

GeneR

April 19, 2009

GeneR

Overview of GeneR package

Description

GeneR packages allow direct use of nucleotide sequences within R software. Functions can be used to read and write sequences from main file formats (Embl, Genbank and Fasta) in order to perform a lot of manipulations and analyses. Main functions are implemented with C extensions.

Details

To summarize, we can split major functions into 4 categories.

I- Reading and writing sequences

GeneR has been designed for fast sequence retrieving even from very large sequence databanks, in Fasta, Embl or Genbank formats. It is also possible to enter sequences directly from a R command line.

There are two ways to store sequences :

- In a C adapted class (buffers) that stores in addition some globals variables, like working strand, size of original sequence and so on.

It is usefull when, for example, we have to work on a subset of a whole chromosome (i.e. a gene). In this case it will be worthwhile to load only the gene in R. Nevertheless, it will remain easy to associate positions on chromosome and positions on gene ...

A complete description on this C class is given in page [globals](#).

- As a character string, the more logical way to store short sequences like "ATGTCGTG". It concerns all functions like "strxxx" (strComp, strMultiExtract etc.).

II- Handling sequences

The usual copy-paste of parts of sequences or other manipulations can be performed by functions using vectors of strands and positions. Annotations from the features field within formatted sequence entries can be extracted and used directly in vectors. By this way, it is easy to extract sequence fragments of interest.

III- Analyzing sequences

Some tools are designed to count oligo-nucleotides, to look for exact word positions or to shuffle sequences (useful for statistical validations).

IV- Genetic tools

Finally, the package also contains functions related to genetic and structural aspects of the sequences : ORF localization, translation, or RNA secondary structure determination (with extension of GeneR: GeneRfold package).

See Also

[globals](#), [readSeq](#), [readEmblLocation](#), [getOrfs](#)

Examples

```
## First of all you can try
demo(GeneR)

## Not run:
demo(hugeGeneR)
## End (Not run)
```

globalSeg

Class manipulating segments

Description

We made a set of tools manipulation segments

Details

The aim of this class is to describe regions on chromosomes that are discontinuous segments on a line like:

```

Region A:      1          10          25          40
              #####          #####
Region B:      #####  #####          #####
Region C:                                #####  #####
```

We made two kind of class

- `segSet`: segments set, is a matrix $n \times 2$ composed of a column of "from", and a column of "to". Used to describe a region like 'A' or 'B' in our example. (A matrix 3×2 to describe region 'B').
- `globalSeg`: a list of `segSet`. It allows the notion of list of discontinuous segments (our use: a list of gene's models as a gene's model is stored as a list of its exons). In our sample, `globalSeg` will be the list of the 3 regions A,B and C. Note that it store more information than just a matrix with 2 columns containing all `segments` of theses 3 regions.

For a better comprehension of other man pages, we introduce this notation:

- a segment is just a part of a line determined by two values (from and to)
- an object of class `segSet` is a set of n segments, determined by a matrix $n \times 2$
- an object of class `globalSeg` is a set of `segSets`, determined by a list of matrix.

Author(s)

Epissage group at CGM.

See Also

See also [as.segSet](#), [as.globalSeg](#), [unionSeg](#)

Examples

```
a = list(
  matrix( c( 1, 15, 17, 5, 45, 38),ncol=2),
  matrix( c( 100 , 120),ncol=2),
  matrix( c( 130, 135, 140, 145),ncol=2),
  matrix( c( 142 , 160),ncol=2))

b = list(
  matrix( c(15, 28, 18, 45),ncol=2),
  matrix( c(1, 15, 25, 10, 20, 40),ncol=2),
  matrix( c(17, 35, 23, 38),ncol=2),
  matrix( c(100, 110, 105, 120),ncol=2))

a = as.globalSeg(a)
b = as.globalSeg(b)

par(mar=c(1,0,1,0))

par(mfrow=c(8,1))
plot(a,xlim=c(1,160),main="A")
plot(b,xlim=c(1,160),main="B")

plot(and(b),xlim=c(1,160),main="and(B)")
plot(or(b),xlim=c(1,160),main="or(B)")
plot(Xor(b),xlim=c(1,160),main="Xor(B)")

plot(a&b,xlim=c(1,160),main="A&B")
plot(a|b,xlim=c(1,160),main="A|B")
plot(Xor(a,b),xlim=c(1,160),main="Xor(A,B)")
```

Globals Variables *Globals variables on Gene sequences*

Description

There are two ways to store sequences in GeneR:

- In a C adapted class (buffers) that stores in addition some globals variables, like working strand, size of original sequence and so on.
It is usefull when, for example, we have to work on a subset of a whole chromosome (i.e. a gene). In this case it will be worthwhile to load only the gene in R. Nevertheless, it will remain easy to associate positions on chromosome and positions on gene ...
- As a character string, the more logical way to store short sequences like "ATGTCGTG". It concerns all functions like "strxxx" ([strComp](#), [strReadFasta](#) etc.).

Details

When GeneR load a subset of a larger sequence stored in a bank file, it will store the following informations in the C adapted class (buffers, by default 100 buffers than can be extended if necessary):

- subsequence (i.e. the succession of A,T,G,C).
- positions of the extremities of the subsequence in the master sequence
- size of the whole sequence in the bank file
- name of the sequence

For specific purposes as renaming a sequence, all these variables can be viewed and carefully changed at any time (here functions [getAccn](#) and [setAccn](#)).

Several sequences can be stored simultaneously and called by their buffer number.

Strand is another global variable which can be set and viewed (functions [getStrand](#) and [setStrand](#)). It is used as input parameter in many functions to analyze complementary strand. It was designed to avoid doing explicitly the complement of the loaded strand then to store it in a buffer with, as consequence, loss of the informations linked to the master sequence.

We have defined 3 types of addresses on a subsequence extracted from a master sequence:

- Absolute addresses i.e. addresses on the master sequence, from the 5' end of the input strand referred as forward (noted A)
- Real addresses, i.e. addresses on the master sequence, from the 5' end of one of strands (noted R)
- Relative addresses, i.e. addresses on working subsequence, from the 5' end of one of strands (noted T).

Let's show an example, if we read sequence from 11 to 20 from a gene of size 40:

```
Strand 0 (Forward strand)
1      11      20      40 Absolute (A)
1      11      20      40 Real (R)
      1      10      Relative (T)
xxxxxxxxxxATGTGTCGTAXxxxxxxxxxxxxxxxxxxxxxxxxxxxx
      10      1      Relative (T)
40     30     21     1 Real (R)
1      11     20     40 Absolute (A)
Strand 1 (Reverse strand)
```

Obviously, when an entire sequence is stored, real and relative addresses will be the same.

Although all functions using positions need and return absolute addresses, 6 functions allow to convert R, A, T into any other type (functions [RtoA](#), [RtoA](#), [RtoA](#), [RtoA](#), [RtoA](#), [RtoA](#)).

A global variable `strand` is used to convert positions (see [setStrand](#) [getStrand](#)).

See Also

[AtoT](#), [AtoR](#), [RtoA](#), [RtoT](#), [TtoA](#), [TtoR](#), [setStrand](#), [getStrand](#), [getParam](#), [setParam](#), [getAccn](#), [setAccn](#)

Examples

```
## Make a dummy sequence
s <- "xxxxxxxxxxATGTGTCGTxxxxxxxxxxxxxxxxxxxxxx"
placeString(s)
writeFasta(file="toto.fa")

indexFasta("toto.fa")
readFasta("toto.fa", from=11, to=20)

getParam()
## $begin
## [1] 11
## $size
## [1] 40
## $strand
## [1] 0
## [...]

## With strand = 0
TtoA(c(1,10))
##[1] 10 19

TtoR(c(1,10))
##[1] 10 19
```

getOrfs

*Gets ORFs from a sequence***Description**

Gets ORFs (Open Reading Frames) from a sequence.

Usage

```
getOrfs(phase = NULL, seqno=0, start="atg",
        stop=c("taa", "tag", "tga"), complete = TRUE, suborfs=TRUE)
maxOrf(seqno=0, phase= NULL, start="atg",
        stop=c("taa", "tag", "tga"), complete = TRUE)
```

Arguments

seqno	Integer/scalar, Sequence number (buffer number)
phase	Integer/scalar, 1,2 or 3, NULL for all three phases
start	string/vector, start codons
stop	string/vector, stop codons
complete	Flag: true returns only complet Orfs, else return all Orfs.
suborfs	Flag: true returns all orfs including subparts of a large orf if it exists "atg" in the phase. False: does not returns sub-orfs.

Value

getOrf returns a table of positions. NULL if no Orfs.
 maxOrf returns the size of the largest Orf, -1 if no Orf.
 All functions return NA if error.

Note

Reverse strand : not implemented

Author(s)

A. Lucas, Emna Marrakchi and Vincent Lefort

Examples

```
s<-"gtcatgcatgctaggtgacagttaaaatgCGTctaggtgacagtctaaca"
placeString(s)

getOrfs(phase = NULL, seqno=0)
maxOrf()

# To get all ORFs on the reverse strand:
sc <- getSeq(0,1)
placeString(sc, seqno=1)
getOrfs(phase = NULL, seqno=1)

# All orfs on both strands :
rbind(getOrfs(seqno=0), getOrfs(seqno=1))
```

AtoR

Conversion of addresses

Description

Converts addresses on a sequence according to the working strand and the address type (A : absolute, T : true, R : relative) .

Usage

```
AtoR(x, seqno=0)
AtoT(x, seqno=0)
RtoA(x, seqno=0)
RtoT(x, seqno=0)
TtoA(x, seqno=0)
TtoR(x, seqno=0)
```

Arguments

x	Integer/vector. Addresses
seqno	Integer/scalar, Sequence number (buffer number)

Details

All details on addresses and global variables are on the page [globals](#).

Value

Integer vector with new addresses

Note

All results depend on the value of Strand. See: [setStrand](#), [getStrand](#).

Author(s)

L. Cottret

See Also

[AtoT](#), [RtoA](#), [RtoT](#), [TtoA](#), [TtoR](#), [setStrand](#), [getStrand](#)

Examples

```
s<-"cgtagtagctagctagctagctagctagc"
placeString (s, seqno=0)
# s of size 30
address <- c(4,20)

# On reverse strand:
setStrand(1)
AtoR(address)
#[1] 26 10

# On forward strand does nothing:
setStrand(0)
AtoR(address)
#[1] 4 20
```

Match

Value Matching

Description

'Match' returns a vector of the positions of (first) matches of its first argument in its second.

Second must be an element of class `segSet` (or a numeric matrix with 2 columns) ordered and without overlapping segments (function `or.segSet` is designed for this purpose).

Usage

```
Match(x, a)
```

Arguments

x a vector of positions
a an element of class segSet

Value

a vector of same size a x. Values are indices corresponding to element 'a', 0 when no segment found.

Note

This can be used for a texture, when we know only regions, described by set of segments.

Author(s)

Antoine Lucas

Examples

```
a = matrix(c(1, 30, 40, 50, 60, 70, 80, 110), ncol=2, byrow=TRUE)
a = or.segSet(a)

## show a:
a

Match(1:40, a)

## texture sample:
b = matrix(c(15, 18, 28, 45,
            1, 10, 15, 20, 25, 40,
            17, 23, 35, 38, 100, 105,
            110, 120), ncol=2, byrow=TRUE)
b = or.segSet(b)

texture = ( Match(1:120, a) > 0 ) + ( Match(1:120, b) > 0 ) * 2

## change numbers to colors
texture <- as.factor(texture)
levels(texture) <- c("red", "blue", "green", "yellow")
texture <- as.character(texture)

plot(1:120, rep(1, 120), col=as.character(texture))
```


Description

computes the eXclusive OR of two objects of class globalSeg a and b, i.e. returns segments which correspond to at least one part of a segment in one set but to nothing in the other set.

When used with only one parameter, Xor(a) returns segments belonging to only one input segment. see the example for a more comprehensive visualisation. .

Usage

```
Xor.globalSeg(a, b = NULL, ...)
```

Arguments

a, b	elements of class globalSeg
...	Unused

Value

An element of class globalSeg

Author(s)

Odile Rogier

See Also

[globalSeg](#), [and.globalSeg](#), [Xor.segSet](#)

Examples

```
a = list(
  matrix( c( 1, 15, 17, 5, 45, 38),ncol=2),
  matrix( c( 100 , 120),ncol=2),
  matrix( c( 130, 135, 140, 145),ncol=2),
  matrix( c( 142 , 160),ncol=2))

b = list(
  matrix( c(15, 28, 18, 45),ncol=2),
  matrix( c(1, 15, 25, 10, 20, 40),ncol=2),
  matrix( c(17, 35, 23, 38),ncol=2),
  matrix( c(100, 110, 105, 120),ncol=2))

a = as.globalSeg(a)
b = as.globalSeg(b)

c = Xor(a,b)
par(mfrow=c(4,1))
plot(a,xlim=c(1,160),main="A")
plot(b,xlim=c(1,160),main="B")
plot(c,xlim=c(1,160),main="Xor(A,B) ")
plot(Xor(b),xlim=c(1,160),main="Xor(B) ")

## show all
c
```

Xor(b)

Xor.segSet *Xor for segments sets*

Description

computes the eXclusive OR of two objects of class segSet a and b, i.e. returns segments which correspond to at least one part of a segment in one set but to nothing in the other set..

XorRecouvr returns segments .which correspond to at least one part of a segment of the envelope but to nothing in the segment set

Usage

```
Xor.segSet(a, b)
xorRecouvr(ranges, envel)
```

Arguments

a, b elements of class segSet, or matrix nx2
 ranges, envel elements of class segSet, or matrices nx2

Value

an element of class segSet.

Author(s)

Antoine Lucas

See Also

[globalSeg](#), [not.globalSeg](#)

Examples

```
a = matrix(c(1, 5, 15, 45, 17, 38,
            100, 120, 130, 140,
            135, 145, 142, 160),
           ncol=2, byrow=TRUE)
b = matrix(c(15, 18, 28, 45,
            1, 10, 15, 20, 25, 40,
            17, 23, 35, 38, 100, 105,
            110, 120), ncol=2, byrow=TRUE)

a <- as.segSet(a)
b <- as.segSet(b)

c = Xor(a, b)
par(mfrow=c(3, 1))
```

```

plot(a,xlim=c(1,160),main="A")
plot(b,xlim=c(1,160),main="B")
plot(c,xlim=c(1,160),main="A Xor B")

## Another sample

a = matrix(c(1,30,40,50,60,70,80,110),ncol=2,byrow=TRUE)
b = matrix(c(1,10,20,30,40,70,80,90,100,110),ncol=2,byrow=TRUE)
a <- as.segSet(a)
b <- as.segSet(b)

c = Xor(a,b)
par(mfrow=c(3,1))
plot(a,xlim=c(1,160),main="A")
plot(b,xlim=c(1,160),main="B")
plot(c,xlim=c(1,160),main="A Xor B")

## Show all
c

```

and.globalSeg *Intersection of global segments*

Description

Computes intersection of two objects of class globalSeg a and b, i.e. returns segments that are both in a and b.

When used with only one object of class globalSeg, it computes intersection of all its segments set (an segments sets is a set of segments; an object of class globalSeg is a set of segments sets). In this case, when global=TRUE, it computes the intersection of all the segments (see example below).

Usage

```
and.globalSeg(a, b = NULL, global = FALSE, byrows = FALSE, relist = TRUE,
  ...)
```

Arguments

a, b	elements of class globalSeg
global	used if b=NULL
byrows	if TRUE, a and b must be lists of same length, and intersection is computed for each elements from a with its homologue in b.
relist	is TRUE, results are re-listed (envelops are union of all a and b).
...	Unused

Value

an element of class globalSeg.

Author(s)

Odile Rogier

See Also[globalSeg, and.segSet](#)**Examples**

```

a = list(
  matrix( c( 1, 15, 17, 5, 45, 38),ncol=2),
  matrix( c( 100 , 120),ncol=2),
  matrix( c( 130, 135, 140, 145),ncol=2),
  matrix( c( 142 , 160),ncol=2))

b = list(
  matrix( c(15, 28, 18, 45),ncol=2),
  matrix( c(1, 15, 25, 10, 20, 40),ncol=2),
  matrix( c(17, 35, 23, 38),ncol=2),
  matrix( c(100, 110, 105, 120),ncol=2))

a = as.globalSeg(a)
b = as.globalSeg(b)

c = and(a,b,byrows=TRUE)
par(mfrow=c(5,1))
plot(a,xlim=c(1,160),main="A")
plot(b,xlim=c(1,160),main="B")
plot(a&b,xlim=c(1,160),main="A&B")
plot(c,xlim=c(1,160),main="A&B, byrow=T")
plot(and(a),xlim=c(1,160),main="and(A) ")

## Show result
a&b
c
and(a)

```

`and.segSet`*Intersection of segments*

Description

`and(a,b)` Computes intersection of two objects of class `segSet` `a` and `b`, i.e. returns segments that are both in `a` and `b`.

When used with only one object of class `segSet` it returns the common segment of all segments of element 'a'.

Usage

```
and.segSet(a, b = NULL, ...)
```

Arguments

a, b	elements of class segSet
...	Unused

Value

an element of class segSet.

Author(s)

Antoine Lucas

See Also

[globalSeg](#), [and.globalSeg](#)

Examples

```

a = matrix(c(1, 5, 15, 45, 17, 38,
            100, 120, 130, 140,
            135, 145, 142, 160),
           ncol=2, byrow=TRUE)
b = matrix(c(15, 18, 28, 45,
            1, 10, 15, 20, 25, 40,
            17, 23, 35, 38, 100, 105,
            110, 120), ncol=2, byrow=TRUE)

a <- as.segSet(a)
b <- as.segSet(b)

c = and(a, b)
par(mfrow=c(3, 1))
plot(a, xlim=c(1, 160))
plot(b, xlim=c(1, 160))
plot(c, xlim=c(1, 160))

## Another sample

a = matrix(c(1, 30, 40, 50, 60, 70, 80, 110), ncol=2, byrow=TRUE)
b = matrix(c(1, 10, 20, 30, 40, 70, 80, 90, 100, 110), ncol=2, byrow=TRUE)
a <- as.segSet(a)
b <- as.segSet(b)

c = and(a, b)
par(mfrow=c(3, 1))
plot(a, xlim=c(1, 160), main="A")
plot(b, xlim=c(1, 160), main="B")
plot(c, xlim=c(1, 160), main="A&B")

## Show result
c
and(a)

```

appendSeq	<i>Appends two sequences</i>
-----------	------------------------------

Description

appends a sequence at the end of another one

Usage

```
appendSeq(destSeqno=0, seqno=1)
```

Arguments

destSeqno	Sequence number of sequence to be appended
seqno	Sequence number of sequence to add at the end of destSeqno

Value

0: no error; NULL if error and a warning if problem in memory allocation.

Note

This function frees the reverse complementary of sequence destSeqno

Author(s)

Antoine Lucas

See Also

[getSeq](#), [concat](#), [assemble](#)

Examples

```
s <- "CTGCGTTTGAAAA"  
placeString(s, seqno=0)  
placeString(s, seqno=1)  
appendSeq(0, 1)  
getSeq(0)
```

as.globalSeg *GlobalSeg manipulation*

Description

These functions implements standards routines for globalSeg manipulations.

Usage

```
as.globalSeg(segments)
as.matrix.globalSeg(x, ...)
range.globalSeg(globalSeg, na.rm = FALSE, global=FALSE, ...)
```

Arguments

x, globalSeg	Element of class globalSeg
segments	Element of class segments
na.rm	logical, indicating if 'NA's should be omitted.
global	Flag, TRUE: range return one value for the whole list; FALSE: range return one value by element of the list.
...	Additional parameters

Value

an element of class globalSeg.

Author(s)

Antoine Lucas

See Also

[globalSeg](#)

as.segSet *Segments Set manipulation*

Description

These functions implements standards routines for segSet manipulations.

Usage

```
as.segSet(x)
as.matrix.segSet(x, ...)
as.data.frame.segSet(x, row.names = NULL, optional = FALSE, ...)
plot.segSet(x, ...)
```

Arguments

<code>x</code>	Element of class <code>segSet</code>
<code>row.names</code>	'NULL' or a character vector giving the row names for the data frame. Missing values are not allowed.
<code>optional</code>	logical. If 'TRUE', setting row names and converting column names (to syntactic names) is optional.
<code>...</code>	Additional parameters

Value

an element of class `segSet`.

Author(s)

Antoine Lucas

See Also

[globalSeg](#)

`assemble`

Concatenates parts of a sequence

Description

concatenates parts of a sequence (in any strand) and puts resulting sequence in destination sequence buffer.

Usage

```
assemble(seqno=0, from=1, to=0, strand=getStrand(), destSeqno=1)
```

Arguments

<code>seqno</code>	Integer/scalar, Sequence number (buffer number)
<code>from, to</code>	Integer/vector, Absolute addresses of the begin and the end of the fragments, (1 means the first nucleotide and 0 conventionally the last one; from must not be larger than to)
<code>strand</code>	Integer/vector, Strand (forward: 0, reverse: 1)
<code>destSeqno</code>	Output sequence number

Value

`destSeqno` or -1 if error.

Author(s)

L.Cottret

See Also

[getSeq](#), [concat](#), [appendSeq](#)

Examples

```
s<-"aaaacgtagctagctagctacccccctagctacgtagattttt"
placeString(s)
x<-c(1,21,39)
y<-c(4,28,0)
assemble(from=x,to=y)
assemble(from=x,to=y,strand=c(0,1,0),destSeqno=2)
getSeq(1)
getSeq(2)
```

bankDensityProfile *Computes density profile(s) of a bank of fasta sequences*

Description

Computes profile(s) of user defined quantities from the beginning or the end of sequence fragments. Profile(s) is(are) constituted of bins of equal size with the mean, the standard deviation and the number of valid events for each bin.

Usage

```
bankDensityProfile (file, seqno=0, fun=seqSkew, fileout= NULL, nbin,
                    sizeBin, allSeq=FALSE, fromEnd=FALSE, name = "all",
                    threshold=0, strand=getStrand(), accu=FALSE, case="all")
```

Arguments

file	Integer/scalar, File name of the bank (fasta file)
seqno	Integer/scalar, Sequence number (buffer number)
strand	Integer/scalar, Strand (forward: 0, reverse: 1)
fun	Function, Function to be used (for example seqSkew)
fileout	String/scalar, If not NULL, a file to write results
nbin	Integer/scalar, Number of bins to be created before the origin
sizeBin	Integer/scalar, Size of the bins
allSeq	Logical/scalar, If TRUE, Input sequence is the whole sequence, if FALSE, input is only the half sequence
fromEnd	Logical/scalar, TRUE: Origin is the end of each sequence, if FALSE: Origin is the beginning of each sequence
name	String/vector, Names of sequences in bank file, "all": uses all sequences of the bank
threshold	Integer/scalar, For each bin, maximum number of N tolerated in the sequence to participate to the computation
case	String/scalar, Case of the letters taken into account ("all", "upper", "lower")
accu	Flag, if true, returns sum, sum of square, and count on demanding function; else returns, mean and standard error on mean.

Value

a list of matrices, with the mean(s), the standard deviation(s) and the number of valid sub-fragments in each bin.

Author(s)

Emna Marrakchi and Antoine Lucas

See Also

[densityProfile](#), [bankSummary](#), [GCcontent](#), [seqSkew](#)

Examples

```
## We create 2 banks
for(i in 1:10)
{
  s=randomSeq(n=100)
  placeString(s, seqno=0)
  writeFasta("toto_norm.fa", append=TRUE, name=i)

  s=randomSeq(prob=c(0.3,1,1,1,0)/3.3, n=100)
  placeString(s, seqno=0)
  writeFasta("toto_lowT.fa", append=TRUE, name=i)
}

densNorm <- bankDensityProfile("toto_norm.fa", nbin=10, sizeBin=10, allSeq=TRUE)
densLowT <- bankDensityProfile("toto_lowT.fa", nbin=10, sizeBin=10, allSeq=TRUE)

par(mfrow=c(1,2))
## Plot skew in normal bank
plot(densNorm$skta, main="TA skew Normal bank", ylim=c(-0.8, 0.3))

## Plot skew in low T bank
plot(densLowT$skta, main="TA skew low T bank", ylim=c(-0.8, 0.3))

## Show numbers
densLowT
```

bankSummary

Computes informations on all a bank file

Description

This functions computes informations (with function `compoSeq`, `GCcontent`, of `seqSkew`) on a complete bank file.

It returns a data frame with one line by sequences in the bank file.

Parameter "name" can be use to limit the exploration to only a few sequences.

Usage

```
bankSummary(file, name = NULL, type = "F", fun = compoSeq, seqno = 0)
```

Arguments

file	Integer/scalar, File name of the bank (fasta file)
seqno	Integer/scalar, Sequence number (buffer number)
name	String/vector, Names of sequences in bank file, NULL: uses all sequences of the bank
type	String/scalar, Bank format ("F" -> fasta. "E" -> embl, "G" -> GenBank)
fun	Function, Function to be used (for example seqSkew)

Value

a data frame with one line by sequence.

Author(s)

Antoine Lucas

See Also

[bankDensityProfile](#)

Examples

```
for(i in 1:8)
{
  s=randomSeq(n=100)
  placeString(s, seqno=0)
  writeFasta("toto_norm.fa", append=TRUE, name=i)
}
bankSummary(file="toto_norm.fa")
bankSummary(file="toto_norm.fa", fun=seqSkew)
```

CompoSeq

Composition in mono, di or trinucleotides of a sequence

Description

Gives composition in mono, di or trinucleotides of fragments of a sequence

Only A, T, G, C are counted, letter as U, X, S, M, N, are counted as 'X'. Flag "case" specify weather we take into account letters in lower, upper or all case.

Usage

```
compoSeq(seqno=0, from=1, to=0, strand=getStrand(),
         wsize=1, p=TRUE, case="all", step=wsize)
strCompoSeq(s, wsize=1, p=TRUE, case="all", step=wsize)
```

Arguments

seqno	Integer/scalar, Sequence number (buffer number)
s	String/scalar, A sequence as character string
wsize	Integer/scalar, size of k-uples.
from,to	Integer/scalar, Absolute addresses of the beginning and the end of the fragment, (1 means the first nucleotide and 0 conventionally the last one; from must not be larger than to and both vectors must be the same size)
strand	Integer/vector, Strand (forward: 0, reverse: 1)
p	Logical/scalar, If p=TRUE, results are in pourcentages, else, results are in counts.
case	String/scalar, Case of the letters taken into account ("all", "upper", "lower")
step	Integer/scalar, Step increment of positions when counting words.

Value

A matrix with nucleotide composition for all regions (NxM where N = 5*wordSize and M = length(from))

If error: NULL

Note

.Use setStrand by default

See Also

[exactWord](#)

Examples

```
s<-"CGTACGTAGTAGCTAGCTAGCTAGCTAGCTAGCTGATCGATGCTAGCTGATCGATGCT"
placeString(s)
x<-c(1,8,15,50)
y<-c(5,12,19,54)
compoSeq(from =x, to = y,wsize=2)

compoSeq(from =x, to = y,wsize=2,strand=1)

compoSeq(from =x, to = y,wsize=2,step=2,p=FALSE)
```

Concat

Concatenation of two sequences

Description

Concatenates a fragment of one sequence with a fragment of another one.

Usage

```
concat(seqno1=0, seqno2=1, destSeqno=2, from1=1, to1=0, strand1=getStrand(),
from2=1, to2=0, strand2=getStrand())
```

Arguments

seqno1	Integer/scalar, Integer/scalar, First sequence number (buffer number)
seqno2	Integer/scalar, Integer/scalar, Second sequence number (buffer number)
destSeqno	Integer/scalar, Output sequence number (buffer number)
strand1	Integer/scalar, Strand of the first fragment (forward: 0, reverse: 1)
from1, to1	Integer/scalar, Absolute addresses of the beginning and the end of the first fragment, (1 means the first nucleotide and 0 conventionally the last one; from must not be larger than to and both vectors must be the same size).
from2, to2	Integer/scalar, Absolute addresses of the beginning and the end of the second fragment, (1 means the first nucleotide and 0 conventionally the last one; from must not be larger than to and both vectors must be the same size)
strand2	Integer/scalar, Strand of the second fragment (forward: 0, reverse: 1)

Value

destSeq or -1 if error

Author(s)

L.Cottret

See Also

[assemble](#), [getSeq](#), [appendSeq](#)

Examples

```
s1<- "aaacgctagcgcg"  
placeString(s1)  
s2<- "ttttctatcag"  
placeString(s2,1)  
concat(seqno1=0, seqno2=1, from1=2, to1=3, from2=8, to2=0, strand1=1)  
getSeq(2)  
#[1] "TTTCAG"
```

deleteCR

Delete carriage return in file

Description

Windows and Mac files does not use same terminating newline. This function convert file to Unix standard with only '\n' char for terminating newline.

Usage

```
deleteCR(file)
```

Arguments

file file name of a file

Value

0 if error, else 1.

Examples

```
seqNcbi("BY608190", file="BY608190.gbk", type="G")
deleteCR("BY608190.gbk")
```

densityProfile *Density profiles*

Description

Computes profile(s) of user defined quantities around sites of interest in sequence fragments. Profile(s) is(are) constituted of bins of equal size around the sites of interest named origins. It produces for each bin, and for each quantity the mean, the standard deviation and the number of valid events.

Usage

```
densityProfile(ori, from, to, seqno = 0, fun = seqSkew,
               fileout = NULL, nbinL, nbinR, sizeBin,
               threshold=0, strand=getStrand(), accu=FALSE, case="all")
plot.profile(x, ylim=NULL, ...)
```

Arguments

ori	Integer/vector, Absolute address of the origins in each fragment
from, to	Integer/vector, Absolute addresses of the beginning and the end of each fragment, (1 means the first nucleotide and 0 conventionally the last one; from must not be larger than to and both vectors must be the same size)
seqno	Integer/scalar, Strand (forward: 0, reverse: 1)
fun	Function, function to be used (by example seqSkew)
fileout	String/scalar, if not NULL, a file name to write results
nbinL	Integer/scalar, number of bins to create before the origin
nbinR	Integer/scalar, number of bins to create after the origin
sizeBin	Integer/scalar, size of the bins
threshold	Integer/scalar, For each bin, maximum number of N tolerated in the sequence to participate to the computation
strand	strand
accu	Flag, if true, returns sum, sum of square, and count on demanding function; else returns, mean and standard error on mean.
case	String/scalar, Case of the letters taken into account ("all", "upper", "lower")

x	An element of class profile
ylim	Range of y axis limits
...	Graphical parameters can be given as arguments to 'plot'.

Value

a list of matrices, with the mean(s), the standard deviation(s) and the number of valid sub-fragments in each bin.

Author(s)

Emna Marrakchi and Antoine Lucas

See Also

[bankDensityProfile](#), [GCcontent](#), [seqSkew](#)

Examples

```
s <- ""
for(i in 1:10)
  s <- paste(s, randomSeq(n=100), randomSeq(prob=c(0.3,1,1,1,0)/3.3,n=100), sep="")

placeString(s, seqno=0)

dens <- densityProfile(ori=200*(1:10)-100, from=200*(0:9)+1, to=200*(1:10),
  seqno=0, fun=seqSkew, nbinL=10, nbinR=10, sizeBin=10)

plot(dens$skta, main="TA skew")

## Example with flagged 'N'

## We create a sequence with a bias every 100 nucleotides
s <- ""
for(i in 1:10)
  s <- paste(s, randomSeq(n=100), randomSeq(prob=c(0.3,1,1,1,0.2)/3.5,n=100), sep="")

placeString(s, seqno=1)

dens2 <- densityProfile(ori=200*(1:10)-100, from=200*(0:9)+1, to=200*(1:10),
  seqno=1, fun=compoSeq, nbinL=10, nbinR=10, sizeBin=10)

plot(dens2$T, main="#T")

## The same but more permissive (allow 4 N in each bin)

dens3 <- densityProfile(ori=200*(1:10)-100, from=200*(0:9)+1, to=200*(1:10),
  seqno=1, fun=compoSeq, nbinL=10, nbinR=10, sizeBin=10, threshold=4)

plot(dens3$T, main="#T")

## Show numbers
dens

dens2
```

dens3

DnaToRna

DNA <-> RNA sequence rewriting

Description

Transformation from T to U in a sequence (dnaToRna), and vice versa (rnaToDna).

Usage

```
dnaToRna(seqno=0, from=1, to=0)
```

```
rnaToDna(seqno=0, from=1, to=0)
```

Arguments

seqno Integer/scalar, Sequence number (buffer number)

from, to Integer/scalar, Absolute addresses of the beginning and the end of the fragment, (1 means the first nucleotide and 0 conventionally the last one; from must not be larger than to and both vectors must be the same size)

Value

seqno or -1 if error

Author(s)

L.Cottret

Examples

```
s<-"atuuutututu"  
placeString(s)  
dnaToRna()  
getSeq()  
#[1] "auuuuuuuuuu"  
rnaToDna()  
getSeq()  
#[1] "atuuuuuuuuu"
```

exactWord	<i>Exact matches of an oligomer</i>
-----------	-------------------------------------

Description

Gets match positions of an oligomer in fragments of a sequence

Usage

```
exactWord(word, seqno=0, from=1, to=0, strand = getStrand(), step=1,
          overlap=TRUE, wNbOcc =-1, case.sensitive=FALSE)
```

Arguments

word	string/scalar, Oligomer sequence
seqno	Integer/scalar, Sequence number (buffer number)
from, to	Integer/scalar, Absolute addresses of the beginning and the end of the fragment, (1 means the first nucleotide and 0 conventionally the last one; from must not be larger than to and both vectors must be the same size)
step	Integer/scalar, Size of the step in which search progression is done
overlap	TRUE: look for overlapping oligomer
wNbOcc	Integer/scalar, Maximum number of occurrences to retrieve sequentially in each fragment (-1 -> all)
strand	Integer/scalar, Strand (forward: 0, reverse: 1)
case.sensitive	T -> difference between lower and upper case letter, F -> no difference

Value

A list of match positions in each fragment.

T -> difference between lower and upper case letter, F -> no difference

If error : NULL

T -> difference between lower and upper case letter, F -> no difference

Note

step with negative values will be implemented soon

Author(s)

L.Cottret

See Also

[getSeq](#)

Examples

```

s<-"cgtagctagctagctagctagctagctagctagctagcta"
placeString(s)
exactWord(word="ag", from=c(3,11,23),to=c(9,17,29))
#[[1]]
#[1] 4 8
#
#[[2]]
#[1] 12 16
#
#[[3]]
#[1] 24 28

placeString("TTTTTTTTTTTT")
exactWord("TTT")
exactWord("TTT", overlap=FALSE, step=2)

```

fastaDescription *Description field reading of a fasta sequence*

Description

Reads the description field of a fasta sequence

Usage

```
fastaDescription(file, name="")
```

Arguments

file	String/scalar, File name of the fasta sequences bank
name	String/scalar, Name of the sequence whose description must be retrieved, (if not specified : first sequence of the file)

Value

The description field (the remaining of the heading line after the first space). if error : NULL

Author(s)

A. Lucas

Examples

```

seqNcbi("BY608190", file="BY608190.fa")

fastaDescription(file="BY608190.fa", name="gi|26943372|gb|BY608190.1|BY608190")

```

Description

nSeq returns the limit of buffers to use, or set this limit to maxBuffers when specified.
freeSeq free sequence seqno and complementary. Free all sequences for freeAllSeq

Usage

```
freeSeq(seqno=0)
freeAllSeq()
nSeq(maxBuffers=NULL)
```

Arguments

seqno	Integer, sequence number to free. (bufseq)
maxBuffers	Integer/Scalar, number of buffers

Value

seqno; 0 for freeAllSeq. Number of sequences for nSeq

Author(s)

A.Viari, L Cotteret, A Lucas

Examples

```
placeString("ATGAGTGATGAGATGATGAG", seqno=0)
placeString("ATGAGTGATGAGATGATGAG", seqno=3)
placeString("ATGAGTGATGAGATGATGAG", seqno=4)
revComp()
revComp(seqno=3)

sizeSeq()
## show size used in all buffers
.seqSize()

## freeSeq(3)
## free buffer 3
.seqSize()

freeAllSeq()
## free all buffers
.seqSize()

## show number of available buffers:
nSeq()
# 100
```

```
## increase number of available buffers
nSeq(150)

.seqSize()
```

getAccn

Reading of a GeneR global variable

Description

Each of these functions returns GeneR global variable associated to a sequence buffer (see [globals](#)), but the global variable `strand` which concerns all the sequence buffers.

Usage

```
getAccn(seqno=0)
getBegSeq(seqno=0)
getSeqSize(seqno=0)
getStrand()
getEndSeq(seqno=0)
```

Arguments

`seqno` Integer/scalar, Sequence number (buffer number)

Details

All details on addresses and GeneR global variables can be found on page [globals](#).

Value

`getAccno` Accession number or name of the sequence
`getBegSeq`, `getEndSeq`, `getSeqSize`
 Beginning or ending position or size of the loaded sequence
`getStrand` Running working strand (0 -> forward, 1 -> reverse)
`if error` : -1

Author(s)

L. Cottret

See Also

[setAccn](#), [getParam](#), [setParam](#), [globals](#)

`getParam`*Reading of all the GeneR global variables*

Description

Gives all the GeneR global variables associated to a sequence

Usage

```
getParam(seqno=0)
```

Arguments

`seqno` Integer/scalar, Sequence number (buffer number)

Details

All details on addresses and global variables can be found on page [globals](#).

Value

A list with

<code>Strand</code>	Running working strand (0 -> forward, 1 -> reverse)
<code>begin</code>	beginning position of the loaded sequence
<code>end</code>	ending position of the loaded sequence
<code>size</code>	size of the parent sequence (to allow computations of relative adresses)
<code>name</code>	name of the loaded sequence

Author(s)

L. Cottret

See Also

[setParam](#)

`getSeq`*Sequence fragments extraction*

Description

Extracts sequence fragments. `getSeq` further converts the fragments in character strings from the GeneR sequence buffer.

Usage

```
getSeq(seqno=0, strand = getStrand(), from=1, to=0)
```

Arguments

seqno	Integer/scalar, Sequence number (buffer number)
from, to	Integer/scalar, Absolute addresses of the begin and the end of the fragment, (1 means the first nucleotide and 0 conventionally the last one; from must not be larger than to and both vectors must be the same size)
strand	Integer/scalar, Strand (forward: 0, reverse: 1)

Value

A vector of character strings. if error : NULL

Author(s)

L.Cottret

See Also

[assemble](#), [concat](#), [appendSeq](#), and for character string manipulation: [substr](#)

Examples

```
s<-"cgtagtagctagctagctagctagctag"
placeString(s, seqno=1)
getSeq(1, from=c(1, 5, 10), to=c(5, 10, 15))
#[1] "CGTAG" "GTAGCT" "TAGCTA"

# And on the reverse:
setStrand(1)
getSeq(1, from=c(1, 5, 10), to=c(5, 10, 15))

## The reverse complement
getSeq(1, strand=1)
```

indexFasta

Index file creation for a sequences bank

Description

These functions create an index file for retrieving quickly a sequence into a fasta, genbank or embl sequence bank file.

Usage

```
indexFasta(file)
indexEmbl(file, index="ix")
indexGbk(file)
```

Arguments

file	String/scalar, File path and name of the sequences file
index	String/scalar, Suffix for the index file

Value

1	Indexfile has been created
-1	Error

Note

These functions create an index file even if it already exists.

Access number larger than 40 characters are skipped (a warning is returned). This can be increasing with variable `MAX_LEN_ACCNO` in file `GeneR_globals.h` (and rebuild GeneR library).

Author(s)

Antoine Lucas

See Also

[makeIndex](#)

insertSeq	<i>Sequence insertion</i>
-----------	---------------------------

Description

Inserts a sequence into another sequence

Usage

```
insertSeq(s, insert, from = 0)
```

Arguments

<code>s</code>	String/scalar, The mother sequence
<code>insert</code>	String/scalar, The sequence to be inserted
<code>from</code>	Integer/scalar, Position of the insertion, 0 = last position, 1 = first

Details

Parameters `insert` and `s` are character string sequences and not GeneR sequences

Author(s)

Antoine Lucas

Examples

```
s<-"gtcatgcgctctaggtcagtca"
insertSeq(s, "aaaaaaaaaaaaaaaa", 7)
```

lowerSeq	<i>Convert upper/lower-case characters in sequence fragments</i>
----------	--

Description

Converts sequence fragments in lower or upper case letters

Usage

```
lowerSeq(seqno=0, from=1, to=0)
upperSeq(seqno=0, from=1, to=0)
```

Arguments

seqno	Integer/scalar, Sequence number (buffer number)
from, to	Integer/scalar, Absolute addresses of the begin and the end of the fragment, (1 means the first nucleotide and 0 conventionally the last one; from must not be larger than to and both vectors must be the same size)

Author(s)

Antoine Lucas

See Also

[posMaskedSeq](#)

Examples

```
s<-"aaaacgtagctagctagctacccccctagctacgtagattttt"
placeString(s, upper=FALSE)
x<-c(1, 21, 39)
y<-c(4, 28, 0)

upperSeq(from=x, to=y)

getSeq()
```

makeIndex	<i>Index file creation for a bank file (internal function)</i>
-----------	--

Description

Makes an index for a bank file (if there wasn't).

Usage

```
makeIndex(file, type="E", index="ix")
```


Arguments

file	string/scalar, file name of bank file
type	Sequence format ("E" for Embl, "G", for Genebank, "F" for Fasta)
index	Suffix for the index file (usually: ix)

Details

Checks if index file exists and is newer than bank file. If not, calls one of `indexFasta`, `indexEmbl`, `indexGbk` functions.

Value

1	Index already exists (and no changes)
0	Index successfully created
-1	Error

Note

Index files are in the form:

```
Accno   deb_feature   deb_sequence   length_sequence
```

with one line by sequence. Number of char must be the same for each line (it is used to search a specific access number) but size used for accno is of 40 char by default. This can be change by setting variable `MAX_LEN_ACCNO` in `GeneR_globals.h` file and recompile the library.

Author(s)

Antoine Lucas

See Also

[indexFasta](#), [indexEmbl](#), [indexGbk](#)

Examples

```
seqNcbi("BY608190", file="BY608190.fa")

# Write index file BY608190.fa.ix ...
indexFasta("BY608190.fa")
```

mask	<i>Mask regions on a sequence</i>
------	-----------------------------------

Description

Mask regions delimited by from and to on a sequence buffer. This function delete the reverse complement if exists (should be recomputed).

Usage

```
mask(seqno = 0, from = 1, to = 0, letter = "N")
```

Arguments

seqno	Integer/scalar, Sequence number (buffer number)
from, to	Integer/scalar, Absolute addresses of the beginning and the end of the fragment, (1 means the first nucleotide and 0 conventionally the last one; from must not be larger than to and both vectors must be the same size)
letter	Letter to use for masking

Value

usually 1; NULL if error.

Author(s)

Antoine Lucas

See Also

See also [posMaskedSeq](#), [upperSeq](#), [lowerSeq](#)

Examples

```
s <- "ATGctgTGTTagtacAATGAGTGAGAGATGTGGGTTTAAAAattt"  
placeString(s, upper=FALSE, seqno=0)  
mask(from=c(10, 20), to=c(15, 22))  
getSeq()
```

not.globalSeg	<i>Substraction of globals segments</i>
---------------	---

Description

Compute substraction of two objects of class globalSeg a and b, i.e. return segments from a that or not in b.

When used with only one parameter, not(A) returns the complementary of each elements of A.

Usage

```
not.globalSeg(a, b = NULL)
```

Arguments

a, b elements of class globalSeg

Value

An element of class globalSeg

Author(s)

Odile Rogier

See Also

[globalSeg, and.globalSeg, not.segSet](#)

Examples

```
a = list(
  matrix( c( 1, 15, 17, 5, 45, 38), ncol=2),
  matrix( c( 100 , 120), ncol=2),
  matrix( c( 130, 135, 140, 145), ncol=2),
  matrix( c( 142 , 160), ncol=2))

b = list(
  matrix( c(15, 28, 18, 45), ncol=2),
  matrix( c(1, 15, 25, 10, 20, 40), ncol=2),
  matrix( c(17, 35, 23, 38), ncol=2),
  matrix( c(100, 110, 105, 120), ncol=2))

a = as.globalSeg(a)
b = as.globalSeg(b)

c = not(a,b)
par(mfrow=c(5,1))
plot(a, xlim=c(1,160), main="A")
plot(b, xlim=c(1,160), main="B")
plot(c, xlim=c(1,160), main="A-B")
plot(not(b,a), xlim=c(1,160), main="B-A")
```

```
plot(not(b), xlim=c(1,160), main="not (B) ")

## Show all
c
not(b, a)
not(b)
```

not.segSet

Substraction of segments sets

Description

This function compute subtraction of two objects of class segSet a and b, i.e. return segments from a that or not in b.

Usage

```
not.segSet(a, b)
```

Arguments

a, b elements of class segSet

Value

an element of class segSet.

Author(s)

Antoine Lucas

See Also

[globalSeg](#), [not.globalSeg](#)

Examples

```
a = matrix(c(1, 5, 15, 45, 17, 38,
            100, 120, 130, 140,
            135, 145, 142, 160),
           ncol=2, byrow=TRUE)
b = matrix(c(15, 18, 28, 45,
            1, 10, 15, 20, 25, 40,
            17, 23, 35, 38, 100, 105,
            110, 120), ncol=2, byrow=TRUE)

a <- as.segSet(a)
b <- as.segSet(b)

c = not(a, b)
par(mfrow=c(3,1))
plot(a, xlim=c(1,160))
```

```

plot(b,xlim=c(1,160))
plot(c,xlim=c(1,160))

## Another sample

a = matrix(c(1,30,40,50,60,70,80,110),ncol=2,byrow=TRUE)
b = matrix(c(1,10,20,30,40,70,80,90,100,110),ncol=2,byrow=TRUE)
a <- as.segSet(a)
b <- as.segSet(b)

c = a - b
par(mfrow=c(3,1))
plot(a,xlim=c(1,160),main="A")
plot(b,xlim=c(1,160),main="B")
plot(c,xlim=c(1,160),main="not (A, B) ")

## Show all
c

```

or.globalSeg	<i>Union of global segments</i>
--------------	---------------------------------

Description

This function compute union of two objects of class globalSeg a and b.

When used with only one parameter, or(A) returns the union of A.

Usage

```
or.globalSeg(a, b = NULL, byrows = FALSE, ...)
```

Arguments

a, b	elements of class globalSeg
byrows	if TRUE, a and b must be lists of same length, and intersection is computed for each elements from a with its homologue in b.
...	Unused

Value

An element of class globalSeg

Author(s)

Odile Rogier

See Also

[globalSeg, and.globalSeg, or.segSet](#)

Examples

```

a = list(
  matrix( c( 1, 15, 17, 5, 45, 38),ncol=2),
  matrix( c( 100 , 120),ncol=2),
  matrix( c( 130, 135, 140, 145),ncol=2),
  matrix( c( 142 , 160),ncol=2))

b = list(
  matrix( c(15, 28, 18, 45),ncol=2),
  matrix( c(1, 15, 25, 10, 20, 40),ncol=2),
  matrix( c(17, 35, 23, 38),ncol=2),
  matrix( c(100, 110, 105, 120),ncol=2))

a = as.globalSeg(a)
b = as.globalSeg(b)

c = or(a,b)
par(mfrow=c(4,1))
plot(a,xlim=c(1,160),main="A")
plot(b,xlim=c(1,160),main="B")
plot(c,xlim=c(1,160),main="or(A,B) ")

plot(or(b),xlim=c(1,160),main="or(B) ")

## Show all
c
or(b)

```

or.segSet

Union of 2 segments sets

Description

Makes union of 2 segments sets

Usage

```

or.segSet(a, b = NULL, simplify=TRUE, ...)
unionSeg(a, b = NULL, simplify=TRUE, ...)

```

Arguments

<code>a, b</code>	Elements of class <code>segSet</code>
<code>simplify</code>	TRUE or FALSE. If TRUE, segments are sorted, and duplicated are removed.
<code>...</code>	Unused

Value

an element of class `segment`.

Author(s)

Antoine Lucas

See Also[globalSeg,or.globalSeg](#)**Examples**

```

from <- c(100,75, 1 ,25,150)
to   <- c(110,120,30,50,170)
or(as.segSet(data.frame(from,to)))
## return:
## from, to
## 1, 50
## 75,120
## 150,170

or.segSet(data.frame(from,to),simplify=FALSE)

## Tip to compute intergenic region
## (imagine: from = genes$start, to=genes$stop

x <- or.segSet(data.frame(from,to))
start <- x[,1]
stop <- x[,2]
n <- length(start)
intergenes <-cbind(stop[1:n-1],start[2:n])
intergenes

```

placeString

Sequence loading in a GeneR sequence buffer.

Description

Puts a character string into a GeneR sequence buffer.

Usage

```
placeString(s, seqno=0, upper=TRUE)
```

Arguments

s	Character string to put into a sequence buffer
seqno	Integer/scalar, Sequence number (buffer number)
upper	upper case conversion (TRUE : conversion of the whole sequence in upper case letters, FALSE : sequence is taken as such)

Value

seqno or -1 if error.

Author(s)

L.Cottret

See Also

[getSeq](#)

Examples

```
s<-"cgtagtagctagctagctagctagctag"
placeString (s, seqno=0)

getSeq(seqno=0)

placeString (s, seqno=1, upper=FALSE)

getSeq(seqno=1)
```

plot.globalSeg *Plot an object of class globalSeg*

Description

Draw rectangles on an axis to represent our segments.

Usage

```
plot.globalSeg(x, xlim = range(x, global = TRUE),
  border = "darkblue", col = "darkblue", density = NULL, at = NULL,
  tick = TRUE, xlabels = TRUE, ...)
```

Arguments

x	Object of class GlobalSeg
xlim	(optional) vector of two elements:
col	color(s) to fill or shade the rectangle(s) with. The default 'NULL', or also 'NA' do not fill, i.e., draw transparent rectangles, unless 'density' is specified.
border	color for rectangle border(s). Can also be 'FALSE' to suppress the border, or 'TRUE' in which case 'col' is used.
density	the density of shading lines, in lines per inch. The default value of 'NULL' means that no shading lines are drawn. A zero value of 'density' means no shading lines whereas negative values (and 'NA') suppress shading (and so allow color filling).

at	the points at which tick-marks are to be drawn. Non-finite (infinite, 'NaN' or 'NA') values are omitted. By default, when 'NULL', tickmark locations are computed.
tick	a logical value specifying whether tickmarks should be drawn on x axis
xlabels	this can either be a logical value specifying whether (numerical) annotations are to be made at the tickmarks, or a vector of character strings to be placed at the tickpoints.
...	other graphical parameters can be given as arguments.

Details

Interval class is described on page [globalSeg](#)

Author(s)

Antoine Lucas

See Also

See also: [globalSeg](#)

Examples

```
a = list(
  matrix( c( 1, 15, 17, 5, 45, 38),ncol=2),
  matrix( c( 100 , 120),ncol=2),
  matrix( c( 130, 135, 140, 145),ncol=2),
  matrix( c( 142 , 160),ncol=2))
a = as.globalSeg(a)
plot(a,at=c(a,recursive=TRUE))
```

posMaskedSeq	<i>Position of masked fragments</i>
--------------	-------------------------------------

Description

These functions return the position of masked fragments within a sequence. Masked fragments are identified by lower case letters or by specific characters (for example: 'N').

Usage

```
posMaskedSeq(seqno=0,from = 1, to = 0, max = 10000,type= "lower")
posMaskedSeqFile(file, name = NA, from = 1, to = 0, max = 10000)
```

Arguments

seqno	Integer/scalar, Sequence number (buffer number)
from, to	Integer/vector, Absolute addresses of the begin and the end of the fragments, (1 means the first nucleotide and 0 conventionally the last one; from must not be larger than to)
file	String/scalar, Path and file name of the Fasta sequence bank
name	Name of the sequence (if NA: the first sequence of the bank)
type	type of masked nucleotide i.e. "lower" or any character like "N" (if lower, posMaskedSeq looks for lower case letters; if a character, posMaskedSeq looks for this character)
max	Maximum number of fragments to retrieve (default: 10000)

Value

a matrix of size n,2.
 gives a warning if there is more than max regions.
 return numeric(0) when no regions found.

Note

This function returns by default only 10000 first regions (default parameter max).

Author(s)

Odile Rogier and Antoine Lucas

See Also

See also [mask](#), [upperSeq](#), [lowerSeq](#)

Examples

```
## Make a dummy sequence
s <- "ATGctgTGTtagtacATNNNNNNNNNNNNNNNTGGGTTaAAAattt"
placeString(s, upper=FALSE, seqno=0)
posMaskedSeq(seqno=0, type="upper")
posMaskedSeq(seqno=0, type="lower")

## Not run:
posMaskedSeq(seqno=0, type="lower", max=2)
## End(Not run)
posMaskedSeq(seqno=0, type="N")

writeFasta(file="toto.fa")
posMaskedSeqFile("toto.fa")
```

randomSeq	<i>Create random sequence</i>
-----------	-------------------------------

Description

Function *randomSeq* creates a random sequence from a distribution of nucleotides, of poly-nucleotides. A real composition of nucleotides can be use from function *compoSeq*, with param `p=TRUE`.

ShuffleSeq creates a sequence while assembling at random specific number of each nucleotides (or poly-nucleotides). These number of nucleotide can be provided by function *compoSeq*, with param `p=FALSE`: it is then a re-assemblage of all nucleotides (or tri-nucleotides, or poly-nucleotides) of a real sequence.

Usage

```
randomSeq(prob = c(0.25, 0.25, 0.25, 0.25, 0), letters = c("T", "C",
"A", "G", "N"), n )
shuffleSeq(count, letters=c("T", "C", "A", "G", "N"))
```

Arguments

<code>prob</code>	A vector of probability weights for obtaining the elements of the vector being sampled or a result from <i>compoSeq</i> function (with option <code>p=TRUE</code>).
<code>count</code>	A vector of number of repetitions for each letters (or bi-tri nucleotides, must be of same length as letters) or a result from <i>compoSeq</i> function (with option <code>p=FALSE</code>).
<code>letters</code>	Letters (or bi-tri nucleotides) to be sampled
<code>n</code>	Integer giving the number of items to choose.

Value

A character string (sequence) or NULL.

Author(s)

A. Lucas

See Also

[compoSeq](#)

Examples

```
## Set seed of your choice (not requiered)
set.seed(3)

#### ---- RANDOMSEQ ----
## Create a sequence of size 30, GC rich
randomSeq(prob = c(0.20, 0.30, 0.20, 0.30), letters = c("T", "C", "A", "G"), n = 30)
## [1] "CTGGAACCGAGGGGTTTCATCCCCCAGTGA"
```

```

## use with bi-nucleotides
randomSeq(prob=rep(0.0625,16),letters = c("TT","TC","TA","TG","CT","CC","CA","CG","AT","A
## [1] "CGCATGATCCCAGGCTAACT"

#### ---- SHUFFLESEQ ----
## Create a sequence with 7 T, 3 C and A, and 4 G.
shuffleSeq(count=c(7,3,3,4,0),letters=c("T","C","A","G","N"))
## [1] "TATCTTTTGTCTGGACGA"

## Same with bi-nucleotides
shuffleSeq(count=c(rep(4,4),rep(2,4),rep(1,4),rep(0,4)),letters = c("TT","TC","TA","TG","
## [1] "TCTTTCATTTCCTTCTAGTGTACCCGTATACGTGTCTGTGTACTTCAACAACCTTAT"

## From a real sequence:
seqNcbi("BY608190",file="BY608190.fa")
readFasta("BY608190.fa")

## create a random sequence from a real tri-nucleotides distribution
## Size of sequence will be 10*3.
randomSeq(compoSeq(ysize=3,p=TRUE),n=10)

## re assemble real tri-nucleotides of a real sequence
shuffleSeq(compoSeq(ysize=3,from=1,to=30,p=FALSE))

```

Read location

Get annotations of a GeneBank or an EMBL sequence

Description

Extracts by keywords, from a GeneBank or an EMBL sequence entry, annotations from the features field

Usage

```

readGbkLocation (file, name = NA, from = 1, to = 0, key = "CDS", subkey
                = "")
readEmblLocation (file, name = NA, from = 1, to = 0, key = "CDS", subkey
                 = "")

```

Arguments

file	String/scalar, File name of the bank
name	String/scalar, Sequence name (default : first sequence)
from, to	Integer/scalar, Absolute addresses of the begin and the end of the fragment, (1 means the first nucleotide and 0 conventionally the last one; from must not be larger than to)
key	String/scalar,. Feature to retrieve. (ex: GENE, CDS...)
subkey	String/scalar, Label to retrieve (ex: locus_tag, note, codon_start...).

Value

A data frame

Author(s)

P.Durosay(C), L.Cottret(R), A. Lucas

See Also

[readEmblLocation](#), [readSeq](#)

Examples

```
## Not run:
# Get SARS Sequence
seqNcbi("NC_004718", file="NC_004718.gbk", type="G")

# Get CDS Positions

readGbkLocation(file="NC_004718.gbk")

# Get peptides...
readGbkLocation(file="NC_004718.gbk", key="mat_peptide", subkey="product")
## End(Not run)

# The same with EMBL...
#
# Get SARS Sequence

## Not run:
download.file("http://bioinfo.hku.hk/sars/AY291451.seq",
destfile="AY291451.seq")

# Get CDS Positions
readEmblLocation(file="AY291451.seq")

# Get peptides...
readEmblLocation(file="AY291451.seq", key="mat_peptide", subkey="product")
## End(Not run)
```

readEmblDescript *Read features from an Embl bank file*

Description

Read features from an Embl bank file

Usage

```
readEmblDescript(file, name = NA, code = "DE")
```

Arguments

file String/scalar, File name of the bank (example: "foo.embl")
 name String/scalar, Sequence name (default : first sequence)
 code String/scalar, feature code to look for (i.e. "CC", "DE", "CC", etc...)

Value

A vector of character string.

Author(s)

Antoine Lucas

Examples

```
## Not run:
download.file("http://bioinfo.hku.hk/sars/AY291451.seq",
destfile="AY291451.seq")

readEmblDescript(file = "AY291451.seq",name = "AY291451",code = "DE")

##_returns: "SARS coronavirus TW1, complete genome."

readEmblDescript(file = "AY291451.seq",name = "AY291451",code = "OC")

##_returns: "Viruses; ssRNA positive-strand viruses, no DNA stage; Nidovirales; Coronavi

readEmblDescript(file = "AY291451.seq",name = "AY291451",code = "RA")
##[1] "Yeh S.-H., Kao C.-L., Tsai C.-Y., Liu C.-J., Chen D.-S., Chen P.-J.;"
##[2] "Yeh S.-H., Kao C.-L., Tsai C.-Y., Liu C.-J., Chen D.-S., Chen P.-J.;"

## End(Not run)
```

readSeq

Sequence extraction from a bank

Description

These functions load a sequence from a Fasta, Embl or GeneBank sequence bank file into a GeneR sequence buffer or as a character string..

readSeq is the generic function, readFasta, readGbk and readEmbl are aliases.

Usage

```
readSeq(file,name=NA, type="F", seqno=0,
        from=1, to=0,upper=TRUE, index="ix")
readFasta(file,name=NA, seqno=0,
          from=1, to=0,upper = TRUE, index="ix")
strReadFasta (file, name = NA, from = 1,
              to = 0,upper = TRUE)
readGbk(file,name=NA, seqno=0, from=1, to=0,upper = TRUE,index="ix")
```

```

strReadGbk (file, name = NA, from = 1,
            to = 0, upper = TRUE)
readEmbl (file, name=NA, seqno=0, from=1, to=0, upper = TRUE, index="ix")
strReadEmbl (file, name = NA, from = 1,
             to = 0, upper = TRUE, index = "ix")

```

Arguments

file	String/scalar, File name of the bank
name	String/scalar, Sequence name (default : first sequence)
upper	upper case letters conversion (TRUE: The whole sequence is converted to upper case letters, FALSE : no conversion (only implemented for Fasta sequence)).
type	String/scalar, Bank format ("F" -> fasta, "E" -> embl, "G" -> GenBank)
seqno	Integer/scalar, Sequence number (buffer number)
from, to	Integer/scalar, Absolute addresses of the begin and the end of the fragment, (1 means the first nucleotide and 0 conventionally the last one; from must not be larger than to)
index	String/scalar, type of index ("ix" -> sequence name is the accession number "id" -> sequence name is the identifier)

Value

integer: 1 if OK; -1 if error. strReadFasta, strReadGbk and strReadEmbl returns the sequence as character string.

Author(s)

L.Cottret, A. Lucas.

See Also

[GeneR](#), [globals](#), [readEmblLocation](#), [readGbkLocation](#), [getSeq](#), [sizeSeqEmbl](#), [sizeSeqFasta](#)

Examples

```

## Get a sequence from Ncbi
seqNcbi ("BY608190", file="BY608190.gbk", type="G")

## Read Gbk file to buffer 0
readGbk ("BY608190.gbk")

## Or to a character string
strReadGbk ("BY608190.gbk")

## Write to Fasta file
writeFasta (file='toto.fa')

## Make an index to this file
indexFasta ('toto.fa')
readFasta (file="toto.fa", seqno=0, from=1, to=159)

## Show seqence on buffer 0

```

```

getSeq()

## Make a Fake file
writeEmblLine(file='toto.embl',code='AC',header='tmp',append=FALSE)
writeEmblSeq(file='toto.embl')

## Make index on file toto.embl
indexEmbl('toto.embl')

## Read Embl file to buffer 0
readEmbl('toto.embl')

## Or read "directly"
strReadEmbl('toto.embl')

```

relist

Group segments into global segments

Description

Relist,Relistage : put all segments into its specific envelope.

Usage

```

relist(ranges, envel)
relistage(ranges, envel)

```

Arguments

envel, ranges elements of class segSet or matrix nx2

Value

Relist a vector with envelopes indices (-1 if none found)
Relistage a element of class globalSeg

Note

Envelops must overlap ranges.

Author(s)

Antoine Lucas and Odile Rogier

Examples

```

from = c(1, 15,17,100,130,135,142)
to   = c(40,45,38,120,140,145,160)
envelfrom = c(1,100,130)
envelto = c(45,120,160)
ranges = as.segSet(data.frame(from,to))

```



```
envel = as.segSet(data.frame(envelfrom, envelto))

relist(ranges, envel)
c = relistage(ranges, envel)

par(mfrow=c(3,1))
plot(ranges,xlim=c(1,160),main="ranges")
plot(envel,xlim=c(1,160),main="Envelopes")
plot(c,xlim=c(1,160),main="relist")
```

Reverse complementary

Performs the reverse of a sequence

Description

These functions perform the reverse/complementary of a sequence. RevComp loads further the result into the transparent complementary buffer.

Usage

```
revComp(seqno=0, force=FALSE)
strComp (s)
```

Arguments

seqno	Integer/scalar, Sequence number (buffer number)
force	Logical/scalar, Force flag (FALSE -> revComp creates the reverse/complementary sequence only if it doesn't exist, TRUE -> revComp creates it anyway)
s	A sequence, string format.

Value

seqno or -1 if error.

Author(s)

L.Cottret

See Also

[getSeq](#)

Examples

```
s<-"cgtagtagctagctagctagctagctag"
placeString (s, seqno=1)
getSeq(1)
# return [1] "CGTAGTAGCTAGCTAGCTAGCTAGCTAG"

revComp(1)      # computes the reverse
getSeq(1,1)
# return [1] "CTAGCTAGCTAGCTAGCTAGCTACTACG"

# Or with strComp
strComp (s)
```

seqSkew

compute the strand asymmetries from one sequence fragment

Description

compute the strand asymmetries from one sequence fragment

Usage

```
seqSkew(seqno = 0, from=1,to=0,strand=getStrand(),case = "all")
GCcontent(seqno = 0, from=1,to=0,case = "all",...)
```

Arguments

seqno	Input sequence number. (bufseq)
from,to	Begining and ending of sequence, can be vectors. 0 represent the last nucleotide and 1 the first one.
case	"all"
strand	strand
...	other parameters

Value

a data frame with 4 columns as follow

TA skew or TA pourcent	$(T-A)/(T+A)$ or $\text{count}(TA) / \text{count}(TAGCN)$
GC skew or GC pourcent	$(G-C)/(G+C)$ or $\text{count}(GC) / \text{count}(TAGCN)$
total skew or C pourcent	TA+GC skews or $\text{count}(C) / \text{count}(TAGCN)$

Author(s)

Yves d'Aubenton and Emna

See Also

[densityProfile](#), [bankDensityProfile](#), [compoSeq](#)

Examples

```
s<-"CGTACGTAGTAGCTAGCTAGCTAGCTAGCTAGCTGATCGATGCTAGCTGATCGATGCT"
placeString(s, seqno=0)
x<-c(1, 8, 15, 50)
y<-c(5, 12, 19, 54)
seqSkew(seqno=0, from=x, to=y, strand=0)

seqSkew(seqno=0, from=x, to=y, strand=1)

GCcontent(seqno=0, from=x, to=y)
```

seqSrs	<i>Download a sequence in Fasta / Genbank / Embl format from Ncbi or Srs</i>
--------	--

Description

Get a sequence in Fasta / Genbank / Embl format trough web with only an accno
seqUrl will get the sequence through a srs web program. seqNcbi will do the same with ncbi web server (I prefer this one).

Usage

```
seqSrs (accno, file="toto.seq", submotif=FALSE,
srs="http://srs.ebi.ac.uk/srsbin/cgi-bin/wgetz",
bank=c("EMBL", "REFSEQ"))
seqNcbi (accno, file="toto.seq", submotif=FALSE, type="fasta")
```

Arguments

accno	Access Number: "dbj BY608190.1 BY608190" or "BY608190"
file	file where seqence will be downloaded
submotif	a logical value indicating whether we look for a subpatern of accno
srs	a url srs web program
type	fasta or genbank for seqNcbi (file format of sequence).
bank	List of banks to search into.

Value

1 if file has been correctly created. A file containing the sequence in file format requested

Note

SeqNcbi returns sometimes Genbank file in a not very valid format (specially with EST sequences). These files will not be computed by readseq. (Use Fasta format, then).

Author(s)

Antoine Lucas, Centre de Genetique moleculaire, CNRS Gif / Yvette

Examples

```
seqNcbi("BY608190", file="BY608190.fa")

## Not run:
# idem:
seqSrs("dbj|BY608190.1|BY608190", file="BY608190.embl", submotif=TRUE, type="embl")

seqSrs("AK129232", type="embl", srs="http://www.infobiogen.fr/srs71bin/cgi-bin/wgetz")
## End(Not run)
```

SetAccn

Set globals variables of a sequence

Description

Globals variables concerning a sequence (see [globals](#)) can be directly set with theses functions. Global variable `strand` concerns all buffers of sequence.

Usage

```
setAccn(name, seqno=0)
setBegSeq(pos=1, seqno=0)
setSeqSize(pos, seqno=0)

setStrand(str=0)
```

Arguments

<code>name</code>	Character: name of sequence
<code>seqno</code>	Integer: sequence number (bufno)
<code>pos</code>	Integer: size, or position.
<code>str</code>	Strand: 0 : watson (forward) 1 : crick (reverse)

Details

All details on addresses and globals variables are on page [globals](#).

Value

The name, or "error" if error.

Author(s)

L. Cottret

See Also

[globals](#), [getAccn](#), [setParam](#), [getParam](#)

SetParam

Set global variables associated to a sequence

Description

Set global variables associated to a sequence

Usage

```
setParam(from = 1, sizeMaster = seqSize(), seqno = 0)
```

Arguments

from	Integer, begin of sequence.
sizeMaster	Integer, size of origin sequence.
seqno	Integer, sequence number (bufseq)

Value

-1 if error or the list.

Author(s)

L. Cottret

See Also

[globals](#), [getParam](#), [setAccn](#), [getAccn](#)

size.globalSeg

Size.globalSeg

Description

Tools manipulating size,max,min,length of elements of class globalSeg

Usage

```
size.globalSeg(a)
Max.globalSeg(a)
Min.globalSeg(a)
Length.globalSeg(a, global=TRUE)
```

Arguments

a elements of class globalSeg
 global if TRUE: return one value (number segments) if FALSE return a vector (number of segments in each elements of the list)

Value

return a vector with min, max, size or length of each element of the list (a element of class globalSeg is a list of elements of class segments).

Author(s)

Odile Rogier

Examples

```
a = list(
  matrix( c( 1, 15, 17, 5, 45, 38),ncol=2),
  matrix( c( 100 , 120),ncol=2),
  matrix( c( 130, 135, 140, 145),ncol=2),
  matrix( c( 142 , 160),ncol=2))
a = as.globalSeg(a)

Length(a)
Length(a,global=TRUE)
size(a)
Max(a)
Min(a)
```

 SizeSeq

Size of a sequence in a buffer

Description

Get size (number of characters) of a sequence. .seqSize returns size allocated in all buffers.

Usage

```
sizeSeq(seqno=0)
.seqSize()
```

Arguments

seqno Integer: sequence number (bufseq)

Value

Integer: size of sequence. A list of two vectors for .seqSize.

Author(s)

L.Cottret

See Also[sizeSeqEmbl](#), [sizeSeqFasta](#)**Examples**

```
freeAllSeq()
s<-"cgtagtagctagctagctagctagctag"
placeString(s, seqno=1)
sizeSeq(1)

.seqSize()
```

SizeSeqFasta*Size of a sequence from a fasta/embl/gbk file*

Description

Give the size of a sequence from a fasta/embl/gbk file.

Usage

```
sizeSeqFasta(file, name=NA)
sizeSeqEmbl(file, name=NA, index="ix")
sizeSeqGbk(file, name=NA)
```

Arguments

file	File name of the bank
name	Name of the sequence in the bank (default: first sequence)
index	character. "ix": Name is accn of sequence. "id" : Le name is id of sequence.

Value

Size of sequence, NULL if error.

Author(s)

P.Durosay(C). L.Cottret(R)

See Also[readSeq](#)

Examples

```

seqNcbi("NC_004718", file="NC_004718.gbk", type="G")
sizeSeqGbk("NC_004718.gbk")
# Or:
sizeSeqGbk("NC_004718.gbk", "NC_004718")

# Idem with fasta format
seqNcbi("NC_004718", file="NC_004718.fa", type="F")
sizeSeqFasta("NC_004718.fa")
# Or:
sizeSeqFasta("NC_004718.fa", "gi|30271926|ref|NC_004718.3|")

## Not run:
# With Embl
download.file("http://bioinfo.hku.hk/sars/AY291451.seq",
destfile="AY291451.seq")
sizeSeqEmbl("AY291451.seq")
## End(Not run)

```

sliceSegment	<i>delineate sub-segments of equal size at both sides of an origin in a segment</i>
--------------	---

Description

delineate sub-segments of equal size at both sides of an origin in a segment

Usage

```
sliceSegment(from = 1, to = 0, ori = 0, nbinL = 0, nbinR = 0, size = 1)
```

Arguments

from	integer, position of the begin of segment
to	integer, position of the end of segment
ori	integer, the position of the origin
nbinL	number of sub-segments to create before the origin
nbinR	number of sub-segments to create after the origin
size	integer, size of the sub-segments

Value

an element of class segSet

Author(s)

Yves d'Aubenton and Emna

See Also

[bankDensityProfile](#), [densityProfile](#), [GCcontent](#), [seqSkew](#)

Examples

```
## If by example we have a gene in positions 150 to 200
## This will create segments to study
## A more complex example is provided with function densityProfile

sliceSegment(from=1,to=200,ori=150,nbinL=15,nbinR=15,size=10)
```

 Translate

Translation from DNA trinucleotides to proteine

Description

Translation tools from DNA trinucleotides to proteine

Usage

```
translate (seqno, from = 1, to = 0, strand = 0, code = 0,
charcode = "")
strTranslate (s, code = 0, charcode = "")
showTable (code = 0, charcode = "")
```

Arguments

seqno	Integer, sequence number (bufno)
s	Sequence as a character string
from,to	Begining and ending of sequence, can be vectors. 0 represent the last nucleotide and 1 the first one.
strand	0: forward, 1: reverse, can be a vector
code	One of the following standard code: 0, standard genetic code; 1 Vertebrate Mitochondrial Code; 2 Yeast Mitochondrial Code; 3 Mold, Protozoan, and Coelenterate Mitochondrial Code and the Mycoplasma/Spiroplasma Code; 4 Invertebrate Mitochondrial Code
charcode	A character string of size 64, like "FFLLSSSSYY**CC*WLLLLPPPHHQ-QRRRRRIIMTTTTNKKSSRRVVVAAAADDEEGGGG" for translation code to use, in the order: TTT TTC TTA TTG TCT TCC TCA ... (for F F L L S S S). Use showTable to be sure!

Value

```
strTranslate return a character string of the protein
translate    return a vector of character string of the protein
showTable    return the table of translation
All return -1 if error.
```

Note

Global value of strand has no effect on this function. (see [globals](#), [getParam](#), [setStrand](#))

Examples

```
s<-"gtcatgcatgcttaggtgacagttaaaatgcgtctaggtgacagtctaaca"

# Simple usage:
strTranslate(s)
#[1] "VMHAR*QLKCV*VTV*Q"

# with buffers
placeString(s)
translate()
# the same...
#[1] "VMHAR*QLKCV*VTV*Q"

# with 2 positions
translate (from=c(1,2),to=c(0,0))
#[1] "VMHAR*QLKCV*VTV*Q" "SCMLGDS*NASR*QSN"

# with 2 strands
translate (from=c(1,2,1),to=c(0,0,0),strand=c(0,0,1))
#[1] "VMHAR*QLKCV*VTV*Q" "SCMLGDS*NASR*QSN" "LLDCHLDAF*LSPSMHD"

# With Invertebrate Mitochondrial Code
translate(code=4)
#[1] "VMHASWQLKCV*VTV*Q"

# With a personal code
translate(charcode="FFLLxxxxYY*CCwwLLLLPPPPHHQRRRRIIMTTTTNNKKSSRRuuuuAAAADDEEGGG")
#[1] "uMHARwQLKCu*uTu*Q"

# Show what is this code...
showTable(charcode="FFLLxxxxYY*CCwwLLLLPPPPHHQRRRRIIMTTTTNNKKSSRRuuuuAAAADDEEGGG")
#      [,1] [,2]
# [1,] "UUU" "F"
# [2,] "UUC" "F"
# [3,] "UUA" "L"
# [4,] "UUG" "L"
# [5,] "UCU" "x"
# [6,] "UCC" "x"
# [7,] "UCA" "x"
# ...

# Show Standard table:
showTable()

# Show Invertebrate Mitochondrial Code
showTable(code=4)
```

Description

Write one or more parts of sequences into Fasta file format.

Usage

```
writeFasta(file = "data.fasta", seqno = 0, from = getBegSeq(seqno),
           to = getEndSeq(seqno), name = getAccn(seqno),
           comment = paste("from", from, "to", to),
           cpl = 60, append = FALSE)
strWriteFasta(s, file="data.fasta", name="Seq_R",
              comment=as.character(nchar(s)),
              cpl=60, append = FALSE)
```

Arguments

seqno	Integer, number of the sequence. (bufseq)
s	A character string containing the sequence
from, to	Begining and ending of sub-sequences to extract. Can be vectors. 0 represent the last nucleotide and 1 the first one.
name	Sequence name.
comment	Comment.
file	Output File.
cpl	Number of caracteres by line.
append	Scalar boolean. TRUE -> sequence is added at the end of the file. FALSE -> file is written over.

Value

1 or -1 if error.

Author(s)

L.Cottret

Examples

```
s<-"cgtagctagctagctagctagctacgtagctagctgactgtcgat"
placeString(s)
from <- c(1,14)
to <- c(10,0)
writeFasta(file="bank.fasta",from =from, to=to, append=FALSE)

strWriteFasta(s="ATGTCGTGGTaaattaatTTGGTCCC", file="bank.fasta", append=TRUE)

## Show file
cat(paste(readLines("bank.fasta"), collapse='\n'))
```

WriteEmblSeq

*Write to Embl file***Description**

Write a sequence, with description into a EMBL file. writeEmblSeq write the sequence; writeEmblLine write a feature line, writeEmblComment write a comment line. CompleteStringWithSpace: internal function

Usage

```
writeEmblSeq (file, seqno = 0)
writeEmblLine (file, code = "", header = "", text = "",
               nextfield = TRUE, append=TRUE)
writeEmblComment(file, code = "", text = "",
                 nextfield = TRUE, append=TRUE)
```

Arguments

file	EMBL File name
code	2 letters: to be written at line beginning
header	First part of line
text	a text to be written
seqno	Integer, sequence number (bufno)
nextfield	TRUE: write XX after line, FALSE: don't write XX.
append	Scalar boolean. TRUE -> line(s) is added at the end of the file. FALSE -> file is written over.

Author(s)

A. Lucas and Vincent Lefort

Examples

```
s<-"gtcatgcatgctaggtgacagttaaaatgctgctaggtgacagtctaaca"
placeString(s)

# Add lines with "CC  bla bla bla" and a line "XX"
writeEmblComment(file="toto.embl",code="CC",text="This is a comment for \
this dummy sequence... I try to be long enough to show that this comment \
will be written on several lines",append=FALSE)

# Add a line with "FT  CDS  bla bla bla"
writeEmblLine(file="toto.embl",code="FT",header="CDS",text="<1..12",
              nextfield = FALSE)
# Add lines with "FT      bla bla bla"
writeEmblLine(file="toto.embl",code="FT",header="",text="/codon_start=2",
              nextfield = FALSE)
writeEmblLine(file="toto.embl",code="FT",header="",text="/gene=\"toto\"",
```

```
        nextfield = FALSE)
writeEmblLine(file="toto.embl",code="FT",header="",text="/note=\"Here is \
what I think about this gene\"",nextfield = FALSE)

## Translation
prot <- translate(seqno=0,from=getOrfs()[1,1],to=getOrfs()[1,2])
writeEmblLine (file="toto.embl",code='FT',header='',
text=paste('/translation=',prot , '\\', sep=' '),nextfield =TRUE)

# Add sequence
writeEmblSeq(file="toto.embl")

## Show file
cat(paste(readLines("toto.embl"),collapse='\\n'))
```

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