Biostrings Quick Overview

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Most but not all functions defined in the Biostrings package are summarized here.

Function	Description
length	Return the number of sequences in an object.
names	Return the names of the sequences in an object.
	Extract sequences from an object.
head, tail	Extract the first or last sequences from an object.
rev	Reverse the order of the sequences in an object.
С	Combine in a single object the sequences from 2 or more objects.
width, nchar	Return the sizes (i.e. number of letters) of all the sequences in an
	object.
==, !=	Element-wise comparison of the sequences in 2 objects.
match, %in%	Analog to match and %in% on character vectors.
duplicated, unique	Analog to duplicated and unique on character vectors.
sort, order	Analog to sort and order on character vectors, except that the ordering of DNA or Amino Acid sequences doesn't depend on the locale.
relist, split, extractList	Analog to relist and split on character vectors, except that the result is a <i>DNAStringSetList</i> or <i>AAStringSetList</i> object. extractList is a generalization of relist and split that supports arbitrary groupings.

Table 1: Low-level manipulation of DNAStringSet and AAStringSet objects.

Function	Description
alphabetFrequency	Tabulate the letters (all the letters in the alphabet for alphabet-
letterFrequency	Frequency, only the specified letters for letterFrequency) in a
	sequence or set of sequences.
uniqueLetters	Extract the unique letters from a sequence or set of sequences.
letterFrequencyInSlidingView	Specialized version of letterFrequency that tallies the requested
	letter frequencies for a fixed-width view that is conceptually slid
	along the input sequence.
consensusMatrix	Computes the consensus matrix of a set of sequences.
dinucleotideFrequency	Fast 2-mer, 3-mer, and k-mer counting for DNA or RNA.
trinucleotideFrequency	
${\tt oligonucleotideFrequency}$	
nucleotideFrequencyAt	Tallies the short sequences formed by extracting the nucleotides
	found at a set of fixed positions from each sequence of a set of
	DNA or RNA sequences.

Table 2: Counting / tabulating.

Function	Description
reverse	Compute the reverse, complement, or reverse-complement, of a set
complement	of DNA sequences.
reverseComplement	
translate	Translate a set of DNA sequences into a set of Amino Acid se-
	quences.
chartr	Replace letters in a sequence or set of sequences.
replaceAmbiguities	
subseq, subseq<-	Extract/replace arbitrary substrings from/in a string or set of
extractAt, replaceAt	strings.
replaceLetterAt	Replace the letters specified by a set of positions by new letters.
padAndClip, stackStrings	Pad and clip strings.
strsplit, unstrsplit	strsplit splits the sequences in a set of sequences according to a
	pattern. unstrsplit is the reverse operation i.e. a fast implemen-
	tation of sapply(x, paste0, collapse=sep) for collapsing the
	list elements of a $DNAStringSetList$ or $AAStringSetList$ object.

Table 3: Sequence transformation and editing.

Function	Description
matchPattern	Find/count all the occurrences of a given pattern (typically short)
countPattern	in a reference sequence (typically long). Support mismatches and
	indels.
vmatchPattern	Find/count all the occurrences of a given pattern (typically short)
vcountPattern	in a set of reference sequences. Support mismatches and indels.
matchPDict	Find/count all the occurrences of a set of patterns in a reference
countPDict	sequence. (whichPDict only identifies which patterns in the set
whichPDict	have at least one match.) Support a small number of mismatches.
vmatchPDict	[Note: vmatchPDict not implemented yet.] Find/count all the
vcountPDict	occurrences of a set of patterns in a set of reference sequences.
vwhichPDict	(whichPDict only identifies for each reference sequence which pat-
	terns in the set have at least one match.) Support a small number
	of mismatches.
pairwiseAlignment	Solve (Needleman-Wunsch) global alignment, (Smith-Waterman)
	local alignment, and (ends-free) overlap alignment problems.
matchPWM	Find/count all the occurrences of a Position Weight Matrix in a
countPWM	reference sequence.
trimLRPatterns	Trim left and/or right flanking patterns from sequences.
matchLRPatterns	Find all paired matches in a reference sequence i.e. matches speci-
	fied by a left and a right pattern, and a maximum distance between
	them.
matchProbePair	Find all the amplicons that match a pair of probes in a reference
	sequence.
findPalindromes	Find palindromic regions in a sequence.

Table 4: String matching / alignments.

Function	Description
readBStringSet	Read ordinary/DNA/RNA/Amino Acid sequences from files
readDNAStringSet	(FASTA or FASTQ format).
readRNAStringSet	
readAAStringSet	
writeXStringSet	Write sequences to a file (FASTA or FASTQ format).
writePairwiseAlignments	Write pairwise alignments (as produced by pairwiseAlignment)
	to a file ("pair" format).
readDNAMultipleAlignment	Read multiple alignments from a file (FASTA, "stockholm", or
${\tt readRNAMultipleAlignment}$	"clustal" format).
${\tt readAAMultipleAlignment}$	
write.phylip	Write multiple alignments to a file (Phylip format).

Table 5: I/O functions.

Function	Description
stringDist	Computes the matrix of Levenshtein edit distances, or Hamming
	distances, or pairwise alignment scores, for a set of strings.

Table 6: Miscellaneous.