

# Package ‘RNASeqR’

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**Type** Package

**Title** RNASeqR: an R package for automated two-group RNA-Seq analysis workflow

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**Description** This R package is designed for case-control RNA-Seq analysis (two-group).

There are six steps: ``RNASeqRParam S4 Object Creation'', ``Environment Setup'', ``Quality Assessment'',

``Reads Alignment & Quantification'', ``Gene-level Differential Analyses'' and ``Functional Analyses''.

Each step corresponds to a function in this package. After running functions in order, a basic RNASeq analysis would be done easily.

**License** Artistic-2.0

**Encoding** UTF-8

**RoxygenNote** 6.1.1

**Depends** R(>= 3.5.0), ggplot2, pathview, edgeR, methods

**Imports** Rsamtools, tools, reticulate, ballgown, gridExtra, rafalib, FactoMineR, factoextra, corrplot, PerformanceAnalytics, reshape2, DESeq2, systemPipeR, systemPipeRdata, clusterProfiler, org.Hs.eg.db, org.Sc.sgd.db, stringr, pheatmap, grDevices, graphics, stats, utils, DOSE, Biostrings, parallel

**Suggests** knitr, png, grid, RNASeqRData

**VignetteBuilder** knitr

**SystemRequirements** RNASeqR only support Linux and macOS. Window is not supported. Python2 is highly recommended. If your machine is Python3, make sure '2to3' command is available.

**BugReports** <https://github.com/HowardChao/RNASeqR/issues>

**URL** <https://github.com/HowardChao/RNASeqR>

**NeedsCompilation** no

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All\_Steps\_Interface      RNASeqEnvironmentSet

## Description

Set up the environment for the following RNA-Seq workflow in R shell

This function do 4 things :

1. Create file directories.
2. Install necessary tools.

3. Export 'RNASeq\_bin/' to the R environment.

4. Check command of tools.

First it will create 'gene\_data/', 'RNASeq\_bin/', 'RNASeq\_results/', 'Rscript/', 'Rscript\_out/' directories.

Afterwards, 'Hisat2', 'Stringtie', 'Gffcompare' will be installed under 'RNASeq\_bin/Download/' and be unpacked under 'RNASeq\_bin/Unpacked/'.

'RNASeq\_bin/' will be added to the R environment and validity of tools will be checked.

Any ERROR occurs will be reported and the program will be terminated.

If you want to set up the environment for the following RNA-Seq workflow in background, please see RNASeqEnvironmentSet\_CMD() function.

## Usage

```
All_Steps_Interface(RNASeqRParam, which.trigger = "OUTSIDE",
INSIDE.path.prefix = NA, RNASeqQualityAssessment.RUN = TRUE,
RNASeqReadProcess.RUN = TRUE, RNASeqDifferentialAnalysis.RUN = TRUE,
RNASeqGoKegg.RUN = TRUE, OrgDb.species, go.level = 3, input.TYPE.ID,
KEGG.organism, check.s4.print = TRUE)
```

## Arguments

RNASeqRParam	S4 object instance of experiment-related parameters
which.trigger	Default value is OUTSIDE. User should not change this value.
INSIDE.path.prefix	Default value is NA. User should not change this value.
RNASeqQualityAssessment.RUN	Default value is TRUE. Set FALSE to skip "Quality Assessment" step.
RNASeqReadProcess.RUN	Default value is TRUE. Set FALSE to skip "RNASeq Read Process" step.
RNASeqDifferentialAnalysis.RUN	Default value is TRUE. Set FALSE to skip "RNASeq Differential Analysis" step.
RNASeqGoKegg.RUN	Default value is TRUE. Set FALSE to skip "RNASeq Go & Kegg" step.
OrgDb.species	the genome wide annotation packages of species on Bioconductor. Currently, there are 19 supported genome wide annotation packages of species.
go.level	the depth of acyclic graph in GO analysis
input.TYPE.ID	The gene name type in OrgDb.species annotation packahge.
KEGG.organism	the species that are supported for KEGG analysis. Currently, there are more than 5000 supported species genome. Check the valid species terms on <a href="https://www.genome.jp/kegg/catalog">https://www.genome.jp/kegg/catalog</a>
check.s4.print	Default TRUE. If TRUE, the result of checking RNASeqRParam will be reported in 'Rscript_out/Environment_Set.Rout'. If FALSE, the result of checking RNASeqRParam will not be in 'Rscript_out/Environment_Set.Rout'.

## Value

None

**Author(s)**

Kuan-Hao Chao

**Examples**

```
data(yeast)
## Not run:
RNASeqEnvironmentSet(RNASeqRParam = yeast)
## End(Not run)
```

*All\_Steps\_Interface\_CMD*

*All\_Steps\_Interface\_CMD*

**Description**

A functios to run all the steps with in one function. This function execute in the background:

1. Create file directories.
2. Install necessary tools.
3. Export 'RNASeq\_bin/' to the R environment.
4. Check command of tools.

First it will create 'gene\_data/', 'RNASeq\_bin/', 'RNASeq\_results/', 'Rscript/', 'Rscript\_out/' directories.

Afterwards, 'Hisat2', 'Stringtie', 'Gffcompare' will be installed under 'RNASeq\_bin/Download/' and be unpacked under 'RNASeq\_bin/Unpacked/'.

'RNASeq\_bin/' will be added to the R environment and validity of tools will be checked.

Any ERROR occurs will be reported and the program will be terminated.

If you want to set up the environment for the following RNA-Seq workflow in R shell, please see RNASeqEnvironmentSet() function.

**Usage**

```
All_Steps_Interface_CMD(RNASeqRParam, RNASeqQualityAssessment.RUN = TRUE,
RNASeqReadProcess.RUN = TRUE, RNASeqDifferentialAnalysis.RUN = TRUE,
RNASeqGoKegg.RUN = TRUE, OrgDb.species, go.level = 3, input.TYPE.ID,
KEGG.organism, run = TRUE, check.s4.print = TRUE)
```

**Arguments**

**RNASeqRParam** S4 object instance of experiment-related parameters

**RNASeqQualityAssessment.RUN**

Default value is TRUE. Set FALSE to skip "Quality Assessment" step.

**RNASeqReadProcess.RUN**

Default value is TRUE. Set FALSE to skip "RNASeq Read Process" step.

RNASeqDifferentialAnalysis.RUN	Default value is TRUE. Set FALSE to skip "RNASeq Differential Analysis" step.
RNASeqGoKegg.RUN	Default value is TRUE. Set FALSE to skip "RNASeq Go & Kegg" step.
OrgDb.species	the genome wide annotation packages of species on Bioconductor. Currently, there are 19 supported genome wide annotation packages of species.
go.level	the depth of acyclic graph in GO analysis
input.TYPE.ID	The gene name type in OrgDb.species annotation packahge.
KEGG.organism	the species that are supported for KEGG analysis. Currently, there are more than 5000 supported species genome. Check the valid species terms on <a href="https://www.genome.jp/kegg/catalog.html">https://www.genome.jp/kegg/catalog.html</a> .
run	Default value is TRUE. If TRUE, 'Rscript/Environment_Set.R' will be created and executed. The output log will be stored in 'Rscript_out/Environment_Set.Rout'. If False, 'Rscript/Environment_Set.R' will be created without executed.
check.s4.print	Default TRUE. If TRUE, the result of checking RNASeqRParam will be reported in 'Rscript_out/Environment_Set.Rout'. If FALSE, the result of checking RNASeqRParam will not be in 'Rscript_out/Environment_Set.Rout'.

**Value**

None

**Author(s)**

Kuan-Hao Chao

**Examples**

```
data(yeast)
## Not run:
RNASeqEnvironmentSet_CMD(yeast)
## End(Not run)
```

CheckToolAll

*CheckToolAll*

**Description**

Check whether 'Hisat2', 'Stringtie' and 'Gffcompare' are installed on the workstation

**Usage**

```
CheckToolAll(path.prefix, print = TRUE)
```

**Arguments**

path.prefix	path prefix of 'gene_data/', 'RNASeq_bin/', 'RNASeq_results/', 'Rscript/' and 'Rscript_out/' directories.
print	If TRUE, detailed information will be printed. If FALSE, detailed information will not be printed.

**Value**

None

**Examples**

```
data(yeast)
## Not run:
CheckToolAll(yeast@path.prefix,
             print=TRUE)
## End(Not run)
```

**RNASeqDifferentialAnalysis**  
*RNASeqDifferentialAnalysis*

**Description**

This function will run differential analysis on ballgown, DESeq2 and edgeR in background.  
This function do following things :

## 1. ballgown analysis

Raw reads are normalized into FPKM values

The main statistic test in ballgown is parametric F-test comparing nested linear models

## 2. DESeq2 analysis

Median of ratios normalization(MRN) is used in DESeq2 for raw reads count normalization.  
Sequencing depth and RNA composition is taken into consideration in this normalization method.

The main statistic test in DESeq2 is negative binomial distribution.

## 3. edgeR analysis

Raw reads are normalized by TMM and library size. (run calcNormFactors() to get a DGEList, and then run cpm() on that DGEList)

The main statistic test in edgeR is trimmed mean of M-values(TMM).

If you want to run differential analysis on ballgown, DESeq2, edgeR for the following RNA-Seq workflow in background, please see RNASeqDifferentialAnalysis() function.

**Usage**

```
RNASeqDifferentialAnalysis(RNASeqRParam, which.trigger = "OUTSIDE",
                           INSIDE.path.prefix = NA, Pre_DE.visualization = TRUE,
                           Post_DE.visualization = TRUE, ballgown.run = TRUE,
                           ballgown.pval = 0.05, ballgown.log2FC = 1, DESeq2.run = TRUE,
                           DESeq2.pval = 0.1, DESeq2.log2FC = 1, edgeR.run = TRUE,
                           edgeR.pval = 0.05, edgeR.log2FC = 1, check.s4.print = TRUE)
```

**Arguments**

**RNASeqRParam** S4 object instance of experiment-related parameters

**which.trigger** Default value is OUTSIDE. User should not change this value.

**INSIDE.path.prefix** Default value is NA. User should not change this value.

**Pre\_DE.visualization** Default TRUE. Whether to visualize pre-DE analysis results.

**Post\_DE.visualization** Default TRUE. Whether to visualize post-DE analysis results.

**ballgown.run** Default TRUE. Logical value whether to run ballgown differential analysis.

**ballgown.pval** Default 0.05. Set the threshold of ballgown p-value to filter out differential expressed gene.

**ballgown.log2FC** Default 1. Set the threshold of ballgown log2 fold change to filter out differential expressed gene.

**DESeq2.run** Default TRUE. Logical value whether to run DESeq2 differential analysis.

**DESeq2.pval** Default 0.05. Set the threshold of DESeq2 p-value to filter out differential expressed gene.

**DESeq2.log2FC** Default 1. Set the threshold of DESeq2 log2 fold change to filter out differential expressed gene.

**edgeR.run** Default TRUE. Logical value whether to run edgeR differential analysis.

**edgeR.pval** Default 0.05. Set the threshold of edgeR p-value to filter out differential expressed gene.

**edgeR.log2FC** Default 1. Set the threshold of edgeR log2 fold change to filter out differential expressed gene.

**check.s4.print** Default TRUE. If TRUE, the result of checking RNASeqRParam will be reported in 'Rscript\_out/Environment\_Set.Rout'. If FALSE, the result of checking RNASeqRParam will not be in 'Rscript\_out/Environment\_Set.Rout'.

**Value**

None

**Author(s)**

Kuan-Hao Chao

**Examples**

```
data(yeast)
## Not run:
RNASeqDifferentialAnalysis(RNASeqRParam = yeast)
## End(Not run)
```

RNASEqDifferentialAnalysis\_CMD  
*RNASEqDifferentialAnalysis\_CMD*

---

## Description

This function will run differential analysis on ballgown, DESeq2 and edgeR in R shell.  
 This function do following things :

### 1. ballgown analysis

Raw reads are normalized into FPKM values

The main statistic test in ballgown is paramatic F-test comparing nested linear models

### 2. DESeq2 analysis

Median of ratios normalization(MRN) is used in DESeq2 for raw reads count normalization.  
 Sequencing depth and RNA composition is taken into consideration in this normalization method.

The main statistic test in DESeq2 is negative binomial distribution.

### 3. edgeR analysis

Raw reads are normalized by TMM and library size. (run calcNormFactors() to get a DGEList, and then run cpm() on that DGEList)

The main statistic test in edgeR is trimmed mean of M-values(TMM).

If you want to run differential analysis on ballgown, DESeq2, edgeR for the following RNA-Seq workflow in R shell, please see RNASEqDifferentialAnalysis() function.

## Usage

```
RNASEqDifferentialAnalysis_CMD(RNASEqRParam, which.trigger = "OUTSIDE",
  INSIDE.path.prefix = NA, Pre_DE.visualization = TRUE,
  Post_DE.visualization = TRUE, ballgown.run = TRUE,
  ballgown.pval = 0.05, ballgown.log2FC = 1, DESeq2.run = TRUE,
  DESeq2.pval = 0.1, DESeq2.log2FC = 1, edgeR.run = TRUE,
  edgeR.pval = 0.05, edgeR.log2FC = 1, run = TRUE,
  check.s4.print = TRUE)
```

## Arguments

RNASEqRParam	S4 object instance of experiment-related parameters
which.trigger	Default value is OUTSIDE. User should not change this value.
INSIDE.path.prefix	Default value is NA. User should not change this value.
Pre_DE.visualization	Default TRUE. Whether to visualize pre-DE analysis results.
Post_DE.visualization	Default TRUE. Whether to visualize post-DE analysis results.
ballgown.run	Default TRUE. Logical value whether to run ballgown differential analysis.

ballgown.pval	Default 0.05. Set the threshold of ballgown p-value to filter out differential expressed gene.
ballgown.log2FC	Default 1. Set the threshold of ballgown log2 fold change to filter out differential expressed gene.
DESeq2.run	Default TRUE. Logical value whether to run DESeq2 differential analysis.
DESeq2.pval	Default 0.05. Set the threshold of DESeq2 p-value to filter out differential expressed gene.
DESeq2.log2FC	Default 1. Set the threshold of DESeq2 log2 fold change to filter out differential expressed gene.
edgeR.run	Default TRUE. Logical value whether to run edgeR differential analysis.
edgeR.pval	Default 0.05. Set the threshold of edgeR p-value to filter out differential expressed gene.
edgeR.log2FC	Default 1. Set the threshold of edgeR log2 fold change to filter out differential expressed gene.
run	Default value is TRUE. If TRUE, 'Rscript/Environment_Set.R' will be created and executed. The output log will be stored in 'Rscript_out/Environment_Set.Rout'. If False, 'Rscript/Environment_Set.R' will be created without executed.
check.s4.print	Default TRUE. If TRUE, the result of checking RNASEqRParam will be reported in 'Rscript_out/Environment_Set.Rout'. If FALSE, the result of checking RNASEqRParam will not be in 'Rscript_out/Environment_Set.Rout'.

**Value**

None

**Author(s)**

Kuan-Hao Chao

**Examples**

```
data(yeast)
## Not run:
RNASEqDifferentialAnalysis_CMD(RNASEqRParam = yeast)
## End(Not run)
```

RNASEqEnvironmentSet    *RNASEqEnvironmentSet***Description**

Set up the environment for the following RNA-Seq workflow in R shell  
 This function do 4 things :

1. Create file directories.

2. Install necessary tools.
3. Export 'RNASeq\_bin/' to the R environment.
4. Check command of tools.

First it will create 'gene\_data/', 'RNASeq\_bin/', 'RNASeq\_results/', 'Rscript/', 'Rscript\_out/' directories.

Afterwards, 'Hisat2', 'Stringtie', 'Gffcompare' will be installed under 'RNASeq\_bin/Download/' and be unpacked under 'RNASeq\_bin/Unpacked/'.

'RNASeq\_bin/' will be added to the R environment and validity of tools will be checked.

Any ERROR occurs will be reported and the program will be terminated.

If you want to set up the environment for the following RNA-Seq workflow in background, please see *RNASeqEnvironmentSet\_CMD()* function.

## **Usage**

```
RNASeqEnvironmentSet(RNASeqRParam, which.trigger = "OUTSIDE",
INSIDE.path.prefix = NA, install.hisat2 = TRUE,
install.STAR = TRUE, install.stringtie = TRUE,
install.gffcompare = TRUE, check.s4.print = TRUE)
```

## **Arguments**

RNASeqRParam	S4 object instance of experiment-related parameters
which.trigger	Default value is OUTSIDE. User should not change this value.
INSIDE.path.prefix	Default value is NA. User should not change this value.
install.hisat2	Whether to install 'HISAT2' in this function step. Default value is TRUE. Set FALSE to skip 'HISAT2' installation.
install.STAR	Whether to install 'STAR' in this function step. Default value is TRUE. Set FALSE to skip 'STAR' installation.
install.stringtie	Whether to install 'StringTie' in this function step. Default value is TRUE. Set FALSE to skip 'StringTie' installation.
install.gffcompare	Whether to install 'Gffcompare' in this function step. Default value is TRUE. Set FALSE to skip 'Gffcompare' installation.
check.s4.print	Default TRUE. If TRUE, the result of checking RNASeqRParam will be reported in 'Rscript_out/Environment_Set.Rout'. If FALSE, the result of checking RNASeqRParam will not be in 'Rscript_out/Environment_Set.Rout'.

## **Value**

None

## **Author(s)**

Kuan-Hao Chao

## Examples

```
data(yeast)
## Not run:
RNASEqEnvironmentSet(RNASEqRParam = yeast)
## End(Not run)
```

RNASEqEnvironmentSet\_CMD

*RNASEqEnvironmentSet\_CMD*

## Description

Set up the environment for the following RNA-Seq workflow in background.  
This function do 4 things :

1. Create file directories.
2. Install necessary tools.
3. Export 'RNASEq\_bin/' to the R environment.
4. Check command of tools.

First it will create 'gene\_data/', 'RNASEq\_bin/', 'RNASEq\_results/', 'Rscript/', 'Rscript\_out/' directories.

Afterwards, 'Hisat2', 'Stringtie', 'Gffcompare' will be installed under 'RNASEq\_bin/Download/' and be unpacked under 'RNASEq\_bin/Unpacked/'.

'RNASEq\_bin/' will be added to the R environment and validity of tools will be checked.

Any ERROR occurs will be reported and the program will be terminated.

If you want to set up the environment for the following RNA-Seq workflow in R shell, please see RNASEqEnvironmentSet() function.

## Usage

```
RNASEqEnvironmentSet_CMD(RNASEqRParam, install.hisat2 = TRUE,
install.STAR = TRUE, install.stringtie = TRUE,
install.gffcompare = TRUE, run = TRUE, check.s4.print = TRUE)
```

## Arguments

RNASEqRParam	S4 object instance of experiment-related parameters
install.hisat2	Whether to install 'HISAT2' in this function step. Default value is TRUE. Set FALSE to skip 'HISAT2' installation.
install.STAR	Whether to install 'STAR' in this function step. Default value is TRUE. Set FALSE to skip 'STAR' installation.
install.stringtie	Whether to install 'StringTie' in this function step. Default value is TRUE. Set FALSE to skip 'StringTie' installation.

install.gffcompare	Whether to install 'Gffcompare' in this function step. Default value is TRUE. SetFALSE to skip 'Gffcompare' installation.
run	Default value is TRUE. If TRUE, 'Rscript/Environment_Set.R' will be created and executed. The output log will be stored in 'Rscript_out/Environment_Set.Rout'. If False, 'Rscript/Environment_Set.R' will be created without executed.
check.s4.print	Default TRUE. If TRUE, the result of checking RNASEqRParam will be reported in 'Rscript_out/Environment_Set.Rout'. If FALSE, the result of checking RNASEqRParam will not be in 'Rscript_out/Environment_Set.Rout'.

**Value**

None

**Author(s)**

Kuan-Hao Chao

**Examples**

```
data(yeast)
## Not run:
RNASEqEnvironmentSet_CMD(yeast)
## End(Not run)
```

RNASEqGoKegg

*RNASEqGoKegg*

**Description**

Run Gene Ontology(GO) and Kyoto Encyclopedia of Genes and Genomes(KEGG) analysis in R shell.

This function do Gene Ontology(GO) and Kyoto Encyclopedia of Genes and Genomes(KEGG) analysis :

1. Gene Ontology(GO) :

- (a) Do GO function classification analysis.
- (b) Do GO function enrichment analysis.
- (c) Visualization : bar plot, dot plot etc.

2. Kyoto Encyclopedia of Genes and Genomes(KEGG) :

- (a) Do KEGG pathway enrichment analysis
- (b) Pathway visualization with pathview package. KEGG webpage pathway url will also be created

If you want to do GO functional analysis and KEGG pathway analysis for the following RNA-Seq workflow in background, please see RNASeqGoKegg\_CMD() function.

## Usage

```
RNASeqGoKegg(RNASeqRParam, which.trigger = "OUTSIDE",
  INSIDE.path.prefix = NA, OrgDb.species, go.level = 3, input.TYPE.ID,
  KEGG.organism, check.s4.print = TRUE)
```

## Arguments

RNASeqRParam	S4 object instance of experiment-related parameters
which.trigger	Default value is OUTSIDE. User should not change this value.
INSIDE.path.prefix	Default value is NA. User should not change this value.
OrgDb.species	the genome wide annotation packages of species on Bioconductor. Currently, there are 19 supported genome wide annotation packages of species.
go.level	the depth of acyclic graph in GO analysis
input.TYPE.ID	The gene name type in OrgDb.species annotation packahge.
KEGG.organism	the species that are supported for KEGG analysis. Currently, there are more than 5000 supported species genome. Check the valid species terms on <a href="https://www.genome.jp/kegg/catalog.html">https://www.genome.jp/kegg/catalog.html</a>
check.s4.print	Default TRUE. If TRUE, the result of checking RNASeqRParam will be reported in 'Rscript_out/Environment_Set.Rout'. If FALSE, the result of checking RNASeqRParam will not be in 'Rscript_out/Environment_Set.Rout'

## Value

None

## Author(s)

Kuan-Hao Chao

## Examples

```
data(yeast)
## Not run:
RNASeqGoKegg(RNASeqRParam = yeast,
  OrgDb.species = "org.Sc.sgd.db",
  go.level = 3,
  input.TYPE.ID = "GENENAME",
  KEGG.organism = "sce")
## End(Not run)
```

---

RNASEqGoKegg\_CMD      *RNASEqGoKegg\_CMD*

---

### Description

Run Gene Ontology(GO) and Kyoto Encyclopedia of Genes and Genomes(KEGG) analysis in background.

This function do Gene Ontology(GO) and Kyoto Encyclopedia of Genes and Genomes(KEGG) analysis :

1. Gene Ontology(GO) :

- (a) Do GO function classification analysis.
- (b) Do GO function enrichment analysis.
- (c) Visualization : bar plot, dot plot etc.

2. Kyoto Encyclopedia of Genes and Genomes(KEGG) :

- (a) Do KEGG pathway enrichment analysis
- (b) Pathway visualization with pathview package. KEGG webpage pathway url will also be created

If you want to do GO functional analysis and KEGG pathway analysis for the following RNA-Seq workflow in R shell, please see RNASEqGoKegg() function.

### Usage

```
RNASEqGoKegg_CMD(RNASEqRParam, OrgDb.species, go.level = 3,
                   input.TYPE.ID, KEGG.organism, run = TRUE, check.s4.print = TRUE)
```

### Arguments

RNASEqRParam	S4 object instance of experiment-related parameters
OrgDb.species	the genome wide annotation packages of species on Bioconductor. Currently, there are 19 supported genome wide annotation packages of species.
go.level	the depth of acyclic graph in GO analysis
input.TYPE.ID	The gene name type in OrgDb.species annotation packahge.
KEGG.organism	the species that are supported for KEGG analysis. Currently, there are more than 5000 supported species genome. Check the valid species terms on <a href="https://www.genome.jp/kegg/catalog">https://www.genome.jp/kegg/catalog</a>
run	Default value is TRUE. If TRUE, 'Rscript/Environment_Set.R' will be created and executed. The output log will be stored in 'Rscript_out/Environment_Set.Rout'. If False, 'Rscript/Environment_Set.R' will be created without executed.
check.s4.print	Default TRUE. If TRUE, the result of checking RNASEqRParam will be reported in 'Rscript_out/Environment_Set.Rout'. If FALSE, the result of checking RNASEqRParam will not be in 'Rscript_out/Environment_Set.Rout'

**Value**

None

**Author(s)**

Kuan-Hao Chao

**Examples**

```
data(yeast)
## Not run:
RNASeqGoKegg_CMD(RNASeqRParam = yeast,
                   OrgDb.species = "org.Sc.sgd.db",
                   go.level = 3,
                   input.TYPE.ID = "GENENAME",
                   KEGG.organism = "sce")

## End(Not run)
```

**Description**

Assess the quality of '.fastq.gz' files for RNA-Seq workflow in R shell. This step is optional in the whole RNA-Seq workflow.

This function reports the quality assessment result in packages systemPipeR For systemPipeR, 'RNASeq\_results/QA\_results/Rqc/systemPipeR/fastqReport.pdf' will be created.

If you want to assess the quality of '.fastq.gz' files for the following RNA-Seq workflow in background, please see RNASeqQualityAssessment\_CMD() function.

**Usage**

```
RNASeqQualityAssessment(RNASeqRParam, which.trigger = "OUTSIDE",
                        INSIDE.path.prefix = NA, check.s4.print = TRUE)
```

**Arguments**

RNASeqRParam	S4 object instance of experiment-related parameters
which.trigger	Default value is OUTSIDE. User should not change this value.
INSIDE.path.prefix	Default value is NA. User should not change this value.
check.s4.print	Default TRUE. If TRUE, the result of checking RNASeqRParam will be reported in 'Rscript_out/Environment_Set.Rout'. If FALSE, the result of checking RNASeqRParam will not be in 'Rscript_out/Environment_Set.Rout'

**Value**

None

**Author(s)**

Kuan-Hao Chao

**Examples**

```
data(yeast)
## Not run:
RNASeqQualityAssessment(RNASeqRParam = yeast)
## End(Not run)
```

**RNASeqQualityAssessment\_CMD**  
*RNASeqQualityAssessment\_CMD*

**Description**

Assess the quality of '.fastq.gz' files for RNA-Seq workflow in background. This step is optional in the whole RNA-Seq workflow.

This function reports the quality assessment result in packages `systemPipeR`. For `systemPipeR`, '`RNASeq_results/QA_results/Rqc/systemPipeR/fastqReport.pdf`' will be created.

If you want to assess the quality of '.fastq.gz' files for the following RNA-Seq workflow in R shell, please see `RNASeqQualityAssessment()` function.

**Usage**

```
RNASeqQualityAssessment_CMD(RNASeqRParam, run = TRUE,
                             check.s4.print = TRUE)
```

**Arguments**

<code>RNASeqRParam</code>	S4 object instance of experiment-related parameters
<code>run</code>	Default value is TRUE. If TRUE, ' <code>Rscript/Environment_Set.R</code> ' will be created and executed. The output log will be stored in ' <code>Rscript_out/Environment_Set.Rout</code> '. If False, ' <code>Rscript/Environment_Set.R</code> ' will be created without executed.
<code>check.s4.print</code>	Default TRUE. If TRUE, the result of checking <code>RNASeqRParam</code> will be reported in ' <code>Rscript_out/Environment_Set.Rout</code> '. If FALSE, the result of checking <code>RNASeqRParam</code> will not be in ' <code>Rscript_out/Environment_Set.Rout</code> '

**Value**

None

**Author(s)**

Kuan-Hao Chao

**Examples**

```
data(yeast)
## Not run:
RNASeqQualityAssessment_CMD(RNASeqRParam = yeast)
## End(Not run)
```

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RNASEqR	<i>RNASEqR-package</i>
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## Description

**RNASEqR-package**

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RNASEqReadProcess	<i>RNASEqReadProcess</i>
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---

## Description

Process raw reads for RNA-Seq workflow in R shell  
This function do 5 things :

1. 'Hisat2' : aligns raw reads to reference genome. If `indices.optional` in `RNASEqRParam` is FALSE, Hisat2 indices will be created.
2. 'Rsamtools': converts '.sam' files to '.bam' files.
3. 'Stringtie': assembles alignments into transcript.
4. 'Gffcompare': examines how transcripts compare with the reference annotation.
5. 'Stringtie': creates input files for ballgown, edgeR and DESeq2.
6. raw reads count: create raw reads count for DESeq2 and edgeR

Before running this function, `RNASEqEnvironmentSet_CMD()` or `RNASEqEnvironmentSet()` must be executed successfully. If you want to process raw reads for the following RNA-Seq workflow in background, please see `RNASEqReadProcess_CMD()` function.

## Usage

```
RNASEqReadProcess(RNASEqRParam, which.trigger = "OUTSIDE",
  INSIDE.path.prefix = NA, SAMtools.or.RSAMtools = "RSAMtools",
  Hisat2.Index.run = TRUE, Hisat2.Index.num.parallel.threads = "1",
  Hisat2.Index.large.index = FALSE,
  Hisat2.Index.local.ftab.chars = "6",
  Hisat2.Index.local.off.rate = "3", Hisat2.Index.ftab.chars = "10",
  Hisat2.Index.off.rate = "4", Hisat2.Alignment.run = TRUE,
  Hisat2.Alignment.num.parallel.threads = "1",
  Hisat2.Alignment.skip = "0", Hisat2.Alignment.trim5 = "0",
  Hisat2.Alignment.trim3 = "0",
  Hisat2.Alignment.n.ceil.1.function.type = "L",
  Hisat2.Alignment.n.ceil.2.constant.term = "0",
```

```

Hisat2.Alignment.n.ceil.3.coefficient = "0.15",
Hisat2.Alignment.mp.MX = "6", Hisat2.Alignment.mp.MN = "2",
Hisat2.Alignment.sp.MX = "2", Hisat2.Alignment.sp.MN = "1",
Hisat2.Alignment.np = "1", Hisat2.Alignment.rdg.1 = "5",
Hisat2.Alignment.rdg.2 = "3", Hisat2.Alignment.rfg.1 = "5",
Hisat2.Alignment.rfg.2 = "3",
Hisat2.Alignment.score.min.1.function.type = "L",
Hisat2.Alignment.score.min.2.constant.term = "0",
Hisat2.Alignment.score.min.3.coefficient = "-0.2",
Hisat2.Alignment.pen.cansplice = "0",
Hisat2.Alignment.penc.noncansplice = "12",
Hisat2.Alignment.pen.canintronlen.1.function.type = "G",
Hisat2.Alignment.pen.canintronlen.2.constant.term = "-8",
Hisat2.Alignment.pen.canintronlen.3.coefficient = "1",
Hisat2.Alignment.pen.noncanintronlen.1.function.type = "G",
Hisat2.Alignment.pen.noncanintronlen.2.constant.term = "-8",
Hisat2.Alignment.pen.noncanintronlen.3.coefficient = "1",
Hisat2.Alignment.min.intronlen = "20",
Hisat2.Alignment.max.intronlen = "500000",
Hisat2.Alignment.rna.strandness = "None", Hisat2.Alignment.k = "5",
Hisat2.Alignment.max.seeds = "5", Hisat2.Alignment.secondary = FALSE,
Hisat2.Alignment.minins = "0", Hisat2.Alignment.maxins = "500",
Hisat2.Alignment.seed = "0", STAR.Index.num.parallel.threads = "1",
STAR.Index.sjdbOverhang.Read.length = "100",
STAR.Index.genomeSAindexNbases = "14",
STAR.Index.genomeChrBinNbites = "18",
STAR.Index.genomeSAsparseD = "1", STAR.Alignment.run = FALSE,
STAR.Alignment.num.parallel.threads = "1",
STAR.Alignment.genomeLoad = "NoSharedMemory",
STAR.Alignment.readMapNumber = "-1",
STAR.Alignment.clip3pNbases = "0", STAR.Alignment.clip5pNbases = "0",
STAR.Alignment.clip3pAdapterSeq = "-",
STAR.Alignment.clip3pAdapterMMp = "0.1",
STAR.Alignment.clip3pAfterAdapterNbases = "0",
STAR.Alignment.limitGenomeGenerateRAM = "31000000000",
STAR.Alignment.limitIObufferSize = "150000000",
STAR.Alignment.limitOutSAMoneReadBytes = "100000",
STAR.Alignment.limitOutSJoneRead = "1000",
STAR.Alignment.limitOutSJcollapsed = "1000000",
STAR.Alignment.limitBAMsortRAM = "0",
STAR.Alignment.outReadsUnmapped = "None",
STAR.Alignment.outQSconversionAdd = "0",
STAR.Alignment.outSAMprimaryFlag = "OneBestScore",
STAR.Alignment.outSAMmapqUnique = "255",
STAR.Alignment.scoreGap = "0", STAR.Alignment.scoreGapNoncan = "-8",
STAR.Alignment.scoreGapGCAG = "-4",
STAR.Alignment.scoreGapATAC = "-8",
STAR.Alignment.scoreGenomicLengthLog2scale = "-0.25",
STAR.Alignment.scoreDelOpen = "-2",
STAR.Alignment.scoreDelBase = "-2",
STAR.Alignment.scoreInsOpen = "-2",
STAR.Alignment.scoreInsBase = "-2",

```

```

STAR.Alignment.scoreStitchSJshift = "1",
STAR.Alignment.seedSearchStartLmax = "50",
STAR.Alignment.seedSearchStartLmaxOverLread = "1.0",
STAR.Alignment.seedSearchLmax = "0",
STAR.Alignment.seedMultimapNmax = "10000",
STAR.Alignment.seedPerReadNmax = "1000",
STAR.Alignment.seedPerWindowNmax = "50",
STAR.Alignment.seedNoneLociPerWindow = "10",
STAR.Alignment.alignIntronMin = "21",
STAR.Alignment.alignIntronMax = "0",
STAR.Alignment.alignMatesGapMax = "0",
STAR.Alignment.alignSJoverhangMin = "5",
STAR.Alignment.alignSJDBoverhangMin = "3",
STAR.Alignment.alignSplicedMateMapLmin = "0",
STAR.Alignment.alignSplicedMateMapLminOverLmate = "0.66",
STAR.Alignment.alignWindowsPerReadNmax = "10000",
STAR.Alignment.alignTranscriptsPerWindowNmax = "100",
STAR.Alignment.alignTranscriptsPerReadNmax = "10000",
STAR.Alignment.alignEndsType = "Local",
STAR.Alignment.winAnchorMultimapNmax = "50",
STAR.Alignment.winBinNbits = "16",
STAR.Alignment.winAnchorDistNbins = "9",
STAR.Alignment.winFlankNbins = "4", Rsamtools.Bam.run = TRUE,
Samtools.Bam.num.parallel.threads = "1", Rsamtools.nCores = "1",
Stringtie.Assembly.run = TRUE,
Stringtie.Assembly.num.parallel.threads = "1",
Stringtie.Assembly.f = "0.1", Stringtie.Assembly.m = "200",
Stringtie.Assembly.c = "2.5", Stringtie.Assembly.g = "50",
Stringtie.Assembly.M = "0.95", Stringtie.Merge.Trans.run = TRUE,
Stringtie.Merge.num.parallel.threads = "1",
Gffcompare.Ref.Sample.run = TRUE, Stringtie.Ballgown.run = TRUE,
Stringtie.2.Ballgown.num.parallel.threads = "1",
PreDECCountTable.run = TRUE, check.s4.print = TRUE)

```

## Arguments

- RNASeqRParam** S4 object instance of experiment-related parameters
- which.trigger** Default value is OUTSIDE. User should not change this value.
- INSIDE.path.prefix**  
Default value is NA. User should not change this value.
- SAMtools.or.Rsamtools**  
Default value is Rsamtools. User can set to SAMtools to use command-line-based 'samtools' instead.
- Hisat2.Index.run**  
Whether to run 'HISAT2 index' step in this function step. Default value is TRUE.  
Set FALSE to skip 'HISAT2 index' step.
- Hisat2.Index.num.parallel.threads**  
Specify the number of processing threads (CPUs) to use for Hisat2 index step.  
The default is "1"
- Hisat2.Index.large.index**  
Hisat2 index terminal '--large-index' option. Default value is FALSE

```

Hisat2.Index.local.ftab.chars
    Hisat2 index terminal '-t/-ftabchars' option. Default value is "6"
Hisat2.Index.local.off.rate
    Hisat2 index terminal '--localoffrate' option. Default value is "3"
Hisat2.Index.ftab.chars
    Hisat2 index terminal '--localftabchars' option. Default value is "10"
Hisat2.Index.off.rate
    Hisat2 index terminal '--offrate' option. Default value is "4"
Hisat2.Alignment.run
    Whether to run 'HISAT2 alignment' step in this function step. Default value is TRUE. Set FALSE to skip 'HISAT2 alignment' step.
Hisat2.Alignment.num.parallel.threads
    Specify the number of processing threads (CPUs) to use for Hisat2 alignment step. The default is "1"
Hisat2.Alignment.skip
    Hisat2 alignment terminal '-s/-skip' option. Default value is "0"
Hisat2.Alignment.trim5
    Hisat2 alignment terminal '-5/-trim5' option. Default value is "0"
Hisat2.Alignment.trim3
    Hisat2 alignment terminal '-3/-trim3' option. Default value is "0"
Hisat2.Alignment.n.ceil.1.function.type
    Hisat2 alignment terminal '--n-ceil' option. Default value is "L"
Hisat2.Alignment.n.ceil.2.constant.term
    Hisat2 alignment terminal '--n-ceil' option. Default value is "0"
Hisat2.Alignment.n.ceil.3.coefficient
    Hisat2 alignment terminal '--n-ceil' option. Default value is "0.15"
Hisat2.Alignment.mp.MX
    Hisat2 alignment terminal '--mp MX' option. Default value is "6"
Hisat2.Alignment.mp.MN
    Hisat2 alignment terminal '--mp MN' option. Default value is "2"
Hisat2.Alignment.sp.MX
    Hisat2 alignment terminal '--sp MX' option. Default value is "2"
Hisat2.Alignment.sp.MN
    Hisat2 alignment terminal '--sp MN' option. Default value is "1"
Hisat2.Alignment.np
    Hisat2 alignment terminal '--np' option. Default value is "1"
Hisat2.Alignment.rdg.1
    Hisat2 alignment terminal '--rdg' first option. Default value is "5"
Hisat2.Alignment.rdg.2
    Hisat2 alignment terminal '--rdg' first option. Default value is "3"
Hisat2.Alignment.rfg.1
    Hisat2 alignment terminal '--rfg' first option. Default value is "5"
Hisat2.Alignment.rfg.2
    Hisat2 alignment terminal '--rfg' first option. Default value is "3"
Hisat2.Alignment.score.min.1.function.type
    Hisat2 alignment terminal '--rdg' first option. Default value is "L"
Hisat2.Alignment.score.min.2.constant.term
    Hisat2 alignment terminal '--rdg' first option. Default value is "0"

```

```
Hisat2.Alignment.score.min.3.coefficient
    Hisat2 alignment terminal '-rdg' first option. Default value is "-0.2"
Hisat2.Alignment.pen.cansplice
    Hisat2 alignment terminal '-pen-cansplice' first option. Default value is "-0"
Hisat2.Alignment.penc.noncansplice
    Hisat2 alignment terminal '-pen-noncansplice' option. Default value is "12"
Hisat2.Alignment.pen.canintronlen.1.function.type
    Hisat2 alignment terminal '-pen-canintronlen' first option. Default value is "G"
Hisat2.Alignment.pen.canintronlen.2.constant.term
    Hisat2 alignment terminal '-pen-canintronlen' second option. Default value is
    "-8"
Hisat2.Alignment.pen.canintronlen.3.coefficient
    Hisat2 alignment terminal '-pen-canintronlen' third option. Default value is "1"
Hisat2.Alignment.pen.noncanintronlen.1.function.type
    Hisat2 alignment terminal '-pen-noncanintronlen' first option. Default value is
    "G"
Hisat2.Alignment.pen.noncanintronlen.2.constant.term
    Hisat2 alignment terminal '-pen-noncanintronlen' second option. Default value
    is "-8"
Hisat2.Alignment.pen.noncanintronlen.3.coefficient
    Hisat2 alignment terminal '-pen-noncanintronlen' third option. Default value is
    "1"
Hisat2.Alignment.min.intronlen
    Hisat2 alignment terminal '-min-intronlen' option. Default value is "20"
Hisat2.Alignment.max.intronlen
    Hisat2 alignment terminal '-max-intronlen' option. Default value is "20"
Hisat2.Alignment.rna.strandness
    Hisat2 alignment terminal '-rna-strandness' option. Default value is "None"
Hisat2.Alignment.k
    Hisat2 alignment terminal '-k' option. Default value is "5"
Hisat2.Alignment.max.seeds
    Hisat2 alignment terminal '-max-seeds' option. Default value is "5"
Hisat2.Alignment.secondary
    Hisat2 alignment terminal '-secondary' option. Default value is "FALSE"
Hisat2.Alignment.minins
    Hisat2 alignment terminal '-I/-minins' option. Default value is "0"
Hisat2.Alignment.maxins
    Hisat2 alignment terminal '-X/-maxins' option. Default value is "500"
Hisat2.Alignment.seed
    Hisat2 alignment terminal '-X/-maxins' option. Default value is "0"
STAR.Index.num.parallel.threads
    Specify the number of processing threads (CPUs) to use for STAR index step.
    The default is "1"
STAR.Index.sjdbOverhang.Read.length
    STAR index terminal '-sjdbOverhang' option. Default value is "100"
STAR.Index.genomeSAindexNbases
    STAR index terminal '-genomeSAindexNbases' option. Default value is "14"
STAR.Index.genomeChrBinNbits
    STAR index terminal '-genomeChrBinNbits' option. Default value is "18"
```

```

STAR.Index.genomeSAsparseD
    STAR index terminal '-genomeSAsparseD' option. Default value is "1"
STAR.Alignment.run
    Whether to run 'STAR index' step in this function step. Default value is FALSE.
    Set TRUE to run STAR alignment step. (need to set Hisat2.Index.run to FALSE)
STAR.Alignment.num.parallel.threads
    Specify the number of processing threads (CPUs) to use for STAR alignment
    step. The default is "1"
STAR.Alignment.genomeLoad
    STAR alignment terminal '-genomeLoad' option. Default value is "NoSharedMemory"
STAR.Alignment.readMapNumber
    STAR alignment terminal '-readMapNumber' option. Default value is "-1"
STAR.Alignment.clip3pNbases
    STAR alignment terminal '-clip3pNbases' option. Default value is "0"
STAR.Alignment.clip5pNbases
    STAR alignment terminal '-clip5pNbases' option. Default value is "0"
STAR.Alignment.clip3pAdapterSeq
    STAR alignment terminal '-clip3pAdapterSeq' option. Default value is "-"
STAR.Alignment.clip3pAdapterMMP
    STAR alignment terminal '-clip3pAdapterMMP' option. Default value is "0.1"
STAR.Alignment.clip3pAfterAdapterNbases
    STAR alignment terminal '-clip3pAfterAdapterNbases' option. Default value
    is "0"
STAR.Alignment.limitGenomeGenerateRAM
    STAR alignment terminal '-limitGenomeGenerateRAM' option. Default value
    is "31000000000"
STAR.Alignment.limitIObufferSize
    STAR alignment terminal '-limitIObufferSize' option. Default value is "150000000"
STAR.Alignment.limitOutSAMoneReadBytes
    STAR alignment terminal '-limitOutSAMoneReadBytes' option. Default value
    is "100000"
STAR.Alignment.limitOutSJoneRead
    STAR alignment terminal '-limitOutSJoneRead' option. Default value is "1000"
STAR.Alignment.limitOutSJcollapsed
    STAR alignment terminal '-limitOutSJcollapsed' option. Default value is "1000000"
STAR.Alignment.limitBAMsortRAM
    STAR alignment terminal '-limitBAMsortRAM' option. Default value is "0"
STAR.Alignment.outReadsUnmapped
    STAR alignment terminal '-outReadsUnmapped' option. Default value is "None"
STAR.Alignment.outQSconversionAdd
    STAR alignment terminal '-outQSconversionAdd' option. Default value is "0"
STAR.Alignment.outSAMprimaryFlag
    STAR alignment terminal '-outSAMprimaryFlag' option. Default value is "OneBestScore"
STAR.Alignment.outSAMmapqUnique
    STAR alignment terminal '-outSAMmapqUnique' option. Default value is "255"
STAR.Alignment.scoreGap
    STAR alignment terminal '-scoreGap' option. Default value is "0"
STAR.Alignment.scoreGapNoncan
    STAR alignment terminal '-scoreGapNoncan' option. Default value is "-8"

```

```
STAR.Alignment.scoreGapGCAG
    STAR alignment terminal '--scoreGapGCAG' option. Default value is "-4"
STAR.Alignment.scoreGapATAC
    STAR alignment terminal '--scoreGapATAC' option. Default value is "-8"
STAR.Alignment.scoreGenomicLengthLog2scale
    STAR alignment terminal '--scoreGenomicLengthLog2scale' option. Default
    value is "-0.25"
STAR.Alignment.scoreDelOpen
    STAR alignment terminal '--scoreDelOpen' option. Default value is "-2"
STAR.Alignment.scoreDelBase
    STAR alignment terminal '--scoreDelBase' option. Default value is "-2"
STAR.Alignment.scoreInsOpen
    STAR alignment terminal '--scoreInsOpen' option. Default value is "-2"
STAR.Alignment.scoreInsBase
    STAR alignment terminal '--scoreInsBase' option. Default value is "-2"
STAR.Alignment.scoreStitchSJshift
    STAR alignment terminal '--scoreStitchSJshift' option. Default value is "1"
STAR.Alignment.seedSearchStartLmax
    STAR alignment terminal '--seedSearchStartLmax' option. Default value is "50"
STAR.Alignment.seedSearchStartLmaxOverLread
    STAR alignment terminal '--seedSearchStartLmaxOverLread' option. Default
    value is "1.0"
STAR.Alignment.seedSearchLmax
    STAR alignment terminal '--seedSearchLmax' option. Default value is "0"
STAR.Alignment.seedMultimapNmax
    STAR alignment terminal '--seedMultimapNmax' option. Default value is "10000"
STAR.Alignment.seedPerReadNmax
    STAR alignment terminal '--seedPerReadNmax' option. Default value is "1000"
STAR.Alignment.seedPerWindowNmax
    STAR alignment terminal '--seedPerWindowNmax' option. Default value is
    "50"
STAR.Alignment.seedNoneLociPerWindow
    STAR alignment terminal '--seedNoneLociPerWindow' option. Default value is
    "10"
STAR.Alignment.alignIntronMin
    STAR alignment terminal '--alignIntronMin' option. Default value is "21"
STAR.Alignment.alignIntronMax
    STAR alignment terminal '--alignIntronMax' option. Default value is "0"
STAR.Alignment.alignMatesGapMax
    STAR alignment terminal '--alignMatesGapMax' option. Default value is "0"
STAR.Alignment.alignSJoverhangMin
    STAR alignment terminal '--alignSJoverhangMin' option. Default value is "5"
STAR.Alignment.alignSJDBoverhangMin
    STAR alignment terminal '--alignSJDBoverhangMin' option. Default value is
    "3"
STAR.Alignment.alignSplicedMateMapLmin
    STAR alignment terminal '--alignSplicedMateMapLmin' option. Default value
    is "0"
```

```

STAR.Alignment.alignSplicedMateMapLminOverLmate
    STAR alignment terminal '-alignSplicedMateMapLminOverLmate' option. Default value is "0.66"
STAR.Alignment.alignWindowsPerReadNmax
    STAR alignment terminal '-alignWindowsPerReadNmax' option. Default value is "10000"
STAR.Alignment.alignTranscriptsPerWindowNmax
    STAR alignment terminal '-alignTranscriptsPerWindowNmax' option. Default value is "100"
STAR.Alignment.alignTranscriptsPerReadNmax
    STAR alignment terminal '-alignTranscriptsPerReadNmax' option. Default value is "10000"
STAR.Alignment.alignEndsType
    STAR alignment terminal '-alignEndsType' option. Default value is "Local"
STAR.Alignment.winAnchorMultimapNmax
    STAR alignment terminal '-winAnchorMultimapNmax' option. Default value is "50"
STAR.Alignment.winBinNbits
    STAR alignment terminal '-winBinNbits' option. Default value is "16"
STAR.Alignment.winAnchorDistNbns
    STAR alignment terminal '-winAnchorDistNbns' option. Default value is "9"
STAR.Alignment.winFlankNbns
    STAR alignment terminal '-winFlankNbns' option. Default value is "4"
Rsamtools.Bam.run
    Whether to run 'Rsamtools SAM to BAM' step in this function step. Default value is TRUE. Set FALSE to skip 'Rsamtools SAM to BAM' step.
Samtools.Bam.num.parallel.threads
    Specify the number of processing threads (CPUs) to use for Samtools sam to bam step. The default is "1"
Rsamtools.nCores
    The number of cores to use when running 'Rsamtools' step. Default value is 1
StringTie.Assemble.run
    Whether to run 'StringTie assembly' step in this function step. Default value is TRUE. Set FALSE to skip 'StringTie assembly' step.
Stringtie.Assembly.num.parallel.threads
    Specify the number of processing threads (CPUs) to use for Stringtie assembly. The default is "1"
Stringtie.Assembly.f
    Stringtie assembly terminal '-f' option. Default value is "0.1"
Stringtie.Assembly.m
    Stringtie assembly terminal '-m' option. Default value is "200"
Stringtie.Assembly.c
    Stringtie assembly terminal '-c' option. Default value is "2.5"
Stringtie.Assembly.g
    Stringtie assembly terminal '-g' option. Default value is "50"
Stringtie.Assembly.M
    Stringtie assembly terminal '-M' option. Default value is "0.95"
StringTie.Merge.Trans.run
    Whether to run 'StringTie GTF merging' step in this function step. Default value is TRUE. Set FALSE to skip 'StringTie GTF merging' step.

```

```

Stringtie.Merge.num.parallel.threads
    Specify the number of processing threads (CPUs) to use for Stringtie merge step.
    The default is "1"
Gffcompare.Ref.Sample.run
    Whether to run 'Gffcompare comparison' step in this function step. Default
    value is TRUE. Set FALSE to skip 'Gffcompare comparison' step.
StringTie.Ballgown.run
    Whether to run 'StringTie ballgown creation' step in this function step. Default
    value is TRUE. Set FALSE to skip 'StringTie ballgown creation' step.
Stringtie.2.Ballgown.num.parallel.threads
    Specify the number of processing threads (CPUs) to use for Stringtie to ball-
    gown step. The default is "1"
PreDECCountTable.run
    Whether to run 'gene raw reads count creation' step in this function step. Default
    value is TRUE. Set FALSE to skip 'gene raw reads count creation' step.
check.s4.print Default TRUE. If TRUE, the result of checking RNASEqRParam will be reported in
    'Rscript_out/Environment_Set.Rout'. If FALSE, the result of checking RNASEqRParam
    will not be in 'Rscript_out/Environment_Set.Rout'.

```

**Value**

None

**Author(s)**

Kuan-Hao Chao

**Examples**

```

data(yeast)
## Not run:
## Before run this function, make sure \code{RNASEqEnvironmentSet_CMD()}
##(or \code{RNASEqEnvironmentSet()}) is executed successfully.
RNASEqReadProcess(RNASEqRParam      = yeast,
                  num.parallel.threads = 10)
## End(Not run)

```

RNASEqReadProcess\_CMD *RNASEqReadProcess\_CMD*

**Description**

Process raw reads for RNA-Seq workflow in background.

This function do 5 things :

1. 'Hisat2' : aligns raw reads to reference genome. If `indices.optional` in `RNASEqRParam` is FALSE, Hisat2 indices will be created.
2. 'Rsamtools': converts '.sam' files to '.bam' files.

3. 'Stringtie': assembles alignments into transcript.
4. 'Gffcompare': examines how transcripts compare with the reference annotation.
5. 'Stringtie': creates input files for ballgown, edgeR and DESeq2.
6. raw reads count: create raw reads count for DESeq2 and edgeR

Before running this function, RNASeqEnvironmentSet\_CMD() or RNASeqEnvironmentSet() must be executed successfully.

If you want to process raw reads for the following RNA-Seq workflow in R shell, please see RNASeqReadProcess() function.

## Usage

```
RNASeqReadProcess_CMD(RNASeqRParam, SAMtools.or.Rsamtools = "Rsamtools",
  Hisat2.Index.run = TRUE, Hisat2.Index.num.parallel.threads = "1",
  Hisat2.Index.large.index = FALSE,
  Hisat2.Index.local.ftab.chars = "6",
  Hisat2.Index.local.off.rate = "3", Hisat2.Index.ftab.chars = "10",
  Hisat2.Index.off.rate = "4", Hisat2.Alignment.run = TRUE,
  Hisat2.Alignment.num.parallel.threads = "1",
  Hisat2.Alignment.skip = "0", Hisat2.Alignment.trim5 = "0",
  Hisat2.Alignment.trim3 = "0",
  Hisat2.Alignment.n.ceil.1.function.type = "L",
  Hisat2.Alignment.n.ceil.2.constant.term = "0",
  Hisat2.Alignment.n.ceil.3.coefficient = "0.15",
  Hisat2.Alignment.mp.MX = "6", Hisat2.Alignment.mp.MN = "2",
  Hisat2.Alignment.sp.MX = "2", Hisat2.Alignment.sp.MN = "1",
  Hisat2.Alignment.np = "1", Hisat2.Alignment.rdg.1 = "5",
  Hisat2.Alignment.rdg.2 = "3", Hisat2.Alignment.rfg.1 = "5",
  Hisat2.Alignment.rfg.2 = "3",
  Hisat2.Alignment.score.min.1.function.type = "L",
  Hisat2.Alignment.score.min.2.constant.term = "0",
  Hisat2.Alignment.score.min.3.coefficient = "-0.2",
  Hisat2.Alignment.pen.cansplice = "0",
  Hisat2.Alignment.penc.noncansplice = "12",
  Hisat2.Alignment.pen.canintronlen.1.function.type = "G",
  Hisat2.Alignment.pen.canintronlen.2.constant.term = "-8",
  Hisat2.Alignment.pen.canintronlen.3.coefficient = "1",
  Hisat2.Alignment.pen.noncanintronlen.1.function.type = "G",
  Hisat2.Alignment.pen.noncanintronlen.2.constant.term = "-8",
  Hisat2.Alignment.pen.noncanintronlen.3.coefficient = "1",
  Hisat2.Alignment.min.intronlen = "20",
  Hisat2.Alignment.max.intronlen = "500000",
  Hisat2.Alignment.rna.strandness = "None", Hisat2.Alignment.k = "5",
  Hisat2.Alignment.max.seeds = "5", Hisat2.Alignment.secondary = FALSE,
  Hisat2.Alignment.minins = "0", Hisat2.Alignment.maxins = "500",
  Hisat2.Alignment.seed = "0", STAR.Index.num.parallel.threads = "1",
  STAR.Index.sjdbOverhang.Read.length = "100",
```

```
STAR.Index.genomeSAindexNbases = "14",
STAR.Index.genomeChrBinNbites = "18",
STAR.Index.genomeSAsparseD = "1", STAR.Alignment.run = FALSE,
STAR.Alignment.num.parallel.threads = "1",
STAR.Alignment.genomeLoad = "NoSharedMemory",
STAR.Alignment.readMapNumber = "-1",
STAR.Alignment.clip3pNbases = "0", STAR.Alignment.clip5pNbases = "0",
STAR.Alignment.clip3pAdapterSeq = "-",
STAR.Alignment.clip3pAdapterMMP = "0.1",
STAR.Alignment.clip3pAfterAdapterNbases = "0",
STAR.Alignment.limitGenomeGenerateRAM = "31000000000",
STAR.Alignment.limitIObufferSize = "150000000",
STAR.Alignment.limitOutSAMoneReadBytes = "100000",
STAR.Alignment.limitOutSJoneRead = "1000",
STAR.Alignment.limitOutSJcollapsed = "1000000",
STAR.Alignment.limitBAMsortRAM = "0",
STAR.Alignment.outReadsUnmapped = "None",
STAR.Alignment.outQSconversionAdd = "0",
STAR.Alignment.outSAMprimaryFlag = "OneBestScore",
STAR.Alignment.outSAMmapqUnique = "255",
STAR.Alignment.scoreGap = "0", STAR.Alignment.scoreGapNoncan = "-8",
STAR.Alignment.scoreGapGCAG = "-4",
STAR.Alignment.scoreGapATAC = "-8",
STAR.Alignment.scoreGenomicLengthLog2scale = "-0.25",
STAR.Alignment.scoreDelOpen = "-2",
STAR.Alignment.scoreDelBase = "-2",
STAR.Alignment.scoreInsOpen = "-2",
STAR.Alignment.scoreInsBase = "-2",
STAR.Alignment.scoreStitchSJshift = "1",
STAR.Alignment.seedSearchStartLmax = "50",
STAR.Alignment.seedSearchStartLmaxOverLread = "1.0",
STAR.Alignment.seedSearchLmax = "0",
STAR.Alignment.seedMultimapNmax = "10000",
STAR.Alignment.seedPerReadNmax = "1000",
STAR.Alignment.seedPerWindowNmax = "50",
STAR.Alignment.seedNoneLociPerWindow = "10",
STAR.Alignment.alignIntronMin = "21",
STAR.Alignment.alignIntronMax = "0",
STAR.Alignment.alignMatesGapMax = "0",
STAR.Alignment.alignSJoverhangMin = "5",
STAR.Alignment.alignSJDBoverhangMin = "3",
STAR.Alignment.alignSplicedMateMapLmin = "0",
STAR.Alignment.alignSplicedMateMapLminOverLmate = "0.66",
STAR.Alignment.alignWindowsPerReadNmax = "10000",
STAR.Alignment.alignTranscriptsPerWindowNmax = "100",
STAR.Alignment.alignTranscriptsPerReadNmax = "10000",
STAR.Alignment.alignEndsType = "Local",
STAR.Alignment.winAnchorMultimapNmax = "50",
STAR.Alignment.winBinNbites = "16",
STAR.Alignment.winAnchorDistNbites = "9",
STAR.Alignment.winFlankNbites = "4", Rsamtools.Bam.run = TRUE,
Samtools.Bam.num.parallel.threads = "1", Rsamtools.nCores = "1",
```

```

StringTie.Assemble.run = TRUE,
Stringtie.Assembly.num.parallel.threads = "1",
Stringtie.Assembly.f = "0.1", Stringtie.Assembly.m = "200",
Stringtie.Assembly.c = "2.5", Stringtie.Assembly.g = "50",
Stringtie.Assembly.M = "0.95", StringTie.Merge.Trans.run = TRUE,
Stringtie.Merge.num.parallel.threads = "1",
Gffcompare.Ref.Sample.run = TRUE, StringTie.Ballgown.run = TRUE,
Stringtie.2.Ballgown.num.parallel.threads = "1",
PreDECCountTable.run = TRUE, run = TRUE, check.s4.print = TRUE)

```

## Arguments

**RNASeqRParam** S4 object instance of experiment-related parameters

**SAMtools.or.Rsamtools**  
Default value is Rsamtools. User can set to SAMtools to use command-line-based 'samtools' instead.

**Hisat2.Index.run**  
Whether to run 'HISAT2 index' step in this function step. Default value is TRUE. Set FALSE to skip 'HISAT2 index' step.

**Hisat2.Index.num.parallel.threads**  
Specify the number of processing threads (CPUs) to use for Hisat2 index step. The default is "1"

**Hisat2.Index.large.index**  
Hisat2 index terminal '--large-index' option. Default value is FALSE

**Hisat2.Index.local.ftab.chars**  
Hisat2 index terminal '-t/-ftabchars' option. Default value is "6"

**Hisat2.Index.local.off.rate**  
Hisat2 index terminal '--localoffrate' option. Default value is "3"

**Hisat2.Index.ftab.chars**  
Hisat2 index terminal '--localftabchars' option. Default value is "10"

**Hisat2.Index.off.rate**  
Hisat2 index terminal '--offrate' option. Default value is "4"

**Hisat2.Alignment.run**  
Whether to run 'HISAT2 alignment' step in this function step. Default value is TRUE. Set FALSE to skip 'HISAT2 alignment' step.

**Hisat2.Alignment.num.parallel.threads**  
Specify the number of processing threads (CPUs) to use for Hisat2 alignment step. The default is "1"

**Hisat2.Alignment.skip**  
Hisat2 alignment terminal '-s/-skip' option. Default value is "0"

**Hisat2.Alignment.trim5**  
Hisat2 alignment terminal '-5/-trim5' option. Default value is "0"

**Hisat2.Alignment.trim3**  
Hisat2 alignment terminal '-3/-trim3' option. Default value is "0"

**Hisat2.Alignment.n.ceil.1.function.type**  
Hisat2 alignment terminal '--n-ceil' option. Default value is "L"

**Hisat2.Alignment.n.ceil.2.constant.term**  
Hisat2 alignment terminal '--n-ceil' option. Default value is "0"

**Hisat2.Alignment.n.ceil.3.coefficient**  
Hisat2 alignment terminal '--n-ceil' option. Default value is "0.15"

```

Hisat2.Alignment.mp.MX
    Hisat2 alignment terminal '-mp MX' option. Default value is "6"
Hisat2.Alignment.mp.MN
    Hisat2 alignment terminal '-mp MN' option. Default value is "2"
Hisat2.Alignment.sp.MX
    Hisat2 alignment terminal '-sp MX' option. Default value is "2"
Hisat2.Alignment.sp.MN
    Hisat2 alignment terminal '-sp MN' option. Default value is "1"
Hisat2.Alignment.np
    Hisat2 alignment terminal '-np' option. Default value is "1"
Hisat2.Alignment.rdg.1
    Hisat2 alignment terminal '-rdg' first option. Default value is "5"
Hisat2.Alignment.rdg.2
    Hisat2 alignment terminal '-rdg' first option. Default value is "3"
Hisat2.Alignment.rfg.1
    Hisat2 alignment terminal '-rfg' first option. Default value is "5"
Hisat2.Alignment.rfg.2
    Hisat2 alignment terminal '-rfg' first option. Default value is "3"
Hisat2.Alignment.score.min.1.function.type
    Hisat2 alignment terminal '-rdg' first option. Default value is "L"
Hisat2.Alignment.score.min.2.constant.term
    Hisat2 alignment terminal '-rdg' first option. Default value is "0"
Hisat2.Alignment.score.min.3.coefficient
    Hisat2 alignment terminal '-rdg' first option. Default value is "-0.2"
Hisat2.Alignment.pen.cansplice
    Hisat2 alignment terminal '-pen-cansplice' first option. Default value is "-0"
Hisat2.Alignment.penc.noncansplice
    Hisat2 alignment terminal '-pen-noncansplice' option. Default value is "12"
Hisat2.Alignment.pen.canintronlen.1.function.type
    Hisat2 alignment terminal '-pen-canintronlen' first option. Default value is "G"
Hisat2.Alignment.pen.canintronlen.2.constant.term
    Hisat2 alignment terminal '-pen-canintronlen' second option. Default value is
    "-8"
Hisat2.Alignment.pen.canintronlen.3.coefficient
    Hisat2 alignment terminal '-pen-canintronlen' third option. Default value is "1"
Hisat2.Alignment.pen.noncanintronlen.1.function.type
    Hisat2 alignment terminal '-pen-noncanintronlen' first option. Default value is
    "G"
Hisat2.Alignment.pen.noncanintronlen.2.constant.term
    Hisat2 alignment terminal '-pen-noncanintronlen' second option. Default value
    is "-8"
Hisat2.Alignment.pen.noncanintronlen.3.coefficient
    Hisat2 alignment terminal '-pen-noncanintronlen' third option. Default value is
    "1"
Hisat2.Alignment.min.intronlen
    Hisat2 alignment terminal '-min-intronlen' option. Default value is "20"
Hisat2.Alignment.max.intronlen
    Hisat2 alignment terminal '-max-intronlen' option. Default value is "20"

```

```

Hisat2.Alignment.rna.strandness
    Hisat2 alignment terminal '-rna-strandness' option. Default value is "None"
Hisat2.Alignment.k
    Hisat2 alignment terminal '-k' option. Default value is "5"
Hisat2.Alignment.max.seeds
    Hisat2 alignment terminal '-max-seeds' option. Default value is "5"
Hisat2.Alignment.secondary
    Hisat2 alignment terminal '-secondary' option. Default value is "FALSE"
Hisat2.Alignment.minins
    Hisat2 alignment terminal '-I/-minins' option. Default value is "0"
Hisat2.Alignment.maxins
    Hisat2 alignment terminal '-X/-maxins' option. Default value is "500"
Hisat2.Alignment.seed
    Hisat2 alignment terminal '-X/-maxins' option. Default value is "0"
STAR.Index.num.parallel.threads
    Specify the number of processing threads (CPUs) to use for STAR index step.
    The default is "1"
STAR.Index.sjdbOverhang.Read.length
    STAR index terminal '-sjdbOverhang' option. Default value is "100"
STAR.Index.genomeSAindexNbases
    STAR index terminal '-genomeSAindexNbases' option. Default value is "14"
STAR.Index.genomeChrBinNbits
    STAR index terminal '-genomeChrBinNbits' option. Default value is "18"
STAR.Index.genomeSAsparseD
    STAR index terminal '-genomeSAsparseD' option. Default value is "1"
STAR.Alignment.run
    Whether to run 'STAR index' step in this function step. Default value is FALSE.
    Set TRUE to run STAR alignment step. (need to set Hisat2.Index.run to FALSE)
STAR.Alignment.num.parallel.threads
    Specify the number of processing threads (CPUs) to use for STAR alignment
    step. The default is "1"
STAR.Alignment.genomeLoad
    STAR alignment terminal '-genomeLoad' option. Default value is "NoSharedMemory"
STAR.Alignment.readMapNumber
    STAR alignment terminal '-readMapNumber' option. Default value is "-1"
STAR.Alignment.clip3pNbases
    STAR alignment terminal '-clip3pNbases' option. Default value is "0"
STAR.Alignment.clip5pNbases
    STAR alignment terminal '-clip5pNbases' option. Default value is "0"
STAR.Alignment.clip3pAdapterSeq
    STAR alignment terminal '-clip3pAdapterSeq' option. Default value is "-"
STAR.Alignment.clip3pAdapterMMP
    STAR alignment terminal '-clip3pAdapterMMP' option. Default value is "0.1"
STAR.Alignment.clip3pAfterAdapterNbases
    STAR alignment terminal '-clip3pAfterAdapterNbases' option. Default value
    is "0"
STAR.Alignment.limitGenomeGenerateRAM
    STAR alignment terminal '-limitGenomeGenerateRAM' option. Default value
    is "31000000000"

```

```
STAR.Alignment.limitIObufferSize
    STAR alignment terminal '-limitIObufferSize' option. Default value is "150000000"
STAR.Alignment.limitOutSAMoneReadBytes
    STAR alignment terminal '-limitOutSAMoneReadBytes' option. Default value
    is "100000"
STAR.Alignment.limitOutSJoneRead
    STAR alignment terminal '-limitOutSJoneRead' option. Default value is "1000"
STAR.Alignment.limitOutSJcollapsed
    STAR alignment terminal '-limitOutSJcollapsed' option. Default value is "1000000"
STAR.Alignment.limitBAMsortRAM
    STAR alignment terminal '-limitBAMsortRAM' option. Default value is "0"
STAR.Alignment.outReadsUnmapped
    STAR alignment terminal '-outReadsUnmapped' option. Default value is "None"
STAR.Alignment.outQsconversionAdd
    STAR alignment terminal '-outQsconversionAdd' option. Default value is "0"
STAR.Alignment.outSAMprimaryFlag
    STAR alignment terminal '-outSAMprimaryFlag' option. Default value is "OneBestScore"
STAR.Alignment.outSAMmapqUnique
    STAR alignment terminal '-outSAMmapqUnique' option. Default value is "255"
STAR.Alignment.scoreGap
    STAR alignment terminal '-scoreGap' option. Default value is "0"
STAR.Alignment.scoreGapNoncan
    STAR alignment terminal '-scoreGapNoncan' option. Default value is "-8"
STAR.Alignment.scoreGapGCAG
    STAR alignment terminal '-scoreGapGCAG' option. Default value is "-4"
STAR.Alignment.scoreGapATAC
    STAR alignment terminal '-scoreGapATAC' option. Default value is "-8"
STAR.Alignment.scoreGenomicLengthLog2scale
    STAR alignment terminal '-scoreGenomicLengthLog2scale' option. Default
    value is "-0.25"
STAR.Alignment.scoreDelOpen
    STAR alignment terminal '-scoreDelOpen' option. Default value is "-2"
STAR.Alignment.scoreDelBase
    STAR alignment terminal '-scoreDelBase' option. Default value is "-2"
STAR.Alignment.scoreInsOpen
    STAR alignment terminal '-scoreInsOpen' option. Default value is "-2"
STAR.Alignment.scoreInsBase
    STAR alignment terminal '-scoreInsBase' option. Default value is "-2"
STAR.Alignment.scoreStitchSJshift
    STAR alignment terminal '-scoreStitchSJshift' option. Default value is "1"
STAR.Alignment.seedSearchStartLmax
    STAR alignment terminal '-scoreStitchSJshift' option. Default value is "50"
STAR.Alignment.seedSearchStartLmaxOverLread
    STAR alignment terminal '-seedSearchStartLmaxOverLread' option. Default
    value is "1.0"
STAR.Alignment.seedSearchLmax
    STAR alignment terminal '-seedSearchLmax' option. Default value is "0"
STAR.Alignment.seedMultimapNmax
    STAR alignment terminal '-seedMultimapNmax' option. Default value is "10000"
```

```

STAR.Alignment.seedPerReadNmax
    STAR alignment terminal '-seedPerReadNmax' option. Default value is "1000"
STAR.Alignment.seedPerWindowNmax
    STAR alignment terminal '-seedPerWindowNmax' option. Default value is
    "50"
STAR.Alignment.seedNoneLociPerWindow
    STAR alignment terminal '-seedNoneLociPerWindow' option. Default value is
    "10"
STAR.Alignment.alignIntronMin
    STAR alignment terminal '-alignIntronMin' option. Default value is "21"
STAR.Alignment.alignIntronMax
    STAR alignment terminal '-alignIntronMax' option. Default value is "0"
STAR.Alignment.alignMatesGapMax
    STAR alignment terminal '-alignMatesGapMax' option. Default value is "0"
STAR.Alignment.alignSJoverhangMin
    STAR alignment terminal '-alignSJoverhangMin' option. Default value is "5"
STAR.Alignment.alignSJDBoverhangMin
    STAR alignment terminal '-alignSJDBoverhangMin' option. Default value is
    "3"
STAR.Alignment.alignSplicedMateMapLmin
    STAR alignment terminal '-alignSplicedMateMapLmin' option. Default value
    is "0"
STAR.Alignment.alignSplicedMateMapLminOverLmate
    STAR alignment terminal '-alignSplicedMateMapLminOverLmate' option. De-
    fault value is "0.66"
STAR.Alignment.alignWindowsPerReadNmax
    STAR alignment terminal '-alignWindowsPerReadNmax' option. Default value
    is "10000"
STAR.Alignment.alignTranscriptsPerWindowNmax
    STAR alignment terminal '-alignTranscriptsPerWindowNmax' option. Default
    value is "100"
STAR.Alignment.alignTranscriptsPerReadNmax
    STAR alignment terminal '-alignTranscriptsPerReadNmax' option. Default value
    is "10000"
STAR.Alignment.alignEndsType
    STAR alignment terminal '-alignEndsType' option. Default value is "Local"
STAR.Alignment.winAnchorMultimapNmax
    STAR alignment terminal '-winAnchorMultimapNmax' option. Default value
    is "50"
STAR.Alignment.winBinNbits
    STAR alignment terminal '-winBinNbits' option. Default value is "16"
STAR.Alignment.winAnchorDistNbins
    STAR alignment terminal '-winAnchorDistNbins' option. Default value is "9"
STAR.Alignment.winFlankNbins
    STAR alignment terminal '-winFlankNbins' option. Default value is "4"
Rsamtools.Bam.run
    Whether to run 'Rsamtools SAM to BAM' step in this function step. Default
    value is TRUE. Set FALSE to skip 'Rsamtools SAM to BAM' step.
Samtools.Bam.num.parallel.threads
    Specify the number of processing threads (CPUs) to use for Samtools sam to
    bam step. The default is "1"

```

```

Rsamtools.nCores
    The number of cores to use when running 'Rsamtools' step. Default value is 1

StringTie.Assemble.run
    Whether to run 'StringTie assembly' step in this function step. Default value is TRUE. Set FALSE to skip 'StringTie assembly' step.

Stringtie.Assembly.num.parallel.threads
    Specify the number of processing threads (CPUs) to use for Stringtie assembly. The default is "1"

Stringtie.Assembly.f
    Stringtie assembly terminal '-f' option. Default value is "0.1"

Stringtie.Assembly.m
    Stringtie assembly terminal '-m' option. Default value is "200"

Stringtie.Assembly.c
    Stringtie assembly terminal '-c' option. Default value is "2.5"

Stringtie.Assembly.g
    Stringtie assembly terminal '-g' option. Default value is "50"

Stringtie.Assembly.M
    Stringtie assembly terminal '-M' option. Default value is "0.95"

StringTie.Merge.Trans.run
    Whether to run 'StringTie GTF merging' step in this function step. Default value is TRUE. Set FALSE to skip 'StringTie GTF merging' step.

Stringtie.Merge.num.parallel.threads
    Specify the number of processing threads (CPUs) to use for Stringtie merge step. The default is "1"

Gffcompare.Ref.Sample.run
    Whether to run 'Gffcompare comparison' step in this function step. Default value is TRUE. Set FALSE to skip 'Gffcompare comparison' step.

StringTie.Ballgown.run
    Whether to run 'StringTie ballgown creation' step in this function step. Default value is TRUE. Set FALSE to skip 'StringTie ballgown creation' step.

Stringtie.2.Ballgown.num.parallel.threads
    Specify the number of processing threads (CPUs) to use for Stringtie to ballgown step. The default is "1"

PreDECCountTable.run
    Whether to run 'gene raw reads count creation' step in this function step. Default value is TRUE. Set FALSE to skip 'gene raw reads count creation' step.

run
    Default value is TRUE. If TRUE, 'Rscript/Environment_Set.R' will be created and executed. The output log will be stored in 'Rscript_out/Environment_Set.Rout'. If False, 'Rscript/Environment_Set.R' will be created without executed.

check.s4.print
    Default TRUE. If TRUE, the result of checking RNASEqRParam will be reported in 'Rscript_out/Environment_Set.Rout'. If FALSE, the result of checking RNASEqRParam will not be in 'Rscript_out/Environment_Set.Rout'.

```

**Value**

None

**Author(s)**

Kuan-Hao Chao

## Examples

```
data(yeast)
## Not run:
## Before run this function, make sure \code{RNASEqEnvironmentSet_CMD()}
## (or \code{RNASEqEnvironmentSet()}) is executed successfully.
RNASEqReadProcess_CMD(RNASEqRParam = yeast,
                      num.parallel.threads = 10)
## End(Not run)
```

*RNASEqRParam-class*      *RNASEqR*

## Description

An S4 class for checking and storing RNA-Seq workflow parameters of this package.

## Slots

`os.type` 'linux' or 'osx'. The operating system type.

`python.variable` A list storing python environment. (`check.answer`, `python.version`)

`python.2to3` Logical value whether 2to3 command is available on the workstation.

`path.prefix` Path prefix of 'gene\_data/' , 'RNASEq\_bin/' , 'RNASEq\_results/' , 'Rscript/' and 'Rscript\_out/' directories.

`input.path.prefix` Path prefix of 'input\_files/' directory,

`genome.name` Variable of genome name defined in this RNA-Seq workflow (ex. `genome.name.fa`, `genome.name.gtf`).

`sample.pattern` Regular expression of paired-end fastq.gz files under 'input\_files/raw\_fastq.gz'. Expression not includes `_[1,2].fastq.gz`.

`independent.variable` Independent variable for the biological. experiment design of two-group RNA-Seq workflow.

`case.group` Group name of the case group.

`control.group` Group name of the control group.

`indices.optional` Logical value whether 'indices/' is exit in 'input\_files/'.

`fastq.gz.type` Specify the fastq.gz file type. 'PE' represents paired-end and 'SE' represents single-end.

## Author(s)

Kuan-Hao Chao

## Examples

```
data(yeast)
"@"(yeast, os.type)
"@"(yeast, python.variable)
"@"(yeast, python.2to3)
"@"(yeast, path.prefix)
"@"(yeast, input.path.prefix)
"@"(yeast, genome.name)
"@"(yeast, sample.pattern)
"@"(yeast, independent.variable)
"@"(yeast, case.group)
"@"(yeast, control.group)
"@"(yeast, indices.optional)
"@"(yeast, fastq.gz.type)
```

## RNASeqRParam-constructor

*RNASeqRParam*

## Description

Constructor function for RNASeqRParam objects

## Usage

```
RNASeqRParam(path.prefix = NA, input.path.prefix = NA,
  genome.name = NA, sample.pattern = NA, independent.variable = NA,
  case.group = NA, control.group = NA, fastq.gz.type = NA)
```

## Arguments

path.prefix	Path prefix of 'gene_data/', 'RNASeq_bin/', 'RNASeq_results/', 'Rscript/' and 'Rscript_out/' directories.
input.path.prefix	Path prefix of 'input_files/' directory.
genome.name	variable of genome name defined in this RNA-Seq workflow (ex. genome.name.fa, genome.name.gtf).
sample.pattern	Regular expression of paired-end fastq.gz files under 'input_files/raw_fastq.gz'. Expression not includes _[1,2].fastq.gz.
independent.variable	Independent variable for the biological experiment design of two-group RNA-Seq workflow.
case.group	Group name of the case group.
control.group	Group name of the control group.
fastq.gz.type	Specify the fastq.gz file type. 'PE' represents paired-end and 'SE' represents single-end.

## Value

an object of class RNASeqRParam

**Author(s)**

kuan-hao Chao  
 Kuan-Hao Chao

**Examples**

```
input_files.path <- system.file("extdata/", package = "RNASeqRData")
rnaseq_result.path <- tempdir(check = TRUE)
exp <- RNASeqRParam(path.prefix      = rnaseq_result.path,
                     input.path.prefix = input_files.path,
                     genome.name       = "Saccharomyces_cerevisiae_XV_Eensembl",
                     sample.pattern    = "SRR[0-9]*_XV",
                     independent.variable = "state",
                     case.group        = "60mins_ID20_amphotericin_B",
                     control.group     = "60mins_ID20_control",
                     fastq.gz.type    = "PE")
```

**RNASeqRParam\_Bam-class***RNASeqRParam\_Bam***Description**

An S4 class for checking and storing RNA-Seq workflow parameters starting with BAM files.

**Slots**

**os.type** 'linux' or 'osx'. The operating system type.  
**python.variable** A list storing python environment. (check.answer,python.version)  
**python.2to3** Logical value whether 2to3 command is available on the workstation.  
**path.prefix** Path prefix of 'gene\_data/','RNASeq\_bin/','RNASeq\_results/','Rscript/' and 'Rscript\_out/' directories.  
**input.path.prefix** Path prefix of 'input\_files/' directory,  
**genome.name** Variable of genome name defined in this RNA-Seq workflow (ex. genome.name.fa, genome.name.gtf).  
**sample.pattern** Regular expression of paired-end fastq.gz files under 'input\_files/raw\_bam'. Expression not includes \_[1,2].fastq.gz.  
**independent.variable** Independent variable for the biological. experiment design of two-group RNA-Seq workflow.  
**case.group** Group name of the case group.  
**control.group** Group name of the control group.

**Author(s)**

Kuan-Hao Chao

## Examples

```
data(yeast)
"@"(yeast, os.type)
"@"(yeast, python.variable)
"@"(yeast, python.2to3)
"@"(yeast, path.prefix)
"@"(yeast, input.path.prefix)
"@"(yeast, genome.name)
"@"(yeast, sample.pattern)
"@"(yeast, independent.variable)
"@"(yeast, case.group)
"@"(yeast, control.group)
```

RNASeqRParam\_Bam-constructor  
*RNASeqR\_Bam*

## Description

Constructor function for RNASeqRParam\_Bam objects

## Usage

```
RNASeqRParam_Bam(path.prefix = NA, input.path.prefix = NA,
  genome.name = NA, sample.pattern = NA, independent.variable = NA,
  case.group = NA, control.group = NA)
```

## Arguments

path.prefix	Path prefix of 'gene_data/', 'RNASeq_bin/', 'RNASeq_results/', 'Rscript/' and 'Rscript_out/' directories.
input.path.prefix	Path prefix of 'input_files/' directory.
genome.name	variable of genome name defined in this RNA-Seq workflow (ex. genome.name.fa, genome.name.gtf).
sample.pattern	Regular expression of paired-end fastq.gz files under 'input_files/raw_bam'. Expression not includes _[1,2].fastq.gz.
independent.variable	Independent variable for the biological experiment design of two-group RNA-Seq workflow.
case.group	Group name of the case group.
control.group	Group name of the control group.

## Value

an object of class RNASeqRParam\_Bam

## Author(s)

Kuan-Hao Chao  
 Kuan-Hao Chao

## Examples

```
input_files.path <- system.file("extdata/", package = "RNASeqRData")
rnaseq_result.path <- tempdir(check = TRUE)
## Not run:
exp <- RNASeqRParam_Bam(path.prefix      = rnaseq_result.path,
                         input.path.prefix = input_files.path,
                         genome.name       = "Saccharomyces_cerevisiae_XV_Eensembl",
                         sample.pattern    = "SRR[0-9]*_XV",
                         independent.variable = "state",
                         case.group        = "60mins_ID20_amphotericin_B",
                         control.group     = "60mins_ID20_control")

## End(Not run)
```

RNASeqRParam\_Sam-class

*RNASeqRParam\_Sam*

## Description

An S4 class for checking and storing RNA-Seq workflow parameters starting with SAM files.

## Slots

os.type 'linux' or 'osx'. The operating system type.  
 python.variable A list storing python environment. (check.answer,python.version)  
 python.2to3 Logical value whether 2to3 command is available on the workstation.  
 path.prefix Path prefix of 'gene\_data/', 'RNASeq\_bin/', 'RNASeq\_results/', 'Rscript/' and 'Rscript\_out/' directories.  
 input.path.prefix Path prefix of 'input\_files/' directory,  
 genome.name Variable of genome name defined in this RNA-Seq workflow (ex. genome.name.fa, genome.name.gtf).  
 sample.pattern Regular expression of paired-end fastq.gz files under 'input\_files/raw\_sam'. Expression not includes \_[1,2].fastq.gz.  
 independent.variable Independent variable for the biological. experiment design of two-group RNA-Seq workflow.  
 case.group Group name of the case group.  
 control.group Group name of the control group.

## Author(s)

Kuan-Hao Chao

## Examples

```
data(yeast)
"@"(yeast, os.type)
"@"(yeast, python.variable)
"@"(yeast, python.2to3)
"@"(yeast, path.prefix)
"@"(yeast, input.path.prefix)
"@"(yeast, genome.name)
"@"(yeast, sample.pattern)
"@"(yeast, independent.variable)
"@"(yeast, case.group)
"@"(yeast, control.group)
```

RNASeqRParam\_Sam-constructor  
*RNASeqR\_Sam*

## Description

Constructor function for RNASeqRParam\_Sam objects

## Usage

```
RNASeqRParam_Sam(path.prefix = NA, input.path.prefix = NA,
  genome.name = NA, sample.pattern = NA, independent.variable = NA,
  case.group = NA, control.group = NA)
```

## Arguments

path.prefix	Path prefix of 'gene_data/', 'RNASeq_bin/', 'RNASeq_results/', 'Rscript/' and 'Rscript_out/' directories.
input.path.prefix	Path prefix of 'input_files/' directory.
genome.name	variable of genome name defined in this RNA-Seq workflow (ex. genome.name.fa, genome.name.gtf).
sample.pattern	Regular expression of paired-end fastq.gz files under 'input_files/raw_sam'. Expression not includes _[1,2].fastq.gz.
independent.variable	Independent variable for the biological experiment design of two-group RNA-Seq workflow.
case.group	Group name of the case group.
control.group	Group name of the control group.

## Value

an object of class RNASeqRParam\_Sam

## Author(s)

Kuan-Hao Chao  
 Kuan-Hao Chao

### Examples

```
input_files.path <- system.file("extdata/", package = "RNASeqRData")
rnaseq_result.path <- tempdir(check = TRUE)
## Not run:
exp <- RNASeqRParam_Sam(path.prefix      = rnaseq_result.path,
                         input.path.prefix = input_files.path,
                         genome.name       = "Saccharomyces_cerevisiae_XV_Ensembl",
                         sample.pattern    = "SRR[0-9]*_XV",
                         independent.variable = "state",
                         case.group        = "60mins_ID20_amphotericin_B",
                         control.group     = "60mins_ID20_control")

## End(Not run)
```

*Update\_Fastq\_gz*

*Update\_Fastq\_gz*

### Description

This function let users update their trimmed fastq.gz files automatically.

### Usage

```
Update_Fastq_gz(RNASeqRParam, prepared_fastq_gz, target_samples = "ALL")
```

### Arguments

RNASeqRParam	S4 object instance of experiment-related parameters
prepared_fastq_gz	absolute path to the prepared 'raw_fastq.gz' directory.
target_samples	list of samples that are going to update. Default value is ALL

### Value

None

### Author(s)

Kuan-Hao Chao

### Examples

```
data(yeast)
## Not run:
RNASeqDifferentialAnalysis(RNASeqRParam = yeast)
## End(Not run)
```

---

yeast

*Toy RNASEqRParam object*

---

### Description

Small RNASEqRParam S4 object created with checked valid parameters for demonstration purposes

### Author(s)

Kuan-Hao Chao

### Examples

```
data(yeast)
yeast
# RNASEqRParam S4 object for example demonstration.
```

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