# **ASpedia-R**

November 22, 2023

asr\_converter

Generate ASpedia-R input format from DAS analysis tools result.

#### **Description**

result of DAS analysis tools convert to ASpedia-R input format.

# Usage

```
asr_converter(
  das.analysis.result = "",
  program = "",
  pvalue.cutoff = 0.05,
  dpsi.cutoff = 0.1,
  gene.model = "Ensembl",
  genome.version = "GRCh38",
  as.type = "",
  gtf.file.name = "",
  ioe.file.name = "")
```

#### **Arguments**

```
das.analysis.result
```

name of DAS analysis tools result file.

program name of DAS analysis tool. one of rMATS, SUPPA, or spliceR

pvalue.cutoff value of pvalue cutoff. default value is 0.05 dpsi.cutoff value of dPSI cutoff. default value is 0.1

gene model of reference. One of Refseq, Ensembl, or GENCODE. (spliceR

only)

genome.version genome version of reference. One of hg18, GRCh19, or GRCh38. (spliceR

only)

as.type AS event type. One of A3SS, A5SS, SE, MXE, or RI. (rMATS only)

gtf.file.name a GTF format file of reference. (SUPPA only)

ioe.file.name name of ioe file generated by SUPPA generateEvents command. (SUPPA)

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#### Value

converter.result

#### **Examples**

rMATS.converter

Generate ASpedia-R input format from rMATS result.

#### **Description**

rMSTS result convert to ASpedia-R input format.

# Usage

```
rMATS.converter(
  rMATS.result = "",
  as.type = "",
  pvalue.cutoff = 0.05,
  dpsi.cutoff = 0.1
)
```

#### **Arguments**

```
rMATS.result name of rMATS result file.

as.type AS event type. One of A3SS, A5SS, SE, MXE, or RI.

pvalue.cutoff value of pvalue cutoff. default value is 0.05

dpsi.cutoff value of dPSI cutoff. default value is 0.1
```

#### Value

converting.result

# Examples

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SUPPA.converter

Generate ASpedia-R input format from SUPPA result.

# Description

SUPPA result convert to ASpedia-R input format.

#### Usage

```
SUPPA.converter(
  SUPPA.result = "",
  pvalue.cutoff = 0.05,
  dpsi.cutoff = 0.1,
  gtf.file.name = "",
  ioe.file.name = ""
)
```

# **Arguments**

```
suppa.result name of suppa result file.

pvalue.cutoff value of pvalue cutoff. default value is 0.05

dpsi.cutoff value of dPSI cutoff. default value is 0.1

gtf.file.name a GTF format file of reference.

ioe.file.name name of ioe file generated by SUPPA generateEvents command.
```

# Value

converting.result

# **Examples**

spliceR.converter

Generate ASpedia-R input format from spliceR result.

# **Description**

spliceR result convert to ASpedia-R input format.

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#### Usage

```
spliceR.converter(
  spliceR.result = "",
  pvalue.cutoff = 0.05,
  gene.model = "Ensembl",
  genome.version = "GRCh38"
)
```

# **Arguments**

```
spliceR.result name of spliceR result file.

pvalue.cutoff value of pvalue cutoff. default value is 0.05

gene.model gene model of reference. One of Refseq, Ensembl, or GENCODE.

genome.version genome version of reference. One of hg18, GRCh19, or GRCh38.
```

#### Value

converting.result

#### **Examples**

asr\_annotation

converting result mapping to ASDB

#### **Description**

DNA and RNA annotation related to aliternative splicing from ASpedia DB(ASDB) add to asr\_converter (rMATS\_converter, SUPPA\_converter, or spliceR\_converter) result. And gene enrichment test result are provided between annotation result gene list and knowledge-based database gene list

# Usage

```
asr_annotation(
  converter.result = "",
  gene.model = "Ensembl",
  genome.version = "GRCh38",
  gsea.gene.list = "",
  result.dir = ""
)
```

#### **Arguments**

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```
genome.version genome version of reference. One of hg18, GRCh19, or GRCh38.

gsea.gene.list optional. reference gene list for gene enrichment test with annotation result gene list and knowledge-based database gene list. If gene list is empty, use all genes in reference.

result.dir directory where annotation result(.tsv file) is saved
```

#### Value

annotation result

#### **Examples**

mining\_gsea

gene enrichment test between annotation gene list and knowledgebased database gene list.

# **Description**

Gene enrichment test result are provided to tsv and plot format between annotation gene list and knowledge-based database gene list.

#### Usage

```
mining_gsea(
  annotation.gene.list = "",
  gsea.gene.list = "",
  gene.model = "",
  genome.version = "",
  result.dir = ""
)
```

#### **Arguments**

```
annotation.gene.list
gene list from asr_annotation function result
gsea.gene.list gene list from reference
gene.model gene model of reference. One of Refseq, Ensembl, or GENCODE.
genome.version genome version of reference. One of hg18, GRCh19, or GRCh38.
result.dir directory where GSEA result(.tsv and .png file) are saved
```

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#### **Examples**

asr\_plot

visualization of annotation result

#### **Description**

DNA and RNA features in result of asr\_annotation function to be visualize.

#### Usage

```
asr_plot(
  annotation.result = "",
  gtf.file.name = "",
  gene.model = "Ensembl",
  genome.version = "GRCh38",
  gene.name = "",
  as.id = "",
  heights.list = "",
  list.of.plot = "",
  result.dir = ""
```

# **Arguments**

```
annotation.result
                  asr_annotation function result.
gtf.file.name a GTF format file of reference.
                  gene model of reference. One of Refseq, Ensembl, or GENCODE.
gene.model
genome.version genome version of reference. One of hg18, GRCh19, or GRCh38.
gene.name
                  gene name to be visualization.
as.id
                  list of AS ID to be visualization.
                  positive integer vectors for track heights include height of ideogram and gene
heights.list
list.of.plot
                  list of DNA or RNA feature to be visualization. choose from "conservation",
                  "NMD", "repeats", "domain", "PTM", or "RBP".
                  directory where plots(.png files) are saved
result.dir
```

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#### **Examples**

asr\_plot\_ppi

visualization of PPI network

# Description

If asr\_annotation function result has protein protein interaction(PPI) information, we provide PPI network plot.

#### Usage

```
asr_plot_ppi(
  annotation.result = "",
  gene.name = "",
  as.id = "",
  result.dir = ""
)
```

#### **Arguments**

#### **Examples**

```
ppi.result.dir <- system.file("extdata/ppi_result", package="ASpediaR")
##using gene name
asr_plot_ppi(annotation.result, gene.name="FGFR2", result.dir=ppi.result.dir)
##using AS ID</pre>
```

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