paper. We presented a zoom-in version in the main paper to the top 20%, because usually there are not many ALI SNPs. Note that the default of the demo only select a subset of the data for illustration purposes. Thus the figure may not an exact replica of the one in the paper. To reproduce the results in the paper, please use the whole dataset

```
cbfpalette <- c(
   "#D55E00",
   "#0072B2",
   "#CC79A7",
   "#009E73",
   "#E69F00",
   "#56B4E9",
   "#F0E442"
)
plotidac <- c(' NPB','EBE','Binom')</pre>
```