# **RMAPPER**

October 25, 2011

hits Hits details

# Description

Function displaying the detailed information on the sequence and other properties of each hit returned from the MAPPER RPC interface query.

# Usage

hits(x)

# Arguments

Х

x – mapperHits object returned from readMAPPER

# References

```
http://genome.ufl.edu/mapper/
```

## See Also

readMAPPER.

# **Examples**

# see readMAPPER

2 mapperHits-class

```
mapperHits-class Class "mapperHits" - holds collection of hits from MAPPER
```

# Description

A data frame and some metadata about a MAPPER query from http://genome.ufl.edu/mapper. The data frame holds the predicted transcription factor binding sites from MAPPER.

# **Objects from the Class**

Objects can be created by calls of the form new("mapperHits", ...). These are annotated data frames.

## Slots

```
query: character string that provides information on the query generating the object
```

hits: Object of class "data.frame" providing information on the sequence and other properties of each hit.

## Methods

```
query signature(x = "mapperHits"): ...
hits signature(x = "mapperHits"): ...
show signature(object = "mapperHits"): ...
```

## Author(s)

VJ Carey <stvjc@channing.harvard.edu>

#### References

```
http://genome.ufl.edu/mapper/
```

#### See Also

```
readMAPPER.
```

# **Examples**

```
# see readMAPPER
```

query 3

query

Character string that provides information on the query generating the

## **Description**

Function displaying the exact query to the MAPPER RPC interface.

## Usage

```
query(x)
```

#### **Arguments**

Χ

x – mapperHits object returned from readMAPPER

## References

```
http://genome.ufl.edu/mapper/
```

#### See Also

```
readMAPPER.
```

#### **Examples**

```
# see readMAPPER
```

readMAPPER

Retrieve a set of predicted transcription factor binding sites from the

# Description

The MAPPER RPC interface allows you to retrieve a set of predicted transcription factor binding sites from the MAPPER database through a simple HTTP request. This package provides a function to retrieve predicted TFBS from R.

## Usage

```
readMAPPER(stub = paste(urlpath, "db-rpc?", sep = ""), ...)
```

# Arguments

```
stub stub – a string giving the URL handle up to the db-rpc selector (optional)
... all the query parameters (see Details).
```

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#### **Details**

Currently (9/10) the RPC interface to MAPPER defines the following parameters. You can use any of these as keywords (with bindings supplied using "=", e.g., gene="CRP") to the readMAPPER function

gene - you can use either NCBI GeneID or mRNA accession number; gene names should work too, but this is not recommended because there often are multiple forms of the same gene. [required, unless 'list' is specified]

models - a comma-separated list of model names (to restrict the output to these models only) [required, unless 'list' is specified]

score - the score threshold (0 by default)

perc - the percentile (one of 50, 80, 85, 90, 95). Only hits with scores above the desired percentile for each model are returned.

eval - the E-value threshold (25 by default)

pbases - how many basepairs to look at (default: 2000). See pstart.

pstart - what pbases is relative to (either T for transcript start or C for coding sequence start - remember that in the db we have hits for the region from 10,000 bp upstream of transcript start to 50bp after coding sequence start)

sort - how to sort the results: either M (by model accession), N (factor name), E (by E-value), S (by score), P (by position, the default)

org - two letter organism code

list - returns a list of all TF names with the corresponding model accession numbers. All other parameters are ignored.

If you issue rmapperHelp(), you will get a help page in text.

# Value

An instance of the mapperHits class.

#### Author(s)

Vince Carey <stvjc@channing.harvard.edu>

# References

```
http://genome.ufl.edu/mapper/
```

#### See Also

```
mapperHits-class query hits
```

#### **Examples**

hits(mh)

```
# Run an example retrieving data from the MAPPER RPC interface for gene ID = NM_009696
mh = readMAPPER(gene="NM_009696", perc="95", models="M00027")
# Display the mapperHits object returned by readMAPPER
mh
# Display the exact query that was issued to the MAPPER RPC interface
query(mh)
# Display the details of hits from the query
```

rmapperFactorTable 5

rmapperFactorTable MAPPER Factor Table

# Description

Function to display a table listing the transcription factor/s known to MAPPER with the corresponding model accession numbers.

#### Usage

```
rmapperFactorTable(tf)
```

## **Arguments**

tf

tf – a specific transcription factor, i.e. "AbaA" or "\*" for all transcription factors

#### References

```
http://genome.ufl.edu/mapper/
```

## **Examples**

```
# Display all transcription factors and their model accession numbers rmapperFactorTable("\star") # Display a specific transcription factor and its model accession numbers rmapperFactorTable("AbaA")
```

# Display transcription factors "Ab..." and their model accession numbers rmapperFactorTable("Ab")

rmapperHelp

Help

# Description

Function to call the MAPPER backdoor interface help page.

#### Usage

```
rmapperHelp()
```

# References

```
http://genome.ufl.edu/mapper/
```

## **Examples**

```
# Display the help page
rmapperHelp()
```

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