

TCGA.SampleSelection Documentation

Description: Retrieve TCGA data from Broad FireBrowse and perform sample selection on the basis of expression levels for specific genes of interest for analysis using GSEA tools.

Author: Anthony S. Castanza, Barbara A. Hill

Contact: genepattern.org/help

Summary: Queries cBioPortal for TCGA samples meeting criteria of mRNA expression z-scores relative to all samples (log RNA Seq V2 RSEM) greater than and less than user supplied thresholds. Outputs a GCT file containing TPM (Transcripts per Million) normalized RNA-seq quantifications suitable for ssGSEA, and a CLS file annotating samples as High or Low expression of the gene of interest.

Parameters:

Name	Description		
TCGA Collection*	TCGA study cohort (tumor types) to query for sample selection.		
Gene Symbol*	The HGNC Gene Symbol to use for classifying samples as high or low expression		
High Expression*	mRNA expression is greater than or equal to this threshold for standard deviations above the mean will be classified as "high" expression of the selected gene. (mRNA expression z-scores relative to all samples) Default = 1		
Low Expression*	mRNA expression is less than or equal to this threshold for standard deviations below the mean will be classified as "low" expression of the selected gene. (mRNA expression z-scores relative to all samples) Default = -1		
Output Type*	Type of RSEM quantifications to output: TPM (transcripts per million, within sample normalization, useful for ssGSEA) Raw counts (unnormalized counts, usable with DESeq2 or other DEG calculations).		
MSigDB Version*	MSigDB version to use for Gene Symbol lookup. This version should match the version of the gene sets intended for all downstream analysis. Supports MSigDB versions 7.1 and higher.		
	Note that the default is 'latest' which queries https://www.gsea-msigdb.org/gsea/msigdb to determine the current latest version of MSigDB. At the time this documentation was written, the latest version was 7.4. A current listing of all MSigDB versions can be found here:		
	https://software.broadinstitute.org/cancer/software/gsea/wiki/index.php/Release_Notes		

^{* =} required

Output File(s): GCT file containing gene expression values in the selected output type for samples that pass the specified thresholds. A CLS file indicating the sample group assignments (high or low expression).

Module Language: R

Source Repository: https://github.com/genepattern/TCGA.SampleSelection/tree/v0.13

Docker image: genepattern/tcga-sampleselection:beta4



Version	Date	Comment
0.14	2021-11-18	Update for cBioPortal assay name change
0.12	2021-07-29	Error handling fixes for invalid and unmappable genes
0.10	2021-05-04	Initial beta release.