# 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. |
| C | 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 order- <br> ing 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 re- <br> sult is a DNAStringSetList or AAStringSetList object. ext ract List is <br> a generalization of relist and split that supports arbitrary group- <br> ings. |

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

| Function | Description |
| :--- | :--- |
| alphabetFrequency <br> letterFrequency | Tabulate the letters (all the letters in the alphabet for <br> alphabetFrequency, only the specified letters for <br> 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 <br> letter frequencies for a fixed-width view that is conceptually slid along <br> the input sequence. |
| consensusMatrix | Computes the consensus matrix of a set of sequences. |
| dinucleotideFrequency <br> trinucleotideFrequency <br> oligonucleotideFrequency | Fast 2-mer, 3-mer, and k-mer counting for DNA or RNA. |
| nucleotideFrequencyAt | Tallies the short sequences formed by extracting the nucleotides found <br> at a set of fixed positions from each sequence of a set of DNA or RNA <br> sequences. |

Table 2: Counting / tabulating.

| Function | Description |
| :--- | :--- |
| reverse <br> complement <br> reverseComplement | Compute the reverse, complement, or reverse-complement, of a set of <br> DNA sequences. |
| translate | Translate a set of DNA sequences into a set of Amino Acid sequences. |
| chartr <br> replaceAmbiguities | Replace letters in a sequence or set of sequences. |
| subseq, subseq<- <br> extractAt, replaceAt | Extract/replace arbitrary substrings from/in a string or set of 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 <br> pattern. unstrsplit is the reverse operation i.e. a fast implementation <br> of sapply (x, paste0, collapse=sep) for collapsing the list <br> elements of a DNAStringSetList or AAStringSetList object. |

Table 3: Sequence transformation and editing.

| Function | Description |
| :--- | :--- |
| matchPattern <br> countPattern | Find/count all the occurrences of a given pattern (typically short) in a <br> reference sequence (typically long). Support mismatches and indels. |
| vmatchPattern | Find/count all the occurrences of a given pattern (typically short) in a set <br> of reference sequences. Support mismatches and indels. <br> vcountPattern |
| matchPDict <br> countPDict <br> whichPDict | Find/count all the occurrences of a set of patterns in a reference sequence. <br> (whichPDict only identifies which patterns in the set have at least one <br> match.) Support a small number of mismatches. |
| vmatchPDict | [Note: vmat chPDict not implemented yet.] Find/count all the occur- <br> vcountPDict <br> rwhichPDict <br> only identifies for each reference sequence which patterns in the set have <br> at least one match.) Support a small number of mismatches. |
| pairwiseAlignment | Solve (Needleman-Wunsch) global alignment, (Smith-Waterman) local <br> alignment, and (ends-free) overlap alignment problems. |
| matchPWM <br> countPWM | Find/count all the occurrences of a Position Weight Matrix in a reference <br> sequence. |
| trimLRPatterns | Trim left and/or right flanking patterns from sequences. |
| matchLRPatterns | Find all paired matches in a reference sequence i.e. matches specified by <br> 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 se- <br> quence. |
| findPalindromes | Find palindromic regions in a sequence. |

Table 4: String matching / alignments.

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

Table 5: I/O functions.

