

A toolkit for DNA sequence analysis and manipulation

D. Pratas (pratas@ua.pt)
J. R. Almeida (joao.rafael.almeida@ua.pt)
A. J. Pinho (ap@ua.pt)

IEETA/DETI, University of Aveiro, Portugal

 $Version\ 1.7.17$

Contents

1	Introduction		
	1.1	Installation	2
	1.2	License	2
2	FASTQ tools		4
	2.1	Program goose-fastq2fasta	5
	2.2	Program goose-fastq2mfasta	6
	2.3	Program goose-FastqExcludeN	8
	2.4	Program goose-FastqExtractQualityScores	9
	2.5	Program goose-FastqInfo	10
	2.6	Program goose-FastqMaximumReadSize	11
3	FASTA tools		
	3.1	Program goose-fasta2seq	13
	3.2	Program goose-fastaextract	14
	3.3	Program goose-fastaextractbyread	15
	3.4	Program goose-fastainfo	17
	3.5	Program goose-mutatefasta	18
	3.6	Program goose-randfastaextrachars	19
4	Ger	nomic sequence tools	2 2
5	Amino acid sequence tools		
	5.1	Program goose-AminoAcidToGroup	23
	5.2	Program goose-ProteinToPseudoDNA	24
6	Ger	neral purpose tools	27
	6.1	Program goose-reverse	27
Bi	Bibliography		

Introduction

Recent advances in DNA sequencing have revolutionized the field of genomics, making it possible for research groups to generate large amounts of sequenced data, very rapidly and at substantially lower cost. Its storage have been made using specific file formats, such as FASTQ and FASTA. Therefore, its analysis and manipulation is crucial [?]. Several frameworks for analysis and manipulation emerged, namely GALAXY [?], GATK [?], HTSeq [?], MEGA [?], among others. In the majority, these frameworks require licenses and do not provide a low level access to the information, since they are commonly approached by scripting or interfaces.

We describe GOOSE, a (free) novel toolkit for analyzing and manipulating FASTA-FASTQ formats and sequences (DNA, amino acids, text), with many complementary tools. The toolkit is for Linux-based systems, built for fast processing. GOOSE supports pipes for easy integration. It includes tools for information display, randomizing, edition, conversion, extraction, searching, calculation and visualization. GOOSE is prepared to deal with very large datasets, typically in the scale Gigabytes or Terabytes.

The toolkit is a command line version, using the prefix "goose-" followed by the suffix with the respective name of the program. GOOSE is implemented in C language and it is available, under GPLv3, at:

```
https://pratas.github.io/goose
```

1.1 Installation

For GOOSE installation, run:

```
git clone https://github.com/pratas/goose.git
cd goose/src/
make
```

1.2 License

The license is **GPLv3**. In resume, everyone is permitted to copy and distribute verbatim copies of this license document, but changing it is not allowed. For details on the license, consult: http://www.gnu.org/

licenses/gpl-3.0.html.

FASTQ tools

Current available tools for FASTQ format analysis and manipulation include:

- 1. goose-fastq2fasta: it converts a FASTQ file format to a pseudo FASTA file.
- 2. goose-fastq2mfasta: it converts a FASTQ file format to a pseudo Multi-FASTA file.
- 3. goose-fastqclustreads
- 4. goose-FastqExcludeN: it discards the FASTQ reads with the minimum number of "N" symbols.
- 5. goose-FastqExtractQualityScores: it extracts all the quality-scores from FASTQ reads.
- 6. goose-FastqInfo: it analyses the basic informations of FASTQ file format.
- 7. goose-FastqMaximumReadSize: it filters the FASTQ reads with the length higher than the value defined.
- $8. \ {\tt goose-FastqMinimumLocalQualityScoreForward}\\$
- $9.\ {\tt goose-FastqMinimumLocalQualityScoreReverse}$
- $10.\ {\tt goose-FastqMinimumQualityScore}$
- 11. goose-FastqMinimumReadSize
- 12. goose-count
- 13. goose-extractreadbypattern
- 14. goose-fastqpack
- 15. goose-fastqsimulation
- 16. goose-FastqSplit
- 17. goose-FastqTrimm

- 18. goose-fastqunpack
- 19. goose-filter
- $20. \; {\tt goose-findnpos}$
- 21. goose-genrandomdna
- 22. goose-getunique
- $23. \; {\tt goose-info}$
- $24.\ {\tt goose-mfmotifcoords}$
- $25.\ {\tt goose-mutatefastq}$
- 26. goose-newlineonnewx
- 27. goose-period
- 28. goose-permuteseqbyblocks
- 29. goose-randfastqextrachars
- 30. goose-real2binthreshold
- 31. goose-reducematrixbythreshold
- 32. goose-renamehumanheaders
- 33. goose-searchphash
- 34. goose-seq2fasta
- $35.\ {\tt goose-seq2fastq}$
- $36.\ {\tt goose-SequenceToGroupSequence}$
- 37. goose-splitreads
- 38. goose-wsearch

2.1 Program goose-fastq2fasta

The goose-fastq2fasta converts a FASTQ file format to a pseudo FASTA file. However, it does not align the sequence. Also, it extracts the sequence and adds a pseudo header. For help type:

```
./goose-fastq2fasta -h
```

In the following subsections, we explain the input and output paramters.

Input parameters

The goose-fastq2fasta program needs two streams for the computation, namely the input and output standard. The input stream is a FASTQ file.

The attribution is given according to:

An example on such an input file is:

Output

The output of the goose-fastq2fasta program a FASTA file.

An example, for the input, is:

```
GGGTGATGGCCGCTGCCGATGGCGTCAAATCCCACCAAGTTACCCTTAACAACTTAAGGGTTTTCAAATAGA
GTTCAGGGATACGACGTTTGTATTTTAAGAATCTGAAGCAGAAGTCGATGATAATACGCGTCGTTTTATCAT
```

2.2 Program goose-fastq2mfasta

The goose-fastq2mfasta onverts a FASTQ file format to a pseudo Multi-FASTA file. However, it does not align the sequence. Also, it extracts the sequence and adds a pseudo header.

For help type:

```
./goose-fastq2mfasta-h
```

In the following subsections, we explain the input and output paramters.

Input parameters

The goose-fastq2mfasta program needs two streams for the computation, namely the input and output standard. The input stream is a FASTQ file.

The attribution is given according to:

```
Usage: ./goose-fastq2mfasta [options] [[--] args]
    or: ./goose-fastq2mfasta [options]

It converts a FASTQ file format to a pseudo Multi-FASTA file.

It does NOT align the sequence.

It extracts the sequence and adds each header in a Multi-FASTA format.

-h, --help show this help message and exit

Basic options
    < input.fastq Input FASTQ file format (stdin)
    > output.mfasta Output Multi-FASTA file format (stdout)

Example: ./goose-fastq2mfasta < input.fastq > output.mfasta
```

An example on such an input file is:

Output

The output of the goose-fastq2mfasta program a Multi-FASTA file.

An example, for the input, is:

```
>SRROO1666.1 071112_SLXA-EAS1_s_7:5:1:817:345 length=72

GGGTGATGGCCGCTGCCGATGGCGTCAAATCCCACCAAGTTACCCTTAACAACTTAAGGGTTTTCAAATAGA

>SRROO1666.2 071112_SLXA-EAS1_s_7:5:1:801:338 length=72

GTTCAGGGATACGACGTTTGTATTTTAAGAATCTGAAGCAGAAGTCGATGATAATACGCGTCGTTTTATCAT
```

2.3 Program goose-FastqExcludeN

The goose-FastqExcludeN discards the FASTQ reads with the minimum number of "N" symbols. Also, if present, it will erase the second header (after +).

For help type:

```
./goose-FastqExcludeN-h
```

In the following subsections, we explain the input and output paramters.

Input parameters

The goose-FastqExcludeN program needs two streams for the computation, namely the input and output standard. The input stream is a FASTQ file.

The attribution is given according to:

```
Usage: ./goose-FastqExcludeN [options] [[--] args]
   or: ./goose-FastqExcludeN [options]
It discards the FASTQ reads with the minimum number of ''N'' symbols. If present,
it will erase the second header (after +).
    -h, --help
                          show this help message and exit
Basic options
   -m, --max=<int>
                          The maximum of of "N" symbols in the read
   < input.fastq</pre>
                          Input FASTQ file format (stdin)
   > output
                          Output read information (stdout)
Example: ./goose-FastqExcludeN < input.fastq > output
Output example :
<FASTQ filtered reads>
Total reads : value
Filtered reads : value
```

An example on such an input file is:

Output

The output of the goose-FastqExcludeN program is a set of all the filtered FASTQ reads, followed by the execution report.

Using the max value as 5, an example for this input, is:

2.4 Program goose-FastqExtractQualityScores

The goose-FastqExtractQualityScores extracts all the quality-scores from FASTQ reads. For help type:

```
./goose-FastqExtractQualityScores -h
```

In the following subsections, we explain the input and output paramters.

Input parameters

The goose-FastqExtractQualityScores program needs two streams for the computation, namely the input and output standard. The input stream is a FASTQ file.

The attribution is given according to:

An example on such an input file is:

Output

The output of the goose-FastqExtractQualityScores program is a set of all the quality scores from the FASTQ reads, followed by the execution report.

An example, for the input, is:

2.5 Program goose-FastqInfo

The goose-FastqInfo analyses the basic informations of FASTQ file format. For help type:

```
./goose-FastqInfo-h
```

In the following subsections, we explain the input and output paramters.

Input parameters

The goose-FastqInfo program needs two streams for the computation, namely the input and output standard. The input stream is a FASTQ file.

The attribution is given according to:

```
Usage: ./goose-FastqInfo [options] [[--] args]
   or: ./goose-FastqInfo [options]
It analyses the basic informations of FASTQ file format.
   -h, --help
                        show this help message and exit
Basic options
   < input.fastq</pre>
                         Input FASTQ file format (stdin)
   > output
                         Output read information (stdout)
Example: ./goose-FastqInfo < input.fastq > output
Output example :
Total reads : value
Max read length : value
Min read length : value
Min QS value : value
Max QS value : value
              : value
QS range
```

An example on such an input file is:

Output

The output of the goose-FastqInfo program is a set of informations related with the file readed. An example, for the input, is:

```
Total reads : 2

Max read length : 72

Min read length : 72

Min QS value : 41

Max QS value : 73

QS range : 33
```

2.6 Program goose-FastqMaximumReadSize

The goose-FastqMaximumReadSize filters the FASTQ reads with the length higher than the value defined. For help type:

```
./goose-FastqMaximumReadSize -h
```

In the following subsections, we explain the input and output paramters.

Input parameters

The goose-FastqMaximumReadSize program needs two streams for the computation, namely the input and output standard. The input stream is a FASTQ file.

The attribution is given according to:

```
Usage: ./goose-FastqMaximumReadSize [options] [[--] args]
or: ./goose-FastqMaximumReadSize [options]

It filters the FASTQ reads with the length higher than the value defined.

If present, it will erase the second header (after +).

-h, --help show this help message and exit

Basic options
-s, --size=<int> The maximum read length
< input.fastq Input FASTQ file format (stdin)
```

```
> output Output read information (stdout)

Example: ./goose-FastqMaximumReadSize < input.fastq > output

Output example:

<FASTQ filtered reads >
Total reads : value
Filtered reads : value
```

An example on such an input file is:

Output

The output of the goose-FastqMaximumReadSize program is a set of all the filtered FASTQ reads, followed by the execution report.

Using the size value as 60, an example for this input, is:

FASTA tools

Current available FASTA tools, for analysis and manipulation, are:

- 1. goose-fasta2seq: it converts a FASTA or Multi-FASTA file format to a seq.
- 2. goose-fastaextract: it extracts sequences from a FASTA file, which the range is defined by the user in the parameters.
- 3. goose-fastaextractbyread: it extracts sequences from each read in a Multi-FASTA file (splited by \n), which the range is defined by the user in the parameters.
- 4. goose-fastainfo: it shows the readed information of a FASTA or Multi-FASTA file format.
- 5. goose-mutatefasta: it reates a synthetic mutation of a fasta file given specific rates of editions, deletions and additions.
- 6. goose-randfastaextrachars: it substitues in the DNA sequence the outside ACGT chars by random ACGT symbols.

3.1 Program goose-fasta2seq

The goose-fasta2seq converts a FASTA or Multi-FASTA file format to a seq. For help type:

```
./goose-fasta2seq -h
```

In the following subsections, we explain the input and output paramters.

Input parameters

The goose-fasta2seq program needs two streams for the computation, namely the input and output standard. The input stream is a FASTA or Multi-FASTA file.

The attribution is given according to:

An example on such an input file is:

Output

The output of the goose-fasta2seq program is a group sequence.

An example, for the input, is:

3.2 Program goose-fastaextract

The goose-fastaextract extracts sequences from a FASTA file, which the range is defined by the user in the parameters.

For help type:

```
./goose-fastaextract -h
```

In the following subsections, we explain the input and output paramters.

Input parameters

The goose-fastaextract program needs two parameters, which defines the begin and the end of the extraction, and two streams for the computation, namely the input and output standard. The input stream is a FASTA file.

The attribution is given according to:

```
Usage: ./goose-fastaextract [options] [[--] args]
or: ./goose-fastaextract [options]

It extracts sequences from a FASTA file.

-h, --help show this help message and exit

Basic options
-i, --init=<int> The first position to start the extraction (default 0)
-e, --end=<int> The last extract position (default 100)
< input.fasta Input FASTA or Multi-FASTA file format (stdin)
> output.seq Output sequence file (stdout)

Example: ./goose-fastaextract -i <init> -e <end> < input.fasta > output.seq
```

An example on such an input file is:

Output

The output of the goose-fastaextract program is a group sequence.

An example, using the value 0 as extraction starting point and the 50 as the end, for the provided input, is:

```
ACAAGACGGCCTCCTGCTGCTGCTCTCCGGGGCCACGGCCCTGGAGG
```

3.3 Program goose-fastaextractbyread

The goose-fastaextractbyread extracts sequences from a FASTA or Multi-FASTA file, which the range is defined by the user in the parameters.

For help type:

```
./goose-fastaextractbyread -h
```

In the following subsections, we explain the input and output paramters.

Input parameters

The goose-fastaextractbyread program needs two parameters, which defines the begin and the end of the extraction, and two streams for the computation, namely the input and output standard. The input stream is a FASTA or Multi-FASTA file.

The attribution is given according to:

```
Usage: ./goose-fastaextractbyread [options] [[--] args]
or: ./goose-fastaextractbyread [options]

It extracts sequences from each read in a Multi-FASTA file (splited by \n)

-h, --help show this help message and exit

Basic options
-i, --init=<int> The first position to start the extraction (default 0)
-e, --end=<int> The last extract position (default 100)
< input.fasta Input FASTA or Multi-FASTA file format (stdin)
> output.fasta Output FASTA or Multi-FASTA file format (stdout)

Example: ./goose-fastaextractbyread -i <init> -e <end> < input.fasta > output.fasta
```

An example on such an input file is:

```
> ABOOO 264 | acc = ABOOO 264 | descr = Homo sapiens mRNA

ACAAGACGGCCTCCTGCTGCTGCTGCTCCCCGGGGCCCACGGCCCTGGAGGGTCCACCGCTGCCCTGCTGCCATTGTCCCC

GGCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCAGGAAGCGGCAGGAA

GTGGTTTGAGTGGACCTCCGGGCCCCTCATAGGAGAGGAAGCTCGGGAGGTGGCCAGGCGGCAGGAAGCAGGCCAGTGCC

GCGAATCCGCGCGCCGGGACAGAATCTCCTGCAAAGCCCTGCAGGAACCTTCTTCTGGAAGACCTTCTCCCACCCCCCCAGC

TAAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAA

> ABOOO 263 | acc = ABOOO 263 | descr = Homo sapiens mRNA

ACAAGATGCCATTGTCCCCCGGGCCTCCTGCTGCTGCTCCTCCGGGGCCACCGCTGCCCTTGCCCCTTGGAGGGT

GGCCCCACCGGCCGAGACAGCATATGCAGGAAGCGGCAGGAATAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTG

GTGGTTTGAGTGGACCTCCCAGGCCAGTGCCGGGCCCCTCATAGGAGAGGAAGCTCGGGAGGTGGCCAGGCGCAGGAAG

GCGCACCCCCCCCAGCAATCCGCGCGCCCGGGACAGAATGCCCTTGCAGAACCTTCTTCTCTCCTCCTCCTCCTGCAAA

TAAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAA
```

Output

The output of the goose-fastaextractbyread program is FASTA or Multi-FASTA file with the extracted sequences.

An example, using the value 0 as extraction starting point and the 50 as the end, for the provided input, is:

```
>AB000264 |acc=AB000264|descr=Homo sapiens mRNA
ACAAGACGGCCTCCTGCTGCTGCTCCTCCGGGGCCACGGCCCTGGAGG
>AB000263 |acc=AB000263|descr=Homo sapiens mRNA
ACAAGATGCCATTGTCCCCCGGGCCTCCTGCTGCTGCTCCCGGGGCC
```

3.4 Program goose-fastainfo

The goose-fastainfo shows the readed information of a FASTA or Multi-FASTA file format. For help type:

```
./goose-fastainfo -h
```

In the following subsections, we explain the input and output paramters.

Input parameters

The goose-fastainfo program needs two streams for the computation, namely the input and output standard. The input stream is a FASTA or Multi-FASTA file.

The attribution is given according to:

```
Usage: ./goose-fastainfo [options] [[--] args]
   or: ./goose-fastainfo [options]
It shows read information of a FASTA or Multi-FASTA file format.
    -h, --help
                         show this help message and exit
Basic options
   < input.fasta</pre>
                        Input FASTA or Multi-FASTA file format (stdin)
    > output
                          Output read information (stdout)
Example: ./goose-fastainfo < input.fasta > output
Output example :
Number of reads
                     : value
Number of bases
                     : value
MIN of bases in read : value
MAX of bases in read : value
AVG of bases in read : value
```

An example on such an input file is:

```
> ABOOO 264 | acc = ABOOO 264 | descr = Homo sapiens mRNA
ACAAGACGGCCTCCTGCTGCTGCTGCTCCTCCGGGGCCACGGCCCTGGAGGGTCCACCGCTGCCCTGCTGCCATTGTCCCC
GGCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCAGGAAGCGGCAGGAA
GTGGTTTGAGTGGACCTCCGGGGCCCCTCATAGGAGAGCTCGGGAGGTGGCCAGGCGGCAGGAAGCAGGCCAGTGCC
GCGAATCCGCGCGCGCGGGACAGAATCTCCTGCAAAGCCCTGCAGGAACTTCTTCTGGAAGACCTTCTCCCACCCCCCCAGC
TAAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAA
> ABOOO 263 | acc = ABOOO 263 | descr = Homo sapiens mRNA
```

Output

The output of the goose-fastainfo program is a set of informations related with the file readed.

An example, for the input, is:

```
Number of reads : 2
Number of bases : 736
MIN of bases in read : 368
MAX of bases in read : 368
AVG of bases in read : 368.0000
```

3.5 Program goose-mutatefasta

The goose-mutatefasta creates a synthetic mutation of a fasta file given specific rates of editions, deletions and additions. All these parameters are defined by the user, and their are optional.

For help type:

```
./goose-mutatefasta-h
```

In the following subsections, we explain the input and output paramters.

Input parameters

The goose-mutatefasta program needs two streams for the computation, namely the input and output standard. However, optional settings can be supplied too, such as the starting point to the random generator, and the edition, deletion and insertion rates. Also, the user can choose to use the ACGTN alphabet in the synthetic mutation. The input stream is a FASTA or Multi-FASTA File.

The attribution is given according to:

```
Usage: ./goose-mutatefasta [options] [[--] args]
or: ./goose-mutatefasta [options]

Creates a synthetic mutation of a fasta file given specific rates of editions, deletions and additions

-h, --help show this help message and exit

Basic options
<input.fasta Input FASTA or Multi-FASTA file format (stdin)
> output.fasta Output FASTA or Multi-FASTA file format (stdout)

Optional
-s, --seed=<int> Starting point to the random generator
```

```
-e, --edit-rate=<dbl>
Defines the edition rate (default 0.0)
-d, --deletion-rate=<dbl>
Defines the deletion rate (default 0.0)
-i, --insertion-rate=<dbl>
Defines the insertion rate (default 0.0)
-a, --ACGTN-alphabet

When active, the application uses the ACGTN alphabet

Example: ./goose-mutatefasta -s <seed> -e <edit rate> -d <deletion rate> -i <insertion rate> -a < input.fast
```

An example on such an input file is:

Output

The output of the goose-mutatefasta program is a FASTA or Multi-FASTA file whith the synthetic mutation of input file.

Using the seed value as 1 and the edition rate as 0.5, an example for this input, is:

```
> ABOOO 264 | acc = ABOOO 264 | descr = Homo sapiens mrna

ACGCAACGNATTCCTGCTGATCATANTGTNCCGCNCCCCNGCGACGGGGNCTCNCNNGCACACATNGTACCATTGTCCAC

NCTTNCANGTNANCGCTAGCAGGCTACNGTTTNTCCTCNCCTANNCCAANCNGGCGTNNNTACACTGGCACGTGCAGGCA

TNGGTCGGCNGGNNCCTCCGGNAACGGCACCGGAGACGAAGCTCGGNGGNTATACAGGTGTCANGAAACATCCCCGGGNC

GNGTGNCCNNGAANCCANAGAGTATCTCACTCACAACCCTGCGTGCACNTCTAGAGNANGACCTTACNCACCNTCCCNTT

NNGTACCACACCAATGAACGCTGCAGAAAGTCTGTTTNNAGGNGNGCA

> ABOOO 263 | acc = ABOOO 263 | descr = Homo sapiens mrna

ATTTGAAGGCAANCGGNCCAGNAATNCGGNGGGTGCNGCTCNTGTNGGCTACGGNCATCGCGGCCCTGCTNTANTAAGCN

TGAACCACCGNTCGNNGCACTTAGCAATNGCGNAANCCGTCGGCACGGCGAGACNAANCCGCTANTNNTTTCCCGCTNA

ATGGNTGTACAAGACCNACTANACCANCCTCCGTCACCACACTGGAGCGCANGATGGNNCGCTGNCTAGNAGNCNNTGAG

GCGCTCCNTCCTANAAANCCGTGGNCGAGCNCCCTATGGNAGNGTGGGGGTTTTACCGGAAGACCNTCGNGCCCTATGGG

AGCAATCANAANCTAGAAAGCTTACNGATGGTGANGAANTAGACTANG
```

3.6 Program goose-randfastaextrachars

The goose-randfastaextrachars substitues in the DNA sequence the outside ACGT chars by random ACGT symbols. It works both in FASTA and Multi-FASTA file formats.

For help type:

```
./goose-randfastaextrachars -h
```

In the following subsections, we explain the input and output paramters.

Input parameters

The goose-randfastaextrachars program needs two streams for the computation, namely the input and output standard. The input stream is a FASTA or Multi-FASTA file.

The attribution is given according to:

```
Usage: ./goose-randfastaextrachars [options] [[--] args]
or: ./goose-randfastaextrachars [options]

It substitues in the DNA sequence the outside ACGT chars by random ACGT symbols.

It works both in FASTA and Multi-FASTA file formats

-h, --help show this help message and exit

Basic options
<input.fasta Input FASTA or Multi-FASTA file format (stdin)
> output.fasta Output FASTA or Multi-FASTA file format (stdout)

Example: ./goose-randfastaextrachars < input.fasta > output.fasta
```

An example on such an input file is:

Output

The output of the goose-randfastaextrachars program is a FASTA or Multi-FASTA file. An example, for the input, is:

```
> ABOOO264 | acc=ABOOO264 | descr=Homo sapiens mRNA
ATAAGACGGCCTCCTGCTGCTGCTCCCGGGGCCACGGCCCTGGAGGGTCCCCCGCTGCCCTGCTGCCATTGTCCCC
TGCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCGGGAAGCGGCAGGAA
GAGGTTTGAGTGGACCTCCCGGCCCCTCATAGGAGAGGAAGCCGGGGAGGTGGCCAGGCGAGCAGGAAGCAGGCCAGTGTC
GCGAATCCGGGCGCCGGGACAGAATCTCCTGCAAAGCCCTGCAGGAACTTCTTCTGGAAGACCTTCTCCACCCCCCCTTG
TAAAAGATCACCCATGAATGCTCACGCAAATTTAATTACAGACCTGAA
> ABOOO263 | acc=ABOOO263 | descr=Homo sapiens mRNA
GCGAATCCGTGCGCCGGGACAGAATCTCCTTCTCCACCCCCCCATCTGCAAAGCCCTGCAGGAACTTCTTCTTGGAAGACC
GGCCCCACCTAAGGAAAAGCAGCCTCCAGGAACTTCTTCCTCGCTTGGGCCGAGCATATGCGGGAAGCGG
AGAAGACGGCCTCCTGCTGCTCCCGGGGCCACGTCCCTGCTCCAGGGTCCTCCGCTGCCCTTGCCATTGC
```

GAGGAAGCGGGGGAGGTGGCCAGGCAGGAAGCAGGCCAGTGGCGCGGTTTGAGTGGACCTCCTGGCCCCTCATAGGA TCACGCAACTTTAATTACAGACCTGAATAAAATGTCACCCATGAATGC

Genomic sequence tools

Current available genomic sequence tools, for analysis and manipulation, are:

- 1. goose-mutatedna
- $2. \ {\tt goose-randseq} {\tt extrachars}$
- 3. goose-geco
- $4.\ {\tt goose-gede}$

Amino acid sequence tools

Current available amino acid sequence tools, for analysis and manipulation, are:

- 1. goose-AminoAcidToGroup: it converts an amino acid sequence to a group sequence.
- 2. goose-ProteinToPseudoDNA: it converts an amino acid (protein) sequence to a pseudo DNA sequence.

5.1 Program goose-AminoAcidToGroup

The goose-AminoAcidToGroup converts an amino acid sequence to a group sequence. For help type:

```
./goose-AminoAcidToGroup-h
```

In the following subsections, we explain the input and output paramters.

Input parameters

The goose-AminoAcidToGroup program needs two streams for the computation, namely the input and output standard. The input stream is an amino acid sequence. The attribution is given according to:

```
Amino acids with electric charged side chains: POSITIVE
Н
K
D
    N
Ε
   N Amino acids with electric charged side chains: NEGATIVE
S
Т
   U Amino acids with electric UNCHARGED side chains
N
Q
    U
C
   S
U
G
   S Special cases
Р
    Н
   Н
V
Ι
L
М
   H Amino acids with hydrophobic side chains
F
   Н
Y
   Н
   Н
    * Others
   X Unknown
```

It can be used to group amino acids by properties, such as electric charge (positive and negative), uncharged side chains, hydrophobic side chains and special cases. An example on such an input file is:

```
IPFLLKKQFALADKLVLSKLRQLLGGRIKMMPCGGAKLEPAIGLFFHAIGINIKLGYGMTETTATVSCWHDFQFNPNSIG
TLMPKAEVKIGENNEILVRGGMVMKGYYKKPEETAQAFTEDGFLKTGDAGEFDEQGNLFITDRIKELMKTSNGKYIAPQY
IESKIGKDKFIEQIAIIADAKKYVSALIVPCFDSLEEYAKQLNIKYHDRLELLKNSDILKMFE
```

Output

The output of the goose-AminoAcidToGroup program is a group sequence.

An example, for the input, is:

HSHHHPPUHHHHNPHHHUPHPUHHSSPHPHHSSSSHPHNSHHSHHPHHSHUHPHSHSHUNUUHUHUSHPNHUHUSUUHS UHHSPHNHPHSNUUNHHHPSSHHHPSHHPPSNNUHUHHUNNSHHPUSNHSNHNNUSUHHHUNPHPNHHPUUUSPHHHSUH HNUPHSPNPHHNUHHHHHNHPPHHUHHHHSSHNUHNNHHPUHUHPHPNPHNHHPUUNHHPHHN

5.2 Program goose-ProteinToPseudoDNA

The goose-ProteinToPseudoDNA converts an amino acid (protein) sequence to a pseudo DNA sequence. For help type:

```
./goose-ProteinToPseudoDNA -h
```

In the following subsections, we explain the input and output paramters.

Input parameters

The goose-ProteinToPseudoDNA program needs two streams for the computation, namely the input and output standard. The input stream is an amino acid sequence. The attribution is given according to:

```
Usage: ./goose-ProteinToPseudoDNA [options] [[--] args]
   or: ./goose-ProteinToPseudoDNA [options]
It converts a protein sequence to a pseudo DNA sequence.
    -h, --help show this help message and exit
Basic options
   < input.prot</pre>
                    Input amino acid sequence file (stdin)
    > output.dna
                    Output DNA sequence file (stdout)
Example: ./goose-ProteinToPseudoDNA < input.prot > output.dna
Table:
Prot
       DNA
Α
   GCA
С
   TGC
D
   GAC
Ε
   GAG
   TTT
F
G
   GGC
   CAT
Н
Ι
   ATC
K
   AAA
   CTG
L
M
   ATG
N
   AAC
P
   CCG
Q
   CAG
R
   CGT
   TCT
T
    ACG
V
   GTA
W
   TGG
Y
   TAC
   TAG
    GGG
```

It can be used to generate pseudo-DNA with characteristics passed by amino acid (protein) sequences. An example on such an input file is:

```
IPFLLKKQFALADKLVLSKLRQLLGGRIKMMPCGGAKLEPAIGLFFHAIGINIKLGYGMTETTATVSCWHDFQFNPNSIG
TLMPKAEVKIGENNEILVRGGMVMKGYYKKPEETAQAFTEDGFLKTGDAGEFDEQGNLFITDRIKELMKTSNGKYIAPQY
IESKIGKDKFIEQIAIIADAKKYVSALIVPCFDSLEEYAKQLNIKYHDRLELLKNSDILKMFE
```

Output

The output of the goose-ProteinToPseudoDNA program is a DNA sequence. An example, for the input, is:

General purpose tools

goose-comparativemap
 goose-BruteForceString
 goose-char2line
 goose-sum
 goose-min
 goose-minus
 goose-max
 goose-extract

6.1 Program goose-reverse

The goose-reverse reverses the order of a sequence file.

10. goose-reverse: it reverses the order of a sequence.

For help type:

9. goose-segment

```
./goose-reverse -h
```

In the following subsections, we explain the input and output paramters.

Input parameters

The goose-reverse program needs two streams for the computation, namely the input and output standard. The input stream is a sequence file.

The attribution is given according to:

An example on such an input file is:

Output

The output of the goose-reverse program is a group sequence.

An example, for the input, is: