# harbChIP

March 23, 2012

allhex

utility function: get all hexamers in upstream sequence for an ORF

# Description

utility function: get all hexamers in upstream sequence for an ORF

# Usage

```
allhex(orf, usobj)
```

# Arguments

orf character string, ORF name

usobj upstreamSeqs object

# **Details**

computes Biostrings Views

# Value

computes Biostrings Views

## Author(s)

Vince Carey <stvjc@channing.harvard.edu>

# **Examples**

```
data(sceUpstr)
allhex("YAL001C", sceUpstr)
```

2 chkMotif4TF

buildUpstreamSeqs2 workflow component - build an upstreamSeqs instance from a FASTA read

## **Description**

workflow component - build an upstreamSeqs instance from a FASTA read

## Usage

```
buildUpstreamSeqs2(fastaRead, organism="sce", provenance="harmen")
```

#### **Arguments**

fastaRead results of a readFASTA from Biostrings

organism string naming organism

provenance string or structure describing provenance

#### Details

generates an instance of upstreamSeqs

#### Value

generates an instance of upstreamSeqs

# Author(s)

Vince Carey <stvjc@channing.harvard.edu>

## **Examples**

```
# x = readFASTA(...)
# y = buildUpstreamSeqs2(x)
```

chkMotif4TF

analyze relationship between motif frequency and binding intensity for selected motif and TF

# Description

analyze relationship between motif frequency and binding intensity for selected motif and TF

# Usage

```
chkMotif4TF(motif, TF, chset, upstr, bthresh=2, countthresh=0)
```

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### **Arguments**

motif	character string in alphabet known to Biostrings
TF	name of a TF (sample name in the ChIP-chip data structure chset
chset	an ExpressionSet instance harboring ChIP-chip data
upstr	an instance of upstreamSeqs
bthresh	threshold for binding intensity results to declare TF 'bound' to the upstream region
countthresh	threshold for motif count to be considered 'present' in upstream region

#### **Details**

Uses countPattern to perform motif count.

#### Value

```
a list with elements call, table, and test, the latter providing the result of fisher.test
```

### Author(s)

Vince Carey <stvjc@channing.harvard.edu>

## **Examples**

```
# slow
## Not run:
data(harbChIP)
data(sceUpstr)
chkMotif4TF("CGGCCG", "RDS1", harbChIP, sceUpstr)
## End(Not run)
```

harbChIP

Experimental Data Package: harbChIP

## **Description**

binding ratios and intergenic region data from a ChIP-chip experiment in yeast

# Usage

```
data(harbChIP)
```

## **Format**

The format is: An ExpressionSetObject with covariates:

• txFac: transcription factor symbol from Harbison website CSV file columnnames

## Note

```
derived from web site jura.wi.mit.edu/young_public/regulatory_code/GWLD.
html, binding ratios
```

4 upstreamSeqs-class

#### **Examples**

```
data(harbChIP)
harbChIP
experimentData(harbChIP)
exprs(harbChIP)[1:6,1:7]
dim(exprs(harbChIP))
pData(featureData(harbChIP))[1:6,]
```

sceUpstr

Biostrings representations of S. cerevisiae upstream regions

## Description

Biostrings representations of S. cerevisiae upstream regions

# Usage

```
data(sceUpstr)
```

#### **Details**

environment-based S4 object with DNAstring elements

#### Value

environment-based S4 object with DNAstring elements

#### Author(s)

Vince Carey <stvjc@channing.harvard.edu>

#### **Examples**

```
data(sceUpstr)
sceUpstr
getUpstream("YAL001C", sceUpstr)
```

```
upstreamSeqs-class Class "upstreamSeqs"
```

#### **Description**

Container for a collection of upstream sequences

# **Objects from the Class**

Objects can be created by calls of the form new ("upstreamSeqs", ...). Environments are used to store collections of DNAstrings.

upstreamSeqs-class 5

## **Slots**

```
seqs: Object of class "environment" ~~
chrom: Object of class "environment" ~~
revComp: Object of class "environment" ~~
type: Object of class "environment" ~~
organism: Object of class "character" ~~
provenance: Object of class "ANY" ~~
```

#### Methods

```
Nmers signature(n = "numeric", orf = "character", usobj = "upstreamSeqs"):
    obtain all subsequences of length n as view elements of a DNA string
keys signature(x = "upstreamSeqs"): ...
organism signature(x = "upstreamSeqs"): ...
seqs signature(x = "upstreamSeqs"): ...
show signature(object = "upstreamSeqs"): ...
```

#### Author(s)

Vince Carey <stvjc@channing.harvard.edu>

#### **Examples**

```
showClass("upstreamSeqs")
data(sceUpstr)
sceUpstr
keys(sceUpstr) [1:5]
```

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