seqmagick Documentation

Release 0.3.1

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March 25, 2014

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Motivation

We often have to convert between sequence formats and do little tasks on them, and it's not worth writing scripts for that. Seqmagick is a kickass little utility built in the spirit of imagemagick to expose the file format conversion in Biopython in a convenient way. Instead of having a big mess of scripts, there is one that takes arguments:

seqmagick convert a.fasta b.phy # convert from fasta to phylip seqmagick mogrify --ungap a.fasta # remove all gaps from a.fasta, in place seqmagick info *.fasta # describe all FASTA files in the current directory

And more.

Installation

First, you'll need to install BioPython. NumPy (which parts of BioPython depend on) is not required for seqmagick to function. Once done, install with:

pip install seqmagick

Get the bleeding edge version here, or clone our repository:

git clone git://github.com/fhcrc/seqmagick.git

Use

Seqmagick can be used to query information about sequence files, convert between types, and modify sequence files. All functions are accessed through subcommands:

seqmagick <subcommand> [options] arguments

Subcommands

4.1 convert and mogrify

Convert and mogrify achieve similar goals. convert performs some operation on a file (from changing format to something more complicated) and writes to a new file. mogrify modifies a file in place, and would not normally be used to convert formats.

The two have similar signatures:

seqmagick convert [options] infile outfile

vs:

seqmagick mogrify [options] infile

Options are shared between convert and mogrify.

4.1.1 Examples

Basic Conversion

convert can be used to convert between any file types BioPython supports (which is many). For a full list of supported types, see the BioPython SeqIO wiki page.

By default, file type is inferred from file extension, so:

seqmagick convert a.fasta a.sto

converts an existing file a.fasta from FASTA to Stockholm format. Neat! But there's more.

Sequence Modification

A wealth of options await you when you're ready to do something slightly more complicated with your sequences.

Let's say I just want a few of my sequences:

<pre>\$ seqmagick converthea</pre>	d 5 example	s/test.fa	sta examp	les/test.	head.fasta
<pre>\$ seqmagick info examples/test*.fasta</pre>					
name	alignment	min_len	max_len	avg_len	num_seqs
examples/test.fasta	FALSE	972	9719	1573.67	15
examples/test.head.fasta	FALSE	978	990	984.00	5

Or I want to remove any gaps, reverse complement, select the last 5 sequences, and remove any duplicates from an alignment in place:

seqmagick mogrify --tail 5 --reverse-complement --ungap --deduplicate-sequences examples/test.fasta @

You can even define your own functions in python and use them via --apply-function.

Note: To maximize flexibility, most transformations passed as options to mogrify and convert are processed *in* order, so:

seqmagick convert --min-length 50 --cut 1:5 a.fasta b.fasta

will work fine, but:

seqmagick convert --cut 1:5 --min-length 50 a.fasta b.fasta

will never return records, since the cutting transformation happens before the minimum length predicate is applied.

4.1.2 Command-line Arguments

The full set of options to mogrify and convert are:

Sequence File Modification

line-wrap N	Adjust line wrap for sequence strings. When N is O,
	all line breaks are removed. Only fasta files are
	supported for the output format.
sort {length-asc,le	ength-desc,name-asc,name-desc}
	Perform sorting by length or name, ascending or
	descending. ASCII sorting is performed for names

Sequence Modification

```
--apply-function /path/to/module.py:function_name
                      Specify a custom function to apply to the input
                      sequences, specified as
                      /path/to/file.py:function_name. Function should accept
                      an iterable of Bio.SeqRecord objects, and yield
                      SeqRecords. Specify more than one to chain.
--cut start:end
                     1-indexed start and end positions for cutting
                      sequences, : separated. Includes last item.
--dash-gap
                     Change . and : into - for all sequences
                     Translate the sequences to lower case
--lower
--reverse
                     Reverse the order of sites in sequences
--reverse-complement Convert sequences into reverse complements
                     Remove any gaps that are present in the same position
--squeeze
                      across all sequences in an alignment (equivalent to
                      --squeeze-threshold=1.0)
--squeeze-threshold PROP
                      Trim columns from an alignment which have gaps in
                      least the specified proportion of sequences.
--transcribe {dna2rna, rna2dna}
                      Transcription and back transcription for generic DNA
                      and RNA. Source sequences must be the correct alphabet
```

	or this action will likely produce incorrect results.
translate {d	<pre>lna2protein,rna2protein,dna2proteinstop,rna2proteinstop}</pre>
	Translate from generic DNA/RNA to proteins. Options
	with "stop" suffix will NOT translate through stop
	codons .Source sequences must be the correct alphabet
	or this action will likely produce incorrect results.
ungap	Remove gaps in the sequence alignment
upper	Translate the sequences to upper case

Record Selection

deduplicate-sequence	es
	Remove any duplicate sequences by sequence content,
	keep the first instance seen
deduplicated-sequen	ces-file FILE
	Write all of the deduplicated sequences to a file
deduplicate-taxa	Remove any duplicate sequences by ID, keep the first
-	instance seen
exclude-from-file F	ILE
	Filter sequences, removing those sequence IDs in the
	specified file
include-from-file F	ILE
	Filter sequences, keeping only those sequence IDs in
	the specified file
head N	Trim down to top N sequences
max-length N	Discard any sequences beyond the specified maximum
-	length. This operation occurs *before* all length-
	changing options such as cut and squeeze.
min-length N	Discard any sequences less than the specified minimum
	length. This operation occurs *before* all length-
	changing options such as cut and squeeze.
min-ungapped-length	N
	Discard any sequences less than the specified minimum
	<pre>length, excluding gaps. This operation occurs *before*</pre>
	all length-changing options such as cut and squeeze.
pattern-include reg	ex
	Filter the sequences by regular expression in name
pattern-exclude reg	ex
	Filter out sequences by regular expression in name
prune-empty	Prune sequences containing only gaps ('-')
seq-pattern-include	regex
	Filter the sequences by regular expression in sequence
seq-pattern-exclude	regex
	Filter out sequences by regular expression in sequence
tail N	Trim down to bottom N sequences

Sequence ID Modification

	"repla	ace_patt	lern"	in sequend	ce ID		
strip-range	Strip	ranges	from	sequences	IDs,	matching	

Format Options

By default, file format is inferred from extension:

```
--input-format Format
Input file format (default: determine from extension)
--output-format Format
Output file format (default: determine from extension)
```

4.2 extract-ids

seqmagick extract-ids is extremely simple - all the IDs from a sequence file are printed to stdout (by default) or the file of your choosing:

4.3 info

seqmagick info describes one or more sequence files

4.3.1 Example

seqmagick info examples/*.fasta				
alignment TRUE	min_len 9797	max_len 9797	avg_len 9797.00	num_seqs 15
TRUE	240	240	240.00	148
TRUE	119	119	119.00	2
FALSE	972	9719	1573.67	15
FALSE	120	237	178.50	2
	alignment TRUE TRUE TRUE FALSE	alignment min_len TRUE 9797 TRUE 240 TRUE 119 FALSE 972	alignmentmin_lenmax_lenTRUE97979797TRUE240240TRUE119119FALSE9729719	alignmentmin_lenmax_lenavg_lenTRUE979797979797.00TRUE240240240.00TRUE119119.00FALSE97297191573.67

Output can be in comma-separated, tab-separated, or aligned formats. See seqmagick info -h for details.

4.4 primer-trim

primer-trim trims an alignment to a region defined by a set of forward and reverse primers. Usage is as follows:

positional arguments:			
source_file	Source alignment file		
output_file	Destination trimmed file		
forward_primer	The forward primer used		
reverse_primer	The reverse primer used. By default the reverse primer		
	is assumed to be a subsequence of the top strand (that		
	is, the reverse complement of an actual downstream PCR		
	primer). Usereverse-is-revcomp if this is not the		
	case.		
optional arguments:			
-h,help	show this help message and exit		
. 1			
1010100 10 10100mp	the top strand (default: False)		
source-format SOURC	• · · · · · · · · · · · · · · · · · · ·		
	Alignment format (default: detect from extension		
output-format OUTPU	-		
	Alignment format (default: detect from extension		
include-primers	Include the primers in the output (default: False)		
max-hamming-distanc	e MAX_HAMMING_DISTANCE		
-	Maximum Hamming distance between primer and alignment		
	site (default: 1). IUPAC ambiguous bases in the primer		
	matching unambiguous bases in the alignment are not		
	penalized		
prune-action {trim,	isolate}		
	Action to take. Options are trim (trim to the region		
	defined by the two primers, decreasing the width of		
	the alignment), or isolate (convert all characters		
	outside the primer-defined area to gaps). default:		
	trim		

4.5 quality-filter

quality-filter truncates and removes sequences that don't match a set of quality criteria. The subcommand takes a FASTA and quality score file, and writes the results to an output file:

```
positional arguments:
 input_fasta
                        Input fasta file
 input_qual
                        The quality scores associated with fasta_file
 output_file
                        Output file. Format determined from extension.
optional arguments:
 -h, --help
                        show this help message and exit
 --min-mean-quality QUALITY
                        Minimum mean quality score for each read [default: 25]
 --min-length LENGTH
                        Minimum length to keep sequence [default: None]
 --quality-window WINDOW_SIZE
                        Window size for truncating sequences. When set to a
                        non-zero value, sequences are truncated where the mean
                        mean quality within the window drops below --min-mean-
                        quality. [default: 0]
 --ambiguous-action {truncate,drop}
                        Action to take on ambiguous base in sequence (N's).
                        [default: no action]
```

Supported File Extensions

Extension	Format
.afa	fasta
.aln	clustal
.fa	fasta
.faa	fasta
.fas	fasta
.fasta	fasta
.fastq	fastq
.ffn	fasta
.fna	fasta
.frn	fasta
.gb	genbank
.gbk	genbank
.needle	emboss
.phy	phylip
.phylip	phylip
.phyx	phylip-relaxed (note: requires building BioPython from the master branch until v1.58 is released)
.qual	qual
.sff	sff
.sth	stockholm
.sto	stockholm

By default, seqmagick infers the file type from extension. Currently mapped extensions are:

If an extension is not listed, you can either rename the file to a supported extension, or specify it manually via --input-format or --output-format.

Acknowledgements

seqmagick is written and maintained by the Matsen Group at the Fred Hutchinson Cancer Research Center.

Contributing

We welcome contributions! Simply fork the repository on GitHub and send a pull request.