Package 'RIPSeekerData'

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Title test data for RIPSeeker

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Author Yue Li

Maintainer Alejandro Reyes <reyes@embl.de>

Description

The RIP-seq data in BAM format are the test data for the package RIPSeeker. The data correspond to two RIP-seq experiments, namely PRC2 and CCNT1. Raw data from NCBI Gene Expression Omnibus under accession numbers GSE17064 for PRC2 and in-house for CCNT1.

biocViews ExperimentData, Genome, StemCell, SequencingData,

RIPSeqData, GEO

License GPL-2

Depends RIPSeeker

NeedsCompilation no

R topics documented:

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RIPSeekerData Alignment data in BAM files from RIP-seq experiments

Description

The RIP-seq data in BAM (binary version of the Sequence Alignment Map (SAM)) format are the test data for the package RIPSeeker. The data correspond to two RIP-seq experiments, namely PRC2 and CCNT1 as detailed below. The BAM data will be imported, processed, and converted into GappedAlignments using RIPSeeker function combineAlignGals.

Usage

RIPSeekerData

Format

BAM

Details

PRC2 RIP-seq dataset: The RIP-seq data from Zhao et al. (2010) for Ezh2 (a PRC2 unique subunit) in mouse embryonic stem cell (mESC) were downloaded from Gene Expression Omnibus (GEO) (GSE17064). Briefly, there are in total five datasets. Two datasets correspond to the non-specific and specific negative controls using the antibody IgG and mutant mESC depleted of Ezh2 (Ezh2 -/-) (MT), respectively. Only the specific negative control was used in our test. The two and one remaining datasets correspond to the libraries constructed from two biological replicates of the wild type mESC. Notably, the library construction and *strand-specific* sequencing generated sequences from the opposite strand of the PRC2-bound RNA (Zhao et al., 2010), consequently, each read was treated as if it were reverse complemented. After the quality control (QC) and alignments, the technical replicates were merged, resulting in three test files - RIP-biorep1, RIP-biorep2, and CTL with 1,022,474, 442,030, and 208,445 reads mapped to unique loci of the mouse reference genome (mm9 build).

CCNT1 RIP-seq dataset: The data for CCNT1 were generated from two RIP-seq experiments. The pilot experiment generated 775,582 and 773,785 strand-specific raw reads, and 5,853 and 4,556 uniquely mapped read remain after the stringent QC for the CCNT1 and GFP control RIP RNA libraries, respectively. Same as in the PRC2 data, the reads came from the second strand of the cDNA synthesis opposite to the original RNA strand. The non-strand-specific library from the second screen has deeper coverage with 1,647,641 and 2,369,271 raw reads, and 26,859 and 45,024 uniquely aligned reads under QC for CCNT1 and GFP, respectively. Since the two experiments were performed with slightly different protocols, we treated them as two separate biological replicates for the following analyses.

For RIP-seq analysis using the RIPSeeker package, both of the PRC2 and CCNT1 BAM datasets will be be imported, processed, and converted into GappedAlignments using RIPSeeker function combineAlignGals.

Source

Raw data from NCBI Gene Expression Omnibus under accession numbers GSE17064 for PRC2 and in-house for CCNT1.

References

Zhao, J., Ohsumi, T. K., Kung, J. T., Ogawa, Y., Grau, D. J., Sarma, K., Song, J. J., et al. (2010). Genome-wide Identification of Polycomb-Associated RNAs by RIP-seq. Molecular Cell, 40(6), 939D953. doi:10.1016/j.molcel.2010.12.011

The RIPSeeker manuscript has been submitted to NAR for review.

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