## DeMixT Log

Cell type-specific deconvolution of heterogeneous tumor samples with two or three components using expression data from RNAseq or microarray platforms.

## Update from version 1.2.1 to 1.2.3

## **Major Changes**

- 1. Added pi01 and pi02 as input values for users to initialize the proportion estimation.
- 2. Added nspikein as an input value in the DeMixT, DeMixT\_S1 and DeMixT\_GS functions to specify how many spike-in normal reference samples need to be generated; Setting nspikein at null as a default value, the number of spike-in normal reference samples equal the  $min(200, 0.3 \times My)$ , where My is the number of mixed samples; By setting nspikein equals 0, no spike-in normal reference will be generated; If the input value of data.N2 is not null, nspikein will be forced to be 0.
- 3. Added DeMixT\_GS function, new proposed gene selection method which applies profile likelihood, for proportion estimation.
- 4. Added simulate \_2comp function for users to simulate test data for 2-component de-convolution.
- 5. Added simulate 3comp function for users to simulate test data for 3-component de-convolution.
- 6. Added row names and column names for all output values.
- 7. Added gene.selection.method as an input value for DeMixT function. The default is 'GS'.
- 8. Added ngene. Profile. selected as an input value for DeMixT function. The default is NA.

## Bug fix

The filter step failed to set the value into input at in the previous version, and it has been resolved in the 1.3.0 version.

```
## filter out genes with constant value across all samples
  inputdata < ifelse(is.null(data.comp2),</pre>
              inputdata[apply(data.comp1, 1, function(x) length(unique(x)) > 1), ],
              inputdata[apply(data.comp1, 1, function(x) length(unique(x)) > 1) &
                         apply(data.comp2, 1, function(x) length(unique(x)) > 1), ])
## filter out genes with constant value across all samples
if (is.null(data.comp2)) {
        if (dim(inputdata)[1] == 1) {
            inputdata <- t(as.matrix(inputdata[apply(data.comp1,</pre>
                1, function(x) length(unique(x)) > 1), ]))
        }
        else {
            inputdata <- inputdata[apply(data.comp1, 1, function(x) length(unique(x)) >
                1), ]
        }
    }
    else {
        if (dim(inputdata)[1] == 1) {
            inputdata <- t(as.matrix(inputdata[apply(data.comp1,</pre>
                1, function(x) length(unique(x)) > 1) & apply(data.comp2,
```