# Package 'CpGFilter'

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Type Package
<b>Title</b> CpG Filtering Method Based on Intra-Class Correlation Coefficients
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<b>Description</b> Filter CpGs based on Intra-class Correlation Coefficients (ICCs) when replicates are available. ICCs are calculated by fitting linear mixed effects models to all samples including the un-replicated samples. Including the large number of un-replicated samples improves ICC estimates dramatically. The method accommodates any replicate design.
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R topics documented:
CpGFilterICC
Index

2 CpGFilterICC

CpGFilterICC

*CpG* filtering method based on intra-class correlation coefficients.

#### **Description**

Filter CpGs based on Intra-class Correlation Coefficients (ICCs). ICCs are calculated by fitting linear mixed effects models to all samples including the un-replicated samples. Including the large number of un-replicated samples improves ICC estimates dramatically. The method accommodates any replicate design.

#### Usage

```
CpGFilterICC(dat, rep.design, REML = FALSE, logit.transform = TRUE, verbose = TRUE)
```

#### **Arguments**

dat a matrix of CpG beta-values, row - CpG, column - sample

rep.design a vector indicating the replicate design, it could be factor, character or numeric

vectors. Example - c(1, 2, 3, 4, 4, 4, 5, 5) OR c('S1', 'S2', 'S2', 'S2', 'S1')

REML If TRUE, Restricted Maximum Likelihood (REML) method will be used; Oth-

erwise, Maximum Likelihood (ML) method will be used. Default is FALSE.

logit.transform

If TRUE, beta-value will be converted into M-value; Default is TRUE.

verbose If TRUE, print run information

#### Value

ICCs for all probes

#### Author(s)

Jun Chen

#### References

Chen J, Just A, et al. CpGFilter:Model-based CpG probe filtering with replicates for epigenome-wide association studies (2016). Bioinformatics, 32(3): 469–471

### Examples

```
require(CpGFilter)
# 10 samples replicated twice, 5 samples replicated four times.
rep.design <- c(1:100, 101:110, 101:110, 111:115, 111:115, 111:115, 111:115)
rho <- CpGFilterICC(matrix(rnorm(140*1000), 1000, 140), rep.design, logit=FALSE)</pre>
```

## **Index**

```
*Topic EWAS
CpGFilterICC, 2
*Topic ICC
CpGFilterICC, 2
*Topic LMM
CpGFilterICC, 2
```

CpGFilterICC, 2