# Package 'PING'

October 17, 2024

| Type Package  |  |  |  |  |
|---|--|--|--|--|
| Title Probabilistic inference for Nucleosome Positioning with<br>MNase-based or Sonicated Short-read Data                             |  |  |  |  |
| <b>Description</b> Probabilistic inference of ChIP-Seq using an empirical Bayes mixture model approach.                               |  |  |  |  |
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| <b>Depends</b> R(>= 3.5.0)  |  |  |  |  |
| Imports methods, PICS, graphics, grDevices, stats, Gviz, fda,<br>BSgenome, stats4, BiocGenerics, IRanges, GenomicRanges,<br>S4Vectors |  |  |  |  |
| Suggests parallel, ShortRead, rtracklayer   |  |  |  |  |
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| License Artistic-2.0  |  |  |  |  |
| biocViews Clustering, StatisticalMethod, Visualization, Sequencing  |  |  |  |  |
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#### Description

Post process Estimation of binding site positions obtained from PING. Refit mixture models with stronger prior in candidate regions contain potential problems, and then convert final result into dataframe.

#### Usage

```
postPING(
  ping,
  seg,
  rho2 = NULL,
  sigmaB2 = NULL,
  alpha2 = NULL,
  beta2 = NULL,
  min.dist = 100,
  paraEM = NULL,
  paraPrior = NULL,
  score = 0.05,
  dataType = "MNase",
  nCores = 1,
  makePlot = FALSE,
  FragmentLength = 100,
  mart = NULL,
  seg.boundary = NULL,
  DupBound = NULL,
  IP = NULL,
  datname = ""
)
```

#### Arguments

| ping                         | A pingList object containing estimation of nucleosome positions as returned<br>by the PING function.   |  |  |  |
|------------------------------|--|--|--|--|
| seg                          | An object of class segmentReadsList containing the results for all pre-processed regions as returned by segmentReads.  |  |  |  |
| rho2, sigmaB2, alpha2, beta2 |  |  |  |  |
|                              | Integer values, the parameters in the prior of mixture models to be re-fitted.   |  |  |  |
| min.dist                     | The minimum distance of two adjacent nucleosomes predicted from different candidate regions, smaller than that will be treated as duplicated predictions for the same nucleosomes. |  |  |  |
| paraEM                       | A list of parameters for the EM algorithm. The default parameters should be good enough for most usages.   |  |  |  |

| paraPrior  | A list of parameters for the prior distribution. The default parameters should be good enough for most usages.  |
|------------|---|
| score      | A numeric. The score threshold used when calling FilterPING.  |
| dataType   | A character that can be set to use selected default parameters for the algorithm.   |
| nCores     | An integer. The number of cores that should be used in parallel by the func-<br>tion.   |
| makePlot   | A logical. Plot a summary of the output.  |
|            | An integer. The length of XSET profile extension<br>ary, DupBound, datname<br>Plotting parameters and options.  |
| IP         | A GRanges object. The reads used in segmentation process.   |
| minK       | An integer. The minimum number of binding events per region. If the value is 0, the minimum number is automatically calculated.                               |
| maxK       | An integer. The maximum number of binding events per region. If the value is 0, the maximum number is automatically calculated.                               |
| tol        | A numeric. The tolerance for the EM algorithm.  |
| В          | An integer. The maximum number of iterations to be used.  |
| mSelect    | A character specifying the information criteria to be used when selecting the number of binding events. Default="AIC3"  |
| mergePeaks | A logical stating whether overlapping binding events should be picked.  |
| mapCorrect | A logical stating whether mappability profiles should be incorporated in the estimation, i.e: missing reads estimated.  |
| xi         | An integer. The average DNA fragment size.  |
| rho        | An integer. A variance parameter for the average DNA fragment size distribu-<br>tion.   |
| alpha      | An integer. First hyperparameter of the inverse Gamma distribution for sigma^2 in the PICS model  |
| beta       | An integer. Second hyperparameter of the inverse Gamma distribution for sigma <sup>2</sup> in the PING model  |
| lambda     | An integer. The lambda control Gaussian Markov Random Field prior on the distance of adjacent nucleosomes, we do not recommend user change the default value. |
| dMu        | An integer. Our best guess for the distance between two neighboring nucleo-<br>somes.   |

#### Value

A data.frame containing the estimated binding site positions

#### Note

Based on our experiemt on a few real data sets, we suggestion to use following values of parameters. For sonication data we use rho1=1.2; sigmaB2=6400; rho=15; alpha1=10; alpha2=98; beta2=200000. For MNase data we use rho1=3; sigmaB2=4900; rho=8; alpha1=20; alpha2=100; beta2=100000. The value of xi depends on specs of sample, since that affect the length of linker-DNA. For example, we use xi=160 for yeast and xi=200 for mouse.

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### See Also

PING, plotSummary

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