

## **ExpressionFileCreator Documentation**

**Module name:** ExpressionFileCreator

**Description:** Creates a res or gct file from a set of Affymetrix CEL files

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**Summary**: Creates a res or gct file from a set of CEL files. The CEL files are given in one zip file. The CEL files contained in the zip file may also be zipped. Algorithms for converting from probe level data to expression level data are RMA, MAS5, and dChip. Note that the results obtained from running dChip and MAS5 differ slightly from the official implementations of these algorithms.

A clm file allows you to change the name of the samples in the expression matrix, reorder the sample names, and create a cls file. By default, sample names are taken from the CEL file names contained in the zip file. A clm file allows you to change the default sample names to your desired sample names. Additionally, the columns in the expression matrix are reordered so that they are in the same order as the sample names appear in the clm file. For example, the zip file my.cel.files.zip contains the files scan1.cel, scan2.cel, and scan3.cel. The clm file could contain the following text:

scan3.cel sample3 tumor scan1.cel sample1 tumor scan2.cel sample2 normal

The column names in the expression matrix would be: sample3, sample1, sample2.

#### References:

- 1. Affymetrix. Affymetrix Microarray Suite User Guide. Affymetrix, Santa Clara, CA, version 5 edition, 2001.
- 2. Rafael A. Irizarry, Bridget Hobbs, Francois Collin, Yasmin D. Beazer-Barclay, Kristen J. Antonellis, Uwe Scherf, and Terence P. Speed. Exploration, normalization, and summaries of high density oligonucleotide array probe level data. Biostatistics, 2003b. To appear.

### Parameters:

Name Description

input filename A zip file containing CEL files or zipped CEL files

method The method to use

quantile normalization (RMA only) Whether to normalize data using quantile

normalization

background correct (RMA only) Whether to background correct using RMA

background correction

compute present absent calls (MAS5 only) Whether to compute Present/Absent calls

normalization The normalization method to apply after expression values are

computed

reference.sample.name: (mean or median scaling only) The sample name to be used as

a reference for normalization. Leave blank to use median scan

as reference.

scale value (MAS5 only) Value at which all arrays will be scaled to

clm .filename tab-delimited text file containing one scan name, sample name,



## and class name per line The base name of the output file

output file

### **Return Value:**

1. gct or res file

2. cls file if clm.filename is supplied

# Platform dependencies:

Preprocess & Utility

Task type: CPU type: any OS: any R Language: