

# Package: lmQCM (via r-universe)

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**Type** Package

**Title** An Algorithm for Gene Co-Expression Analysis

**Version** 0.2.4

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**Description** Implementation based on Zhang, Jie & Huang, Kun (2014)  
<[doi:10.4137/CIN.S14021](https://doi.org/10.4137/CIN.S14021)> Normalized lmQCM: An Algorithm for Detecting Weak Quasi-Cliques in Weighted Graph with Applications in Gene Co-Expression Module Discovery in Cancers. Cancer informatics, 13, CIN-S14021.

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**Encoding** UTF-8

**Depends** genefilter, Biobase, progress, stats, methods

**Suggests** devtools, roxygen2

**LazyData** true

**RoxygenNote** 7.2.1

**URL** <https://github.com/huangzhii/lmQCM/>

**BugReports** <https://github.com/huangzhii/lmQCM/issues/>

**Repository** <https://huangzhii.r-universe.dev>

**RemoteUrl** <https://github.com/huangzhii/lmqcm>

**RemoteRef** HEAD

**RemoteSha** 586da1915f2a28b9ab2387c72bb17e54a962b04c

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**fastFilter***fastFilter: Subroutine for filtering expression matrix***Description**

Author: Zhi Huang

**Usage**

```
fastFilter(
  RNA,
  lowest_percentile_mean = 0.2,
  lowest_percentile_variance = 0.2,
  var_func = "var"
)
```

**Arguments**

RNA	an expression matrix (rows: genes; columns: samples)
lowest_percentile_mean	a float value range 0-1
lowest_percentile_variance	a float value range 0-1
var_func	specify variance function

**Value**

An filtered expression matrix

**lmQCM***lmQCM: Main Routine for Gene Co-expression Analysis***Description**

Author: Zhi Huang

**Usage**

```
lmQCM(  
  data_in,  
  gamma = 0.55,  
  t = 1,  
  lambda = 1,  
  beta = 0.4,  
  minClusterSize = 10,  
  CCmethod = "pearson",  
  positiveCorrelation = F,  
  normalization = F  
)
```

**Arguments**

data_in	real-valued expression matrix with rownames indicating gene ID or gene symbol
gamma	gamma value (default = 0.55)
t	t value (default = 1)
lambda	lambda value (default = 1)
beta	beta value (default = 0.4)
minClusterSize	minimum length of cluster to retain (default = 10)
CCmethod	Methods for correlation coefficient calculation (default = "pearson"). Users can also pick "spearman".
positiveCorrelation	This determines if correlation matrix should convert to positive (with abs function) or not.
normalization	Determine if normalization is needed on massive correlation coefficient matrix.

**Value**

QCObject - An S4 Class with lmQCM results

**Examples**

```
library(lmQCM)  
library(BioBase)  
data(sample.ExpressionSet)  
data = assayData(sample.ExpressionSet)$exprs  
data = fastFilter(data, 0.2, 0.2)  
lmQCM(data)
```

**localMaximumQCM***localMaximumQCM: Subroutine for Creating Gene Clusters***Description**

Author: Zhi Huang

**Usage**

```
localMaximumQCM(cMatrix, gamma = 0.55, t = 1, lambda = 1)
```

**Arguments**

cMatrix	a correlation matrix
gamma	gamma value (default = 0.55)
t	t value (default = 1)
lambda	lambda value (default = 1)

**Value**

An unmerged clusters group 'C'

**merging\_lmQCM***merging\_lmQCM: Subroutine for Merging Gene Clusters***Description**

Author: Zhi Huang

**Usage**

```
merging_lmQCM(C, beta = 0.4, minClusterSize = 10)
```

**Arguments**

C	Resulting clusters
beta	beta value (default = 0.4)
minClusterSize	minimum length of cluster to retain (default = 10)

**Value**

mergedCluster - An merged clusters group

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