# rGMAP

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Type Package
Title Call hierarchical chromatin domains from HiC matrix by GMAP
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<b>Description</b> Call hierarchical chromatin domains from HiC contact matrix by Gaussian Mixture model And Proportion test, for more details see paper: Yu, W., He, B., & Tan, K. (2017). Identifying topologically associating domains and subdomains by Gaussian Mixture model And Proportion test. Nature Communications, 8, 535.
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LazyData TRUE
Imports data.table, ggplot2, mclust, EMD, caTools, Matrix, Rcpp (>= 0.12.5)
LinkingTo Rcpp
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data\_simu

generate simulated hic\_mat and true tads

#### **Description**

generate simulated hic\_mat and true tads

#### Usage

```
data_simu(stype = "poisson-dist", nratio = 2.5, mu0 = 20, resl = 1)
```

#### **Arguments**

stype One of four types of simulated data in the manuscript: poission-dist, poission-

dist-hier, nb-dist, nb-dist-hier; poission- or nb- indicates poission distribution or negative bionomial distribution -hier indicated subtads are generated nestly

nratio The effect size between intra- and inter domain, larger means higher intra-tad

contacts

mu0 The mean parameter, default 20 res1 Resolution, default set to 1

#### Value

A list includes following elements:

hic\_mat n by n contact matrix
hierTads True heirarchical domains

tads\_true True TADs

hic\_rao\_IMR90\_chr15

Normalized Hi-C data for IMR90, chr15 with resolution 10kb.

# Description

Normalized Hi-C data for IMR90, chr15 with resolution 10kb.

# Usage

```
hic_rao_IMR90_chr15
```

#### **Format**

A data table with 3 variables:

**n1** bin 1

**n2** bin 2

count normalized counts

plotdom 3

#### **Source**

Rao et al., Cell 2014, A 3D map of the human genome at kilobase resolution reveals principles of chromatin looping

plotdom

visualize hierarchical domains

#### **Description**

visualize hierarchical domains

#### Usage

```
plotdom(hic_dat, hiertads_gmap, start_bin, end_bin, cthr = 20, kb_resl = 10)
```

### **Arguments**

hiertads\_gmap

the hierarchical domains called by gmap

start\_bin the start bin of the genome end\_bin the end bin of the genome

cthr the count threshold for color, default 20

kb\_resl reslution of Hi-C data in kb

hic\_mat with 3 columns matrix or data.frame with columns: bin1, bin2, counts,

in which bin1 and bin2, from 1 to m, are the bin

rGMAP

Detect hierarchical choromotin domains by GMAP

#### **Description**

Detect hierarchical choromotin domains by GMAP

## Usage

```
rGMAP(hic_mat, resl = 10 * 10^3, logt = T, dom_order = 2, maxDistInBin = min(200, 2 * 10^6/resl), min_d = 25, max_d = 100, min_dp = 5, max_dp = 10, hthr = 0.95, t1thr = 0.5)
```

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#### **Arguments**

hic_mat	Either a 3 columns Hi-C contact matrix for a given chromosome, with each row corrsponding to the start bin, end bin and the contact number; or a n by n matrix, n is the number of bins for a given chromosom
resl	The resolution (bin size), default 10kb
logt	Do log-transformation or not, default TRUE
dom_order	Maximum level of hierarchical structures, default 2 (call TADs and subTADs)
maxDistInBin	Only consider contact whose distance is not greater than maxDistInBIn bins, default 200 bins (or 2Mb) $$
min_d	The minimum d (d: window size), default 25
max_d	The maximum d (d: window size), default 100
min_dp	The minmum dp (dp: lower bound of tad size), defalt 5
max_dp	The maximum dp (dp: lower bound of tad size), defalt 10. min_d, max_d, min_dp and max_dp should be specified in number of bins
hthr	The lower bound cutoff for posterior probability, default 0.95
t1thr	Lower bound for t1 for calling TAD, default $0.5$ quantile of test statistics of TADs, $0.9$ of subTADs

#### Value

#### A list includes following elements:

A data frame with columns start, end indicates the start and end coordinates of

each domain, respectively

hierTads A data frame with columns start, end, dom\_order, where dom\_order indicates

the hierarchical status of a domain, 1 indicates tads, 2 indicates subtads, and so

on

params A data frame gives the final parameters for calling TADs

# **Examples**

```
## On simulated data:
library(rGMAP)
simu_res = data_simu('poisson-dist-hier')
true_domains = simu_res$hierTads
simu_mat = simu_res$hie_mat
predicted_domains = rGMAP(simu_mat, resl = 1)$hierTads
true_domains
predicted_domains

## On an real data example
hic_rao_IMR90_chr15  # normalized Hi-C data for IMR90, chr15 with resolution 10kb
res = rGMAP(hic_rao_IMR90_chr15, resl = 10 * 1000, dom_order = 2)
names(res)
#quickly visualize some hierarchical domains
pp = plotdom(hic_rao_IMR90, res$hierTads, 6000, 7000, 30, 10)
pp$p2
```

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