

Translate fasta headers

Translate long fasta headers to short - and back!

Your alignment program X doesn't allow strings longer than n characters, but all your info is in the fasta headers of your file. What to do?

Use `translate_fasta_headers.pl` on your fasta file to create short labels and a translation table. Run your program X, and then back-translate your fasta headers by running `translate_fasta_headers.pl` again!

And if you created a tree with the short labels, try to back-translate using `replace_taxon_labels_in_newick.pl`!

If you only wish to transform your long fasta headers to short, without keeping the information about how they were translated, the quick solution might be to use `awk`:

```
awk '/>/{ $0=">Seq_" ++n } 1' long.fas
```

But, if you want to be able to back-translate, read on!

DESCRIPTION

Replace fasta headers with headers taken from tab delimited file. If no tab file is given, the (potentially long) fasta headers are replaced by short labels "Seq_1", "Seq_2", etc, and the short and original headers are printed to a translation file.

If you wish, you may choose your own prefix (instead of `Seq_`). This could be handy if, for example, you wish to concatenate files.

The script for translating labels in Newick trees is somewhat limited in capacity due to the restrictions of the Newick tree format. Use with caution.

USAGE

```
./translate_fasta_headers.pl [options] <file>
```

From long to short labels:

```
./translate_fasta_headers.pl --out=short.fas long.fas
```

And back, using a translation table:

```
./translate_fasta_headers.pl --tabfile=short.fas.translation.tab short.fas
```

Slightly shorter version (see note about the `--out` option below):

```
./translate_fasta_headers.pl long.fas > short.fas  
./translate_fasta_headers.pl -t long.fas.translation.tab short.fas
```

Use your own prefix:

```
./translate_fasta_headers.pl --prefix='Own_' long.fas
```

Translate short seq labels in Newick tree to long:

```
./replace_taxon_labels_in_newick.pl -t long.fas.translation.tab short.fas.phy
```

Use your own prefix:

```
./translate_fasta_headers.pl --prefix='Own_' long.fas
```

Translate short seq labels in Newick tree to long:

```
./replace_taxon_labels_in_newick.pl -t long.fas.translation.tab short.fas.phy
```

OPTIONS

`translate_fasta_headers.pl`

- `-t, --tabfile=<filename>` Specify tab-separated translation file with unique “short” labels to the left, and “long” names to the right. Translation will be from left to right.
- `-o, --out=<filename>` Specify output file for the fasta sequences. **Note:** If `--out=<filename>` is specified, the translation file will be named `<filename>.translation.tab`. This simplifies back translation. If, on the other hand, `--out` is not used, the translation file will be named after the infile!
- `-i, --in=<filename>` Specify name of fasta file. Can be skipped as script reads files from STDIN.
- `-n, --notab` Do not create a translation file.
- `-p, --prefix=<string>` User your own prefix (default is `Seq_`). A numerical will be added to the labels (e.g. `Own_1`, `Own_2`, ...)
- `-f, --forceorder` [NOT YET IMPLEMENTED!] translate in order of appearance in the fasta file, and use the same order as in the tabfile - without rigid checking of the names! This allows non-unique labels in the left column.
- `-h, --help` Show this help text and quit.

`replace_taxon_labels_in_newick.pl`

- `-t, --table=<translation.tab>` file with table describing what will be translated with what.
- `-h, --help` Help text.
- `-o, --out=<out.file>` Print to outfile `out.file`, else to STDOUT.

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FILES

- `translate_fasta_headers.pl` Perl script
- `replace_taxon_labels_in_newick.pl` Perl script
- `long.fas` Example file with long fasta headers
- `short.fas.translation.tab` Example translation table
- `short.fas` Example output with short fasta headers
- `short.fas.phy` Example Newick tree with short labels
- `README.md` Documentation, markdown format
- `README.pdf` Documentation, PDF format

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