

The Genomics Toolkit

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Chapter 1

Introduction

Recent advances in DNA sequencing, specifically in next-generation sequencing (NGS), revolutionised the field of genomics, making possible the generation of large amounts of sequencing data very rapidly and at substantially low cost [1]. This new technology also brought with it several challenges, namely in what concerns the analysis, storage, and transmission of the generated sequences [2, 3]. As a consequence, several specialised tools were developed throughout the years in order to deal with these challenges.

Firstly, the storage of the raw data generated by NGS experiments is possible by using several file formats, the FASTQ and FASTA are the most commonly used [4]. FASTQ is an extension of the FASTA format, that besides the nucleotide sequence, also stores associated per base quality score and it is considered the standard format for sequencing data storage and exchange [5].

Regarding the analysis and manipulation of these sequencing data files many software applications emerged, including fqtools [6], FASTX-Toolkit [7], GALAXY [8], GATK [9], MEGA [10], SeqKit [11], among others. Fqtools is a suite of tools to view, manipulate and summarise FASTQ data. This software also identifies invalid FASTQ files [6]. GALAXY, in its turn, is an open, web-based scientific platform for analysing genomic data [12]. This platform integrates several specialised sets of tools, e.g. for manipulating FASTQ files [13]. FASTX-Toolkit is a collection of command-line tools to process FASTA and FASTQ files. This toolkit is available in two forms: as a command-line, or integrated into the web-based platform GALAXY [7]. SeqKit is another toolkit used to process FASTA and FASTQ files and is available for all major operating systems [11]. The Genome Analysis Toolkit (GATK) was designed as a structured programming framework to simplify the development of analysis tools. However, nowadays, it is a suite of tools focused on variant discovering and genotyping [14]. More towards the evolutionary perspectives, Molecular Evolutionary Genetics Analysis (MEGA) software provides tools to analyse DNA and protein sequences statistically [15]. Several of these frameworks lack on variety, namely the ability to perform multiple tasks using only one toolkit.

Compression is another important aspect when dealing with high-throughput sequencing data, as it reduces storage space and accelerates data transmission. A survey on DNA compressors and amino acid sequence compression can be found in [16]. Currently, the DNA sequence compressors HiRGC [17], iDo-Comp [18], GeCo [19], and GDC [20] are considered to have the best performance [21]. Of these four

approaches, GeCo is the only one that can be used for reference-free and reference-based compression. Furthermore, GeCo can be used as an analysis tool to determine absolute measures for many distance computations and local measures [19].

Amino acid sequences are known to be very hard to compress [22], however, Hosseini et al. [23] recently developed AC, a state-of-the-art for lossless amino acid sequence compression. In [24] the authors compared the performance of AC, in terms of bit-rate, to several general-purpose lossless compressors and several protein compressors, using different proteomes. They concluded that in average AC provides the best bit-rates.

Another relevant subject is genomic data simulation. Read simulations tools are fundamental for the development, testing and evaluation of methods and computational tools [25, 26]. Despite the availability of a large number of real sequence reads, read simulation data is necessary due to the inability to know the ground truth of real data [27]. Escalona et al. [28], recently, reviewed 23 NGS simulation tools. XS [29], a FASTQ read simulation tool, stands out in relation to the other 22 simulation tools because it is the only one that does not need a reference sequence. Furthermore, XS is the only open-source tool for simulation of FASTQ reads produced by the four most used sequencing machines, Roche-454, Illumina, ABI SOLiD and Ion Torrent.

Although a large number of tools are available for analysing, compressing, and simulation, these tools are specialised in only a specific task. Besides, in many cases the output of one tool cannot be used directly as input for another tool, e.g. the output of a simulation tool cannot always be used directly as input for an analysis tool. Thus, unique software that includes several specialised tools is necessary.

In this document, we describe GTO, a complete toolkit for genomics and proteomics, namely for FASTQ, FASTA and SEQ formats, with many complementary tools. The toolkit is for Unix-based systems, built for ultra-fast computations. GTO supports pipes for easy integration with the sub-programs belonging to GTO as well as external tools. GTO works as LEGOs, since it allows the construction of multiple pipelines with many combinations.

GTO includes tools for information display, randomisation, edition, conversion, extraction, search, calculation, compression, simulation and visualisation. GTO is prepared to deal with very large datasets, typically in the scale of Gigabytes or Terabytes (but not limited). The complete toolkit is an optimised command-line version, using the prefix "gto_" followed by the suffix with the respective name of the program. GTO is implemented in C language and it is available, under the MIT license, at:

```
http://bioinformatics.ua.pt/gto
```

1.1 Installation

To install GTO through the GitHub repository:

```
git clone https://github.com/cobilab/gto.git
cd gto/src/
make
```

Or by installing them directly using the Cobilab channel from Conda:

```
conda install -c cobilab gto --yes
```

1.2 Testing

The examples provided in this document are available in the repository. Therefore, each example can be easily reproduced, which it will also test and validate each tool. To replicate those tests, it can be done in two different ways:

• Running one test for a specific tool:

```
cd gto/tester/gto_{tool}
sh runExample.sh
```

• Running the batch of tests for all the tools:

```
cd gto/tester/
sh runAllTests.sh
```

Some of these tests require internet connection to download external files and it will create new files.

1.3 Execution control

The quality control in Unix/Linux pipelines using GTO's tools is made in three ways:

- 1. Input verification: where the tools verify the format of the input file;
- 2. Stderr logs: Some execution errors are directly sent for the stderr channel.
- 3. Scripting validation: In complex pipelines, the verification of all the tools in the pipeline were executed properly, it is used the PIPESTATUS variable, e.g.:

```
gto_fasta_rand_extra_chars < input.fa | gto_fasta_to_seq > output.seq
echo "${PIPESTATUS[0]} ${PIPESTATUS[1]}"
0 0
```

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Chapter 2

FASTQ tools

The toolkit has a set of tools dedicated to manipulating FASTQ files. Some of these tools allow the data conversion to/from different formats, i. e., there are tools designed to convert a FASTQ file into a sequence or a FASTA/Multi-FASTA format, or converting DNA in some of those formats to FASTQ.

There are also tools for data manipulation in this format, which are designed to exclude 'N', remove low quality scored reads, following different metrics and randomize DNA sequences. Succeeding the manipulation, it is also possible to perform analyses over these files, simulations and mutations. The current available tools for FASTQ format analysis and manipulation include:

- 1. gto_fastq_to_fasta: it converts a FASTQ file format to a pseudo FASTA file.
- 2. gto_fastq_to_mfasta: it converts a FASTQ file format to a pseudo Multi-FASTA file.
- 3. gto_fastq_exclude_n: it discards the FASTQ reads with the minimum number of "N" symbols.
- gto_fastq_extract_quality_scores: it extracts all the quality-scores from FASTQ reads.
- 5. gto_fastq_info: it analyses the basic information of FASTQ file format.
- 6. gto_fastq_maximum_read_size: it filters the FASTQ reads with the length higher than the value defined.
- 7. gto_fastq_minimum_quality_score: it discards reads with average quality-score below of the defined.
- 8. gto_fastq_minimum_read_size: it filters the FASTQ reads with the length smaller than the value defined.
- 9. gto_fastq_rand_extra_chars: it substitues in the FASTQ files, the DNA sequence the outside ACGT chars by random ACGT symbols.
- 10. gto_fastq_from_seq: it converts a genomic sequence to pseudo FASTQ file format.
- 11. gto_fastq_mutate: it creates a synthetic mutation of a FASTQ file given specific rates of mutations, deletions and additions.

- 12. gto_fastq_split: it splits Paired End files according to the direction of the strand ('/1' or '/2').
- 13. gto_fastq_pack: it packages each FASTQ read in a single line.
- 14. gto_fastq_unpack: it unpacks the FASTQ reads packaged using the gto_fastq_pack tool.
- 15. gto_fastq_quality_score_info: it analyses the quality-scores of a FASTQ file.
- 16. gto_fastq_quality_score_min: it analyses the minimal quality-scores of a FASTQ file.
- 17. gto_fastq_quality_score_max: it analyses the maximal quality-scores of a FASTQ file.
- 18. gto_fastq_cut: it cuts read sequences in a FASTQ file.
- 19. gto_fastq_minimum_local_quality_score_forward: it filters the reads considering the quality score average of a defined window size of bases.
- 20. gto_fastq_minimum_local_quality_score_reverse: it filters the reverse reads, considering the average window size score defined by the bases.
- 21. gto_fastq_xs: it is a skilled FASTQ read simulation tool, flexible, portable and tunable in terms of sequence complexity.
- 22. gto_fastq_clust_reads: it agroups reads and creates an index file.
- 23. gto_fastq_complement: it replaces the ACGT bases with their complements in a FASTQ file format.
- 24. gto_fastq_reverse: it reverses the ACGT bases order for each read in a FASTQ file format.
- 25. gto_fastq_variation_map: it identifies the variation that occours in the sequences relative to the reads or a set of reads.
- 26. gto_fastq_variation_filter: it filters and segments the regions of singularity from the output of gto_fastq_variation_map.
- 27. gto_fastq_variation_visual: it depites the regions of singularity using the output from gto_fastq_variation_into an SVG image.
- 28. gto_fastq_metagenomics: it measures similarity between any FASTQ file, independently from the size, against any multi-FASTA database (also under the alias gto fastq falcon).

2.1 Program gto_fastq_to_fasta

The gto_fastq_to_fasta 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:

```
./gto_fastq_to_fasta -h
```

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

Input parameters

The gto_fastq_to_fasta 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 of such an input file is:

Output

The output of the gto_fastq_to_fasta program a FASTA file.

Using the input above, an output example for this is the following:

```
> Computed with Fastq2Fasta
GGGTGATGGCCGCTGCCGATGGCGTCAAATCCCACCAAGTTACCCTTAACAACTTAAGGGTTTTCAAATAGA
GTTCAGGGATACGACGTTTGTATTTTAAGAATCTGAAGCAGAAGTCGATGATAATACGCGTCGTTTTATCAT
```

2.2 Program gto fastq to mfasta

The gto_fastq_to_mfasta 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:

```
./gto_fastq_to_mfasta -h
```

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

Input parameters

The gto_fastq_to_mfasta 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: ./gto_fastq_to_mfasta [options] [[--] args]
    or: ./gto_fastq_to_mfasta [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: ./gto_fastq_to_mfasta < input.fastq > output.mfasta
```

An example of such an input file is:

Output

The output of the gto_fastq_to_mfasta program a Multi-FASTA file. Using the input above, an output example for this is the following:

```
>SRR001666.1 071112_SLXA-EAS1_s_7:5:1:817:345 length=72
GGGTGATGGCCGCTGCCGATGGCGTCAAATCCCACCAAGTTACCCTTAACAACTTAAGGGTTTTCAAATAGA
>SRR001666.2 071112_SLXA-EAS1_s_7:5:1:801:338 length=72
GTTCAGGGATACGACGTTTGTATTTTAAGAATCTGAAGCAGAAGTCGATGATAATACGCGTCGTTTTATCAT
```

2.3 Program gto fastq exclude n

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

For help type:

```
./gto_fastq_exclude_n -h
```

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

Input parameters

The gto_fastq_exclude_n 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: ./gto_fastq_exclude_n [options] [[--] args]
   or: ./gto_fastq_exclude_n [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
                         Input FASTQ file format (stdin)
   > output.fastq
                          Output FASTQ file format (stdout)
Example: ./gto_fastq_exclude_n -m <max> < input.fastq > output.fastq
Console output example :
<FASTQ non-filtered reads>
             : value
Total reads
Filtered reads : value
```

An example of such an input file is:

Output

The output of the gto_fastq_exclude_n program is a set of all the filtered FASTQ reads, followed by the execution report. The execution report only appears in the console.

Using the input above with the max value as 5, an output example for this is the following:

$2.4 \quad Program\ gto\ _fastq_extract_quality_scores$

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

```
./gto_fastq_extract_quality_scores -h
```

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

Input parameters

The gto_fastq_extract_quality_scores 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 of such an input file is:

Output

The output of the gto_fastq_extract_quality_scores program is a set of all the quality scores from the FASTQ reads, followed by the execution report. The execution report only appears in the console.

Using the input above, an output example for this is the following:

2.5 Program gto fastq info

The gto_fastq_info analyses the basic information of FASTQ file format.

For help type:

```
./gto_fastq_info -h
```

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

Input parameters

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

```
Usage: ./gto_fastq_info [options] [[--] args]
   or: ./gto_fastq_info [options]
It analyses the basic information 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: ./gto_fastq_info < input.fastq > output
Output example:
Total reads : value
Max read length : value
Min read length : value
Min QS value : value
Max QS value : value
QS range : value
```

Output

The output of the gto_fastq_info program is a set of information related to the file read.

Using the input above, an output example for this is the following:

```
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 gto fastq maximum read size

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

```
./gto_fastq_maximum_read_size -h
```

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

Input parameters

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

```
Usage: ./gto_fastq_maximum_read_size [options] [[--] args]
    or: ./gto_fastq_maximum_read_size [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
```

Output

The output of the gto_fastq_maximum_read_size program is a set of all the filtered FASTQ reads, followed by the execution report. The execution report only appears in the console.

Using the input above with the size values as 60, an output example for this is the following:

2.7 Program gto_fastq_minimum_quality_score

The gto_fastq_minimum_quality_score discards reads with average quality-score below of the defined. For help type:

```
./gto_fastq_minimum_quality_score -h
```

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

Input parameters

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

```
Usage: ./gto_fastq_minimum_quality_score [options] [[--] args]
   or: ./gto_fastq_minimum_quality_score [options]
It discards reads with average quality-score below value.
    -h, --help
                          show this help message and exit
Basic options
   -m, --min=<int>
                          The minimum average quality-score (Value 25 or 30 is commonly used)
    < input.fastq</pre>
                        Input FASTQ file format (stdin)
    > output.fastq
                          Output FASTQ file format (stdout)
Example: ./gto_fastq_minimum_quality_score -m <min> < input.fastq > output.fastq
Console output example:
<FASTQ non-filtered reads>
Total reads
              : value
Filtered reads : value
```

Output

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

Using the input above with the minimum averge value as 30, an output example for this is the following:

$2.8 \quad Program\ gto_fastq_minimum_read_size$

The gto_fastq_minimum_read_size filters the FASTQ reads with the length smaller than the value defined. For help type:

```
./gto_fastq_minimum_read_size -h
```

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

Input parameters

The gto_fastq_minimum_read_size 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: ./gto_fastq_minimum_read_size [options] [[--] args]
   or: ./gto_fastq_minimum_read_size [options]
It filters the FASTQ reads with the length smaller 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 minimum read length
    < input.fastq</pre>
                          Input FASTQ file format (stdin)
                          Output FASTQ file format (stdout)
   > output.fastq
Example: ./gto_fastq_minimum_read_size -s <size> < input.fastq > output.fastq
Console output example:
<FASTQ non-filtered reads>
Total reads
            : value
Filtered reads : value
```

An example of such an input file is:

Output

The output of the gto_fastq_minimum_read_size program is a set of all the filtered FASTQ reads, followed by the execution report. The execution report only appears in the console.

Using the input above with the size values as 65, an output example for this is the following:

2.9 Program gto fastq rand extra chars

The gto_fastq_rand_extra_chars substitues in the FASTQ files, the DNA sequence the outside ACGT chars by random ACGT symbols.

For help type:

```
./gto_fastq_rand_extra_chars -h
```

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

Input parameters

The gto_fastq_rand_extra_chars 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 of such an input file is:

Output

The output of the gto_fastq_rand_extra_chars program is a FASTQ file.

Using the input above, an output example for this is the following:

2.10 Program gto fastq from seq

The gto_fastq_from_seq converts a genomic sequence to pseudo FASTQ file format. For help type:

```
./gto_fastq_from_seq -h
```

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

Input parameters

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

The attribution is given according to:

An example of such an input file is:

Output

The output of the gto_fastq_from_seq program is a pseudo FASTQ file.

An example, using the size line as 80 and the read's header as "SeqToFastq", for the input, is:

SeqToFastq1 CAAGACGGCCTCCTGCTGCTGCTGCTCTCCGGGGCCACGGCCCTGGAGGGTCCACCGCTGCCCTC FFFFFFFFFFFFFFFFFFFFFFFFFFF	FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	
CAAGACGGCCTCCTGCTGCTGCTGCTCCCGGGGCCACGGCCCTGGAGGGTCCACCGCTGCCCTC FFFFFFFFFFFFFFFFFFFFFFFFFFF	FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	
FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	
SeqToFastq2 GCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCA FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	AGGAAGCGGCAGGAA FFFFFFFFFFFFFFFFFFFFFFF	
SeqToFastq2 GCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCA FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	AGGAAGCGGCAGGAA FFFFFFFFFFFFFFFFFFFFFFF	
GCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCA FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	
FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	
FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	AAGCAGGCCAGTGCC FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	
SeqToFastq3 TGGTTTGAGTGGACCTCCGGGCCCCTCATAGGAGAGGAAGCTCGGGAGGTGGCCAGGCGGCAGGA FFFFFFFFFFFFFFFFFFFFFFFF	AAGCAGGCCAGTGCC FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	
TGGTTTGAGTGGACCTCCGGGCCCCTCATAGGAGAGGAAGCTCGGGAGGTGGCCAGGCGGCAGGAGGTTGGCTAGGAGAGATTTGAGGAGAGGTTGAGGAGAGAGA	CTCCACCCCCCAGC CFFFFFFFFFFFFFFFFFFFFFFFFFF	
FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	CTCCACCCCCCAGC CFFFFFFFFFFFFFFFFFFFFFFFFFF	
FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	CTCCACCCCCCAGC FFFFFFFFFFFFFFFFFFFFFFFFFFF	
SeqToFastq4 CGAATCCGCGCGCGCGGACAGAATCTCCTGCAAAGCCCTGCAGGAACTTCTTCTGGAAGACCTTC FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	CTCCACCCCCCAGC FFFFFFFFFFFFFFFFFFFFFFFFFFF	
CGAATCCGCGCGCCGGGACAGAATCTCCTGCAAAGCCCTGCAGGAACTTCTTCTGGAAGACCTTC FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	CCCGGCCTCCTGCTG	
FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	CCCGGCCTCCTGCTG	
FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	CCCGGCCTCCTGCTG	
SeqToFastq5 AAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAAACAAGATGCCATTGTCC FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	CCCGGCCTCCTGCTG	
AAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAAACAAGATGCCATTGTCCC		
FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF		
	FFFFFFFFFFFF	
	FFFFFFFFFFFF	
South Frank		
Seq10rastq0		
TGCTGCTCTCCGGGGCCACGGCCACCGCTGCCCTGCAGGGTGGCCCCACCGGCCGAGA	CAGCGAGCATATGCA	
FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	FFFFFFFFFFFF	
SeqToFastq7		
GAAGCGGCAGGAATAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGTGGTTTGAGTGGACCT	CCCAGGCCAGTGCCG	
FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	FFFFFFFFFFFF	
SeqToFastq8		
GCCCCTCATAGGAGAGGAAGCTCGGGAGGTGGCCAGGGGGGGAAGGCGCACCCCCCAGCAA	TCCGCGCGCGGGAC	
	FFFFFFFFFFFF	
SeqToFastq9		
GAATGCCCTGCAGGAACTTCTTCTGGAAGACCTTCTCCTCCTGCAAATAAAACCTCACCCATGAA	ATGCTCACGCAAGTT	
FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF		

```
@SeqToFastq10
TAATTACAGACCTGAA
+
FFFFFFFFFFFFFFF
```

2.11 Program gto fastq mutate

The gto_fastq_mutate creates a synthetic mutation of a FASTQ file given specific rates of mutations, deletions and additions. All these parameters are defined by the user, and their are optional. For help type:

```
./gto_fastq_mutate -h
```

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

Input parameters

The gto_fastq_mutate 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 FASTQ File.

The attribution is given according to:

```
Usage: ./gto_fastq_mutate [options] [[--] args]
  or: ./gto_fastq_mutate [options]
Creates a synthetic mutation of a FASTQ file given specific rates of mutations,
deletions and additions
   -h, --help
                              show this help message and exit
Basic options
                              Input FASTQ file format (stdin)
   < input.fasta
   > output.fasta
                              Output FASTQ file format (stdout)
Optional
   -s, --seed=<int>
                             Starting point to the random generator
   -m, --mutation-rate=<dbl>
                             Defines the mutation 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
<insertion rate> -a < input.fastq > output.fastq
```

An example of such an input file is:

Output

The output of the gto_fastq_mutate program is a FASTQ file whith the synthetic mutation of input file. Using the input above with the seed value as 1 and the mutation rate as 0.5, an output example for this is the following:

2.12 Program gto fastq split

The gto_fastq_split splits Paired End files according to the direction of the strand ('/1' or '/2'). It writes by default singleton reads as forward stands.

For help type:

```
./gto_fastq_split -h
```

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

Input parameters

The gto_fastq_split program needs a stream for the computation, namely the input standard. The input stream is a FASTQ file.

```
Usage: ./gto_fastq_split [options] [[--] args]
  or: ./gto_fastq_split [options]

It writes by default singleton reads as forward stands.

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

Output

The output of the gto_fastq_split program is a set of information related to the file read. Using the input above, an output example for this is the following:

```
Total reads : 2
Singleton reads : 0
Forward reads : 65536
Reverse reads : 1
```

Also, this program generates two FASTQ files, with the reverse and forward reads.

An example of the forward reads, for the input, is:

2.13 Program gto fastq pack

The gto_fastq_pack packages each FASTQ read in a single line. It can show the read score first or the dna sequence, depending on the execution mode.

For help type:

```
./gto_fastq_pack -h
```

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

Input parameters

The gto_fastq_pack 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 of such an input file is:

Output

The output of the gto_fastq_pack program is a packaged FASTQ file.

```
Using the input above, an output example for this is the following:

GNNTGATGGCCGCTGCCGATGGCGNANAATCCCACCAANATACCCTTAACAACTTAAGGGTTNTCAAATAGA
```

Another example for the same input, but using the scores first (flag "s"), is:

2.14 Program gto fastq unpack

The gto_fastq_unpack unpacks the FASTQ reads packaged using the gto_fastq_pack tool. For help type:

```
./gto_fastq_unpack -h
```

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

Input parameters

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

The attribution is given according to:

An example of such an input file is:

Output

The output of the gto_fastq_unpack program is a FASTQ file.

Using the input above, an output example for this is the following:

2.15 Program gto fastq quality score info

The gto_fastq_quality_score_info analyses the quality-scores of a FASTQ file. For help type:

```
./gto_fastq_quality_score_info -h
```

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

Input parameters

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

```
Usage: ./gto_fastq_quality_score_info [options] [[--] args]
   or: ./gto_fastq_quality_score_info [options]
It analyses the quality-scores of a FASTQ file.
    -h, --help
                         show this help message and exit
Basic options
   < input.fastq</pre>
                     Input FASTQ file format (stdin)
   > output
                          Output read information (stdout)
Optional
   -m, --max=<int>
                         The lenght of the maximum window
Example: ./gto_fastq_quality_score_info -m <max> < input.fastq > output
Output example :
              : value
Total reads
Max read length : value
Min read length : value
Min QS value : value
```

```
Max QS value : value
QS range : value
```

Output

The output of the gto_fastq_quality_score_info program is a set of information related to the file read. Using the input above with the max window value as 30, an output example for this is the following:

2.16 Program gto fastq quality score max

The gto_fastq_quality_score_max analyses the maximal quality-scores of a FASTQ file. For help type:

```
./gto_fastq_quality_score_max -h
```

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

Input parameters

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

```
Usage: ./gto_fastq_quality_score_max [options] [[--] args]
  or: ./gto_fastq_quality_score_max [options]

It analyses the maximal quality-scores of a FASTQ file.
```

Output

The output of the gto_fastq_quality_score_max program is a set of information related to the file read, considering the maximal quality scores.

Using the input above with the max window value as 30, an output example for this is the following:

$2.17 \quad Program\ gto_fastq_quality_score_min$

The gto_fastq_quality_score_min analyses the minimal quality-scores of a FASTQ file. For help type:

```
./gto_fastq_quality_score_min -h
```

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

Input parameters

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

Output

The output of the gto_fastq_quality_score_min program is a set of information related to the file read, considering the minimum quality scores.

Using the input above with the max window value as 30, an output example for this is the following:

2.18 Program gto_fastq_cut

The gto_fastq_cut cuts read sequences in a FASTQ file. It requires that the initial and end positions for the cut.

For help type:

```
./gto_fastq_cut -h
```

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

Input parameters

The gto_fastq_cut 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 of such an input file is:

Output

The output of the gto_fastq_cut program is a FASTQ file cut.

Using the initial value as 10 and the end value as 30, an example for this input, is:

```
@SRR001666.1 071112_SLXA-EAS1_s_7:5:1:817:345 length=72
CGCTGCCGATGGCGTCAAATC
+
IIIIIIIIIIIIIIIIII
@SRR001666.2 071112_SLXA-EAS1_s_7:5:1:801:338 length=72
ACGACGTTTGTATTTTAAGAA
+
IIIIIIIIIIIIIIIIIIIII
```

2.19 Program gto fastq minimum local quality score forward

The gto_fastq_minimum_local_quality_score_forward filters the reads considering the quality score average of a defined window size of bases.

For help type:

```
./gto_fastq_minimum_local_quality_score_forward -h
```

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

Input parameters

The gto_fastq_minimum_local_quality_score_forward 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: ./gto_fastq_minimum_local_quality_score_forward [options] [[--] args]
   or: ./gto_fastq_minimum_local_quality_score_forward [options]
It filters the reads considering the quality score average of a defined window size
of bases.
   -h, --help
                             show this help message and exit
Basic options
   -k, --windowsize=<int> The window size of bases (default 5)
   -w, --minavg=<int>
                            The minimum average of quality score (default 25)
   -m, --minqs=<int>
                            The minimum value of the quality score (default 33)
   < input.fastq</pre>
                             Input FASTQ file format (stdin)
                             Output FASTQ file format (stdout)
   > output.fastq
Example: ./gto_fastq_minimum_local_quality_score_forward -k <windowsize> -w <minavg>
-m <minqs> < input.fastq > output.fastq
Console output example:
Minimum QS
            : value
<FASTQ output>
Total reads
               : value
Trimmed reads
                : value
```

An example of such an input file is:

Output

The output of the gto_fastq_minimum_local_quality_score_forward program is a FASTQ file with the reads filtered following a quality score average of a defined window of bases. The execution report only

appears in the console.

Using the input above with the default values, an output example for this is the following:

2.20 Program gto fastq minimum local quality score reverse

The gto_fastq_minimum_local_quality_score_reverse filters the reverse reads, considering the quality score average of a defined window size of bases.

For help type:

```
./gto_fastq_minimum_local_quality_score_reverse -h
```

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

Input parameters

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

```
Usage: ./gto_fastq_minimum_local_quality_score_reverse [options] [[--] args]
   or: ./gto_fastq_minimum_local_quality_score_reverse [options]
It filters the reverse reads, considering the quality score average of a defined
window size of bases.
    -h, --help
                              show this help message and exit
Basic options
   -k, --windowsize=<int>
                              The window size of bases (default 5)
   -w, --minavg=<int>
                             The minimum average of quality score (default 25)
    -m, --minqs=<int>
                             The minimum value of the quality score (default 33)
    < input.fastq</pre>
                              Input FASTQ file format (stdin)
                              Output FASTQ file format (stdout)
    > output.fastq
Example: ./gto_fastq_minimum_local_quality_score_reverse -k <windowsize> -w <minavg>
-m <minqs> < input.fastq > output.fastq
```

```
Console output example:
Minimum QS : value
<FASTQ output>
Total reads : value
Trimmed reads : value
```

Output

The output of the gto_fastq_minimum_local_quality_score_reverse program is a FASTQ file with the reads filtered following a quality score average of a defined window of bases. The execution report only appears in the console.

Using the input above with the default values, an output example for this is the following:

2.21 Program gto fastq xs

The gto_fastq_xs is a skilled FASTQ read simulation tool, flexible, portable (does not need a reference sequence) and tunable in terms of sequence complexity. XS handles Ion Torrent, Roche-454, Illumina and ABI-SOLiD simulation sequencing types. It has several running modes, depending on the time and memory available, and is aimed at testing computing infrastructures, namely cloud computing of large-scale projects, and testing FASTQ compression algorithms. Moreover, XS offers the possibility of simulating the three main FASTQ components individually (headers, DNA sequences and quality-scores). Quality-scores can be simulated using uniform and Gaussian distributions.

For help type:

```
./gto_fastq_xs -h
```

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

Input parameters

The gto_fastq_xs program needs a FASTQ file to compute.

The attribution is given according to:

```
[OPTION]... [FILE]
Usage: ./gto_fastq_xs
System options:
                          give this help
 - 17
                          verbose mode
Main FASTQ options:
-t <sequencingType>
                          type: 1=Roche-454, 2=Illumina, 3=ABI SOLiD, 4=Ion Torrent
-hf <headerFormat>
                        header format: 1=Length appendix, 2=Pair End
 -i n=<instrumentName> the unique instrument name (use n= before name)
                          use the same header in third line of the read
-ls <lineSize>
                          static line (bases/quality scores) size
 -ld <minSize>:<maxSize> dynamic line (bases/quality scores) size
-n <numberOfReads>
                         number of reads per file
DNA options:
-f \langle A \rangle, \langle C \rangle, \langle G \rangle, \langle T \rangle, \langle N \rangle symbols frequency
-rn <numberOfRepeats> repeats: number (default: 0)
 -ri <repeatsMinSize>
                          repeats: minimum size
-ra <repeatsMaxSize> repeats: maximum size
-rm <mutationRate>
                         repeats: mutation frequency
                          repeats: use reverse complement repeats
Quality scores options:
-qt <assignmentType>
                          quality scores distribution: 1=uniform, 2=gaussian
-qf <statsFile>
                         load file: mean, standard deviation (when: -qt 2)
-qc <template>
                          custom template ascii alphabet
Filtering options:
 -eh
                          excludes the use of headers from output
                          excludes the use of optional headers (+) from output
 -eo
                          excludes the use of DNA bases from output
 -ed
 -edb
                          excludes '\n' when DNA bases line size is reached
 -es
                           excludes the use of quality scores from output
Stochastic options:
-s <seed>
                          generation seed
<genFile>
                         simulated output file
Common usage:
./XS -v -t 1 -i n=MySeq -ld 30:80 -n 20000 -qt=1 -qc 33,36,39:43 File
 ./XS -v -ls 100 -n 10000 -eh -eo -es -edb -f 0.3,0.2,0.2,0.3,0.0 -rn 50 -ri 300 -ra 3000 -rm 0.1 File
```

An example of such an input file is:

```
@SRR001666.1 071112_SLXA-EAS1_s_7:5:1:817:345 length=60
GGGTGATGGCCGCTGCCGATGGCGTCAAATCCCACCAAGTTACCCTTAACAACTTAAGGG
+
```

Output

The output of the gto_fastq_xs program is a FASTQ file

Using the input above using the common usage with 5 reads (-n 5), an output example for this is the following:

2.22 Program gto fastq clust reads

The gto_fastq_clust_reads agroups reads and creates an index file. It cluster reads in therms of Seq k-mer Lexicographical order.

For help type:

```
./gto_fastq_clust_reads -h
```

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

Input parameters

The gto_fastq_clust_reads program needs two streams for the computation, namely the input and output standard. The input stream is a FASTQ file. The program sorts the FASTQ reads according to the lexicographic order of the genomic sequences.

```
Usage: ./gto_fastq_clust_reads [options] [[--] args]
    or: ./gto_fastq_clust_reads [options]

It agroups reads and creates an index file.
It cluster reads in therms of Seq k-mer Lexicographical order

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

Basic options
    -c, --ctx=<int>
    < input.fastq Input FASTQ file format (stdin)
    > output.fastq Output FASTQ file format (stdout)

Example: ./gto_fastq_clust_reads -c <ctx> < input.fastq > output.fastq
```

Output

The output of the gto_fastq_clust_reads program is a FASTQ file with clustered reads in therms of the genomic sequence k-mer Lexicographical order. An example, for the output, is:

2.23 Program gto fastq complement

The gto_fastq_complement replaces the ACGT bases with their complements in a FASTQ file format. For help type:

```
./gto_fastq_complement -h
```

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

Input parameters

The gto_fastq_complement 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 of such an input file is:

Output

The output of the gto_fastq_complement program is the FASTQ file with the ACGT base complements. Using the input above, an output example for this is the following:

2.24 Program gto fastq reverse

The gto_fastq_reverse reverses the ACGT bases order for each read in a FASTQ file format. For help type:

```
./gto_fastq_reverse -h
```

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

Input parameters

The gto_fastq_reverse 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 of such an input file is:

Output

The output of the gto_fastq_reverse program is the FASTQ file complement with the flag "(Reversed)" added in the header.

Using the input above, an output example for this is the following:

2.25 Program gto fastq variation map

The gto_fastq_variation_map identifies the variation that occours in the sequences relative to the reads or a set of reads.

For help type:

```
./gto_fastq_variation_map -h
```

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

Input parameters

The gto_fastq_variation_map program needs FASTQ, FASTA or SEQ files to be used as reference and target files.

```
Usage: ./gto_fastq_variation_map <OPTIONS>... [FILE]:<...> [FILE]:<...>
./gto_fastq_variation_map: a tool to map relative singularity regions
The (probabilistic) Bloom filter is automatically set.
  - v
                           verbose mode,
                           about CHESTER,
  - a
  -s <value>
                           bloom size,
 - i
                           use inversions,
                           show positions/words,
 -p
  -k <value>
                           k-mer size (up to 30),
  [rFile1]: <rFile2>: <...> reference file(s),
  [tFile1]:<tFile2>:<...> target file(s).
The reference files may be FASTA, FASTQ or DNA SEQ,
while the target files may be FASTA OR DNA SEQ.
Report bugs to <{pratas,raquelsilva,ap,pjf}@ua.pt>.
```

An example of a reference file (Multi-FASTA format) is:

An example for the target file (FASTQ format) is:

Output

The output of the gto_fastq_variation_map program is a text file identifying the relative regions. Using the inputs above, an output example for this is the following:

2.26 Program gto fastq variation filter

The gto_fastq_variation_filter filters and segments the regions of singularity from the output of gto_fastq_variation_map.

For help type:

```
./gto_fastq_variation_filter -h
```

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

Input parameters

The gto_fastq_variation_filter program needs a the output of gto_fastq_variation_map to compute. The attribution is given according to:

Output

The output of the gto_fastq_variation_filter program is a text file with the coordenates of the segmented regions.

Using the inputs above, an output example for this is the following:

```
#132#132
30:60
90:130
```

$2.27 \quad {\bf Program~gto_fastq_variation_visual}$

The gto_fastq_variation_visual depites the regions of singularity using the output from gto_fastq_variation_filt into an SVG image.

For help type:

```
./gto_fastq_variation_visual -h
```

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

Input parameters

The gto_fastq_variation_visual program needs a the output of gto_fastq_variation_filter to compute.

```
#132#132
30:60
90:130
```

Output

The output of the gto_fastq_variation_visual program is a SVG plot with the maps. In the Figure 2.1 is represented the plot using the input above.



Figure 2.1: gto_fastq_variation_visual execution plot.

2.28 Program gto fastq metagenomics

The gto_fastq_metagenomics is an ultra-fast method to infer metagenomic composition of sequenced reads relative to a database. gto_fastq_metagenomics measures similarity between any FASTQ file (or FASTA), independently from the size, against any multi-FASTA database, such as the entire set of complete genomes from the NCBI. gto_fastq_metagenomics supports single reads, paired-end reads, and compositions of both. It has been tested in many plataforms, such as Illumina MySeq, HiSeq, Novaseq, IonTorrent.

gto_fastq_metagenomics is efficient to detect the presence and authenticate a given species in the FASTQ reads. The core of the method is based on relative data compression. gto_fastq_metagenomics uses variable multi-threading, without multiplying the memory for each thread, being able to run efficiently in a common laptop.

For help type:

```
./gto_fastq_metagenomics -h
```

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

Input parameters

The gto_fastq_metagenomics program needs a FASTQ file to compute.

The attribution is given according to:

```
NAME
      {\tt gto\_fastq\_metagenomics}\ \ {\tt v3.1:}\ \ {\tt a}\ \ {\tt tool}\ \ {\tt to}\ \ {\tt infer}\ \ {\tt metagenomic}\ \ {\tt composition}.
SYNOPSIS
      gto_fastq_metagenomics [OPTION]... [FILE1]:[FILE2]:... [FILE]
SAMPLE
      gto_fastq_metagenomics -v -F -1 47 -Z -y pro.com reads1.fq:reads2.fq DB.fa
DESCRIPTION
      It infers metagenomic sample composition of sequenced reads.
      The core of the method uses a cooperation between multiple
      context and tolerant context models with several depths.
      The reference sequences must be in a multi-FASTA format.
      The sequenced reads must be trimmed and in FASTQ format.
      Non-mandatory arguments:
      - h
                            give this help,
      - F
                            force mode (overwrites top file),
                           display version number,
      – V
      – v
                            verbose mode (more information),
      - Z
                            database local similarity,
                            show compression levels,
      - s
      -1 <level>
                            compression level [1;47],
                          subsampling (default: 1),
      -p <sample>
      -t <top>
                           top of similarity (default: 20),
      -n <nThreads>
                            number of threads (default: 2),
      -x <FILE>
                            similarity top filename,
      -y <FILE>
                            profile filename (-Z must be on).
      Mandatory arguments:
      [FILE1]:[FILE2]:... metagenomic filename (FASTQ),
                            Use ":" for splitting files.
      [FILE]
                            database filename (Multi-FASTA).
COPYRIGHT
      Copyright (C) 2014-2019, IEETA, University of Aveiro.
      This is a Free software, under GPLv3. You may redistribute
      copies of it under the terms of the GNU - General Public
      License v3 <http://www.gnu.org/licenses/gpl.html>.
```

An example of such an input file is:

Output

The output of the gto_fastq_metagenomics program is a CSV file (top.csv) with the highest probability of being contained in the samples. An example for this CSV file is the following:

```
1 66725
           12.263
                    NC_037703.1_Saccharomycodes_ludwigii_strain_Y-8871_mitochondrion
2 66725
           12.263
                    {\tt NC\_037703.1\_Saccharomycodes\_ludwigii\_strain\_Y-8871\_mitochondrion}
3 107123 11.492
                    {\tt NC\_012621.1\_Nakaseomyces\_bacillisporus\_mitochondrion}
                    {\tt NC\_012621.1\_Nakaseomyces\_bacillisporus\_mitochondrion}
4 107123 11.492
                    {\tt NC\_024030.1\_Equus\_przewalskii\_mitochondrial\_DNA}
5 16592 11.153
6 14583 10.851
                    NC_021120.1_Bursaphelenchus_mucronatus_mitochondrion
7 162504 10.607
                    NC_018415.1_Candidatus_Carsonella_ruddii_CS_isolate_Thao2000
8 10315
          10.586
                    NC_016117.1_Mnemiopsis_leidyi_mitochondrion
9 162589 10.550
                    NC_018414.1_Candidatus_Carsonella_ruddii_CE_isolate_Thao2000
10 166163 10.476
                    {\tt NC\_018416.1\_Candidatus\_Carsonella\_ruddii\_HC\_isolate\_Thao2000}
```

Chapter 3

FASTA tools

The FASTA tool subset has similar goals to the FASTQ tools. With these tools, it is possible convert data from different formats to the FASTA and multi-FASTA files, or the opposite. In these tools, there are also features to extract and filter reads based on patterns, which can solve specific problems in genomic analytic workflows. The currently available FASTA tools, for analysis and manipulation, are:

- 1. gto_fasta_to_seq: it converts a FASTA or Multi-FASTA file format to a seq.
- 2. gto_fasta_from_seq: it converts a genomic sequence to pseudo FASTA file format.
- 3. gto_fasta_extract: it extracts sequences from a FASTA file, which the range is defined by the user in the parameters.
- 4. gto_fasta_extract_by_read: 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.
- 5. gto_fasta_info: it shows the readed information of a FASTA or Multi-FASTA file format.
- 6. gto_fasta_mutate: it creates a synthetic mutation of a FASTA file given specific rates of editions, deletions and additions.
- gto_fasta_rand_extra_chars: it substitues in the DNA sequence the outside ACGT chars by random ACGT symbols.
- 8. gto_fasta_extract_read_by_pattern: it extracts reads from a Multi-FASTA file format given a pattern in the header.
- 9. gto_fasta_find_n_pos: it reports the "N" regions in a sequence or FASTA (seq) file.
- 10. gto_fasta_split_reads: it splits a Multi-FASTA file to multiple FASTA files.
- 11. gto_fasta_rename_human_headers: it changes the headers of FASTA or Multi-FASTA file to simple chrX by order, where X is the number.

- 12. gto_fasta_extract_pattern_coords: it extracts the header and coordinates from a Multi-FASTA file format given a pattern/motif in the sequence.
- 13. gto_fasta_complement: it replaces the ACGT bases with their complements in FASTA or Multi-FASTA file format.
- 14. gto_fasta_reverse: it reverses the order of a FASTA or Multi-FASTA file format.
- 15. gto_fasta_variation_map: this tool is an alias to gto_fastq_variation_map tool. Please check the documentation of this tool in the in the section of FASTQ tools.
- 16. gto_fasta_variation_filter: this tool is an alias to gto_fastq_variation_filter tool. Please check the documentation of this tool in the in the section of FASTQ tools.
- 17. gto_fasta_variation_visual: this tool is an alias to gto_fastq_variation_visual tool. Please check the documentation of this tool in the in the section of FASTQ tools.
- 18. gto_fasta_split_streams: it splits and writes a FASTA file into three channels of information: headers, extra and DNA.
- 19. gto_fasta_merge_streams: it merges the three channels of information (headers, extra and DNA) and writes it into a FASTA file.

3.1 Program gto fasta to seq

The gto_fasta_to_seq converts a FASTA or Multi-FASTA file format to a sequence. For help type:

```
./gto_fasta_to_seq -h
```

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

Input parameters

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

```
> output.seq Output sequence file (stdout)

Example: ./gto_fasta_to_seq < input.mfasta > output.seq
```

```
>ABO00264 |acc=ABO00264|descr=Homo sapiens mRNA
ACAAGACGGCCTCCTGCTGCTGCTGCTCCCCGGGGCCACGGCCCTGGAGGGTCCACCGCTGCCCTGCCATTGTCCCC
GGCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCAGGAAGCGGCAGGAA
GTGGTTTGAGTGGACCTCCGGGGCCCCTCATAGGAGAGGCAGGAAGCTCGGGAGGTGGCCAGGCGGCAGGAAGCAGGCCAGTGCC
GCGAATCCGCGCGCGCGGGACAGAATCTCCTGCAAAGCCCTGCAGGAACTTCTTCTGGAAGACCTTCTCCACCCCCCAGC
TAAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAA
>ABO00263 |acc=ABO00263|descr=Homo sapiens mRNA
ACAAGATGCCATTGTCCCCCGGGCCTCCTGCTGCTGCTCCCGGGGCCACCGCCACCGCTGCCCTGCCCCTGGAGGGT
GGCCCCACCGGCCGAGACAGCGAGCATATGCAGGAAGCGGCAGGAATAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTG
GTGGTTTGAGTGGACCTCCCAGGCCAGTGCCGGGCCCCTCATAGGAGAGACTCCGGGAAGCTGCCAGGCCAGGCAGAAG
GCGCACCCCCCCAGCAATCCGCGGGCCCGGGACAGAATGCCCTGCAGGAACTTCTTCTCTCCTCCTCCTGCAAA
TAAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAA
```

Output

The output of the gto_fasta_to_seq program is a group sequence. Using the input above, an output example for this is the following:

ACAAGACGCCTCCTGCTGCTGCTGCTCTCCCGGGGCCACGGCCCTGAAGGTCCACCGCTGCCCTGCTGCCATTGTCCCC
GGCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCAGGAAGCGGCAGGAA
GTGGTTTGAGTGGACCTCCGGGCCCCTCATAGGAGAGGAAGCTCGGGAGGTGGCCAGGCGAGAACCAGGCCAGTGCC
GCGAATCCGCGCGCCGGGACAGAATCTCCTGCAAAGCCCTGCAGGAACTTCTTCTGGAAGACCTTCTCCCACCCCCCAGC
TAAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAAACAAGATGCCATTGTCCCCCGGCCTCCTGCTG
CTGCTGCTCTCCGGGGCCACCGCCACCGCTGCCCCTGGAGGGTGGCCCCACCGGCCGAGACAGCGAGCATATGCA
GGAAGCGGCAGGAATAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGTGGTTTGAGTGGACCTCCCAGGCCAGTGCCG
GGCCCCTCATAGGAGAAGACTCCGGGAAGTGCCCAGGCGAGAATCCGCCCCCCGGGAC
AGAATGCCCTGCAGGAACTTCTTCTGGAAGACCTTCTCCTCCTGCAAATAAAACCTCACCCATGAATGCTCACGCAAGTT
TAATTACAGACCTGAA

3.2 Program gto fasta from seq

The gto_fasta_from_seq converts a genomic sequence to pseudo FASTA file format. For help type:

```
./gto_fasta_from_seq -h
```

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

Input parameters

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

Output

The output of the gto_fasta_from_seq program is a pseudo FASTA file.

Using the input above with the size line as 80 and the read's header as "SeqToFasta", an output example for this is the following:

3.3 Program gto fasta extract

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

For help type:

```
./gto_fasta_extract -h
```

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

Input parameters

The gto_fasta_extract 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:

An example of such an input file is:

Output

The output of the gto_fasta_extract program is a group sequence.

Using the input above with the value 0 as the extraction starting point and the 50 as the ending, an output example for this is the following:

3.4 Program gto fasta extract by read

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

For help type:

```
./gto_fasta_extract_by_read -h
```

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

Input parameters

The gto_fasta_extract_by_read 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:

An example of such an input file is:

```
>ABOOO264 |acc=ABOOO264|descr=Homo sapiens mRNA
ACAAGACGGCCTCCTGCTGCTGCTCCTCCGGGGCCACGGCCCTGGAGGGTCCACCGCTGCCCTGCCATTGTCCCC
GGCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCAGGAAGCGGCAGGAA
GTGGTTTGAGTGGACCTCCGGGCCCCTCATAGGAGAGGAAGCTCGGGAGGTGGCCAGGCGAGAAGCAGGCCAGTGCC
GCGAATCCGCGCGCGCGGGACAGAATCTCCTGCAAAGCCCTGCAGGAACCTTCTTCTGGAAGACCTTCTCCACCCCCCAGC
TAAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAA
>ABOOO263 |acc=ABOOO263|descr=Homo sapiens mRNA
ACAAGATGCCATTGTCCCCCGGGCCTCCTGCTGCTGCTCTCCGGGGCCACCGCCACCGCTGCCCTTGCCCCTGGAGGGT
GGCCCCACCGGCCGAGACATATGCAGGAAGCGGCAGGAATAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTG
GTGGTTTGAGTGGACCTCCCAGGCCAGTGCCGGGCCCCCTGAGGAGAG
```

Output

The output of the gto_fasta_extract_by_read program is FASTA or Multi-FASTA file wiht the extracted sequences.

Using the input above with the value 0 as the extraction starting point and the 50 as the ending, an output example for this is the following:

```
>AB000264 |acc=AB000264|descr=Homo sapiens mRNA
ACAAGACGGCCTCCTGCTGCTGCTCCCGGGGCCACGGCCCTGGAGG
>AB000263 |acc=AB000263|descr=Homo sapiens mRNA
ACAAGATGCCATTGTCCCCCGGCCTCCTGCTGCTGCTGCTCCCCGGGGCC
```

3.5 Program gto fasta info

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

```
./gto_fasta_info -h
```

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

Input parameters

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

```
Usage: ./gto_fasta_info [options] [[--] args]
   or: ./gto_fasta_info [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: ./gto_fasta_info < input.mfasta > 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
```

```
>ABOOO264 |acc=ABOOO264|descr=Homo sapiens mRNA
ACAAGACGGCCTCCTGCTGCTGCTGCTCCCGGGGCCACGGCCCTGGAGGGTCCACCGCTGCCCTGCCATTGTCCCC
GGCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCAGGAAGCGGCAGGAA
GTGGTTTGAGTGGACCTCCGGGCCCCTCATAGGAGAGGAAGCTCGGGAGGTGGCCAGGCGGCAGGAAGCAGGCCAGTGCC
GCGAATCCGCGCGCGCGGGACAGAATCTCCTGCAAAGCCCTGCAGGAACCTTCTTCTGGAAGACCTTCTCCACCCCCCCAGC
TAAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAA
>ABOOO263 |acc=ABOOO263|descr=Homo sapiens mRNA
ACAAGATGCCATTGTCCCCCGGGCCTCCTGCTGCTGCTCCTCCGGGGCCACCGCCACCGCTGCCCTGGAGGGT
GGCCCCACCGGCCGAGACAGCATATGCAGGAAGCCGCACGGCAACAGCCTCCTGACTTTCCTCGCTTG
GTGGTTTGAGTGGACCTCCCAGGCCAGTGCCGGGCCCCCTCATAGGAAGAGCTCCGGGAGGTGGCCAGGCCAGGCAGAAG
GCGCACCCCCCCAGCAATCCGCGCGCCCGGGACAGAATGCCCTGCAGGAACTTCTTCTTCTCGCAAA
```

Output

The output of the gto_fasta_info program is a set of information related to the file read. Using the input above, an output example for this is the following:

```
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
```

3.6 Program gto_fasta_mutate

The gto_fasta_mutate 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:

```
./gto_fasta_mutate -h
```

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

Input parameters

The gto_fasta_mutate 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.

```
Usage: ./gto_fasta_mutate [options] [[--] args]
   or: ./gto_fasta_mutate [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</pre>
                                  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: ./gto_fasta_mutate -s <seed> -e <edit rate> -d <deletion rate> -i
<insertion rate> -a < input.mfasta > output.fasta
```

```
>ABO00264 |acc=AB000264|descr=Homo sapiens mRNA
ACAAGACGGCCTCCTGCTGCTGCTGCTCCCCGGGGCCACGGCCCTGGAGGGTCCACCGCTGCCCTGCCATTGTCCCC
GGCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCAGGAAGCGGCAGGAA
GTGGTTTGAGTGGACCTCCGGGCCCCTCATAGGAAGAGCACCTCGGGAGGTGGCCAGGCGCAGGAAGCAGCCAGTGCC
GCGAATCCGCGCGCGGGACAGAATCTCCTGCAAAGCCCTGCAGGAACTTCTTCTGGAAGACCTTCTCCACCCCCCCAGC
TAAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAA
>AB000263 |acc=AB000263|descr=Homo sapiens mRNA
ACAAGATGCCATTGTCCCCCGGCCTCCTGCTGCTGCTCTCCCGGGGCCACCGCCACCGCTGCCCTGGAGGGT
GGCCCCACCGGCCGAGACAGCGAGCATATGCAGGAAGCGCAGGAATAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTG
GTGGTTTGAGTGGACCTCCCAGGCCAGTGCCGGGCCCCCTCATAGGAGAGACTCCGGGAGGTGGCCAGGCGAGGAAG
GCGCACCCCCCCAGCAATCCGCGGCCCGGGACAGAATGCCCTGCAGGAACTTCTTCTTGGAAGACCTTCTCCTCCTCCCAAA
TAAAACCTCACCCATGAATCCTCACGCAAGTTTAATTACAGACCTGAA
```

Output

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

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

```
>ABOOO264 |acc=ABOOO264|descr=Homo sapiens mRNA
ACGCAACGNATTCCTGCTGATCATANTGTNCCGCNCCCCNGCGACGGGGNCTCNCNNGCACACATNGTACCATTGTCCAC
NCTTNCANGTNANCGCTAGCAGGCTACNGTTTNTCCTCNCCTANNCCAANCNGGCGTNNNTACACTGGCACGTGCAGGCA
TNGGTCGGCNGGNNCCTCCGGNAACGGCACCGGAGACCTCGGNGGNTATACAGGTGTCANGAAACATCCCCGCGNC
GNGTGNCCNNGAANCCANAGAGTATCTCACTCACAACCCTGCGTGCACNTCTAGAGNANGACCTTACNCACCNTCCCNTT
NNGTACCACACCAATGAACGCTGCAGAAAGTCTGTTTNNAGGNGNGCA
>ABOOO263 |acc=ABOOO263|descr=Homo sapiens mRNA
```

ATTTGAAGGCAANCGGNCCAGNAATNCGGNGGGTGCNGCTCNTGTNGGCTACGGNCATCGCGGCCCTGCTNTANTAAGCN
TGAACCACCGNTCGNNGCACTTAGCAATNGCGNAANCCGTCGGCACGGCGGAGACNAANCCGCTANTNNTTTCCCGCTNA
ATGGNTGTACAAGACCNACTANACCANCCTCCGTCACCACACTGGAGCGCANGATGGNNCGCTGNCTAGNAGNCNNTGAG
GCGCTCCNTCCTANAAANCCGTGGNCGAGCNCCCTATGGNAGNGTGGGGGTTTTACCGGAAGACCNTCGNGCCCTATGGG
AGCAATCANAANCTAGAAAGCTTACNGATGGTGANGAANTAGACTANG

3.7 Program gto fasta rand extra chars

The gto_fasta_rand_extra_chars 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:

```
./gto_fasta_rand_extra_chars -h
```

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

Input parameters

The gto_fasta_rand_extra_chars 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 of such an input file is:

```
>ABOOO264 |acc=ABOOO264|descr=Homo sapiens mRNA
ANAAGACGGCCTCCTGCTGCTGCTCCCGGGGCCACGNCCCTGAGGGTCCNCCGCTGCCCTGCTGCCATTGNCNCC
NGCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCNGGAAGCGGCAGGAA
GNGGTTTGAGTGGACCTCCNGGCCCCTCATAGGAGAGGAAGCNNGGGAGGTGGCCAGGCGAGCAGGAAGCAGGCCAGTGNC
GCGAATCCGNGCGCCGGGACAGAATCTCCTGCAAAGCCCTGCAGGAACTTCTTCTGGAAGACCTTCTCCACCCCCCCNNN
TAAANNNTCACCCATGAATGCTCACGCAANTTTAATTACAGACCTGAA
>ABOOO263 |acc=ABOOO263|descr=Homo sapiens mRNA
GCGAATCCGNGCGCCGGGACAGAATCTCCTTCTCCACCCCCCCNNNTGCAAAGCCCTGCAGGAACTTCTTCTGGAAGACC
NGCCCCACCTAAGGAAAAGCAGCCTCCAGGAACTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCNGGAAGCCG
ANAAGACGGCCTCCTGCTGCTGCTCCCGGGGCCACGNCCCTGGCNCCAGGGTCCNCCGCTGCCCTGCTGCCATTGN
GAGGAAGCNNGGGAGGTGGCCAGGCCAGGAAGCAGGCCAGTGNCGNGGTTTGAGTGGACCTCCNGGCCCCTCATAGGA
```

Output

The output of the gto_fasta_rand_extra_chars program is a FASTA or Multi-FASTA file. Using the input above, an output example for this is the following:

3.8 Program gto fasta extract read by pattern

The gto_fasta_extract_read_by_pattern extracts reads from a Multi-FASTA file format given a pattern in the header. Also, this pattern is case insensitive.

For help type:

```
./gto_fasta_extract_read_by_pattern -h
```

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

Input parameters

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

```
>ABOOO264 |acc=ABOOO264|descr=Homo sapiens mRNA
ACAAGACGGCCTCCTGCTGCTGCTCTCCCGGGGCCACGGCCCTGGAGGGTCCACCGCTGCCCTGCTGCCATTGTCCCC
GGCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCAGGAAGCGGCAGGAA
GTGGTTTGAGTGGACCTCCGGGCCCCTCATAGGAGAGGAAGCTCGGGAGGTGGCCAGGCGAGGAAGCAGGCCAGTGCC
GCGAATCCGCGCGCGCGGGACAGATCTCCTGCAAAGCCCTGCAGGAACTTCTTCTGGAAGACCTTCTCCACCCCCCCAGC
TAAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAA
>ABOOO263 |acc=ABOOO263|descr=Homo sapiens mRNA
ACAAGATGCCATTGTCCCCCGGGCCTCCTGCTGCTGCTCCTCCGGGGCCACCGCTGCCCTGCCCCTGGAGGGT
GGCCCCACCGGCCGAGACAGCATATGCAGGAAGCGGCAGGAATAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTG
GTGGTTTGAGTGGACCTCCCAGGCCAGTGCCGGGCCCCTCTATAGGAGAGAGCTCGGGAGGTGGCCAGGCGCAGGAAG
GCGCACCCCCCCAGGAATCCGCGCGCCGGGACAGAATGCCCTGCAGAACTTCTTCTCGGAAGACCTTCTCCTCCTCCTGCAAA
TAAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAA
```

Output

The output of the gto_fasta_extract_read_by_pattern program is a Multi-FASTA file.

Using the input above with the pattern value as "264", an output example for this is the following:

```
>ABOOO264 |acc=ABOOO264|descr=Homo sapiens mRNA
ACAAGACGGCCTCCTGCTGCTGCTCCTCCGGGGCCACGGCCCTGGAGGGTCCACCGCTGCCCTGCTGCCATTGTCCCC
GGCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCAGGAAGCGGCAGGAA
GTGGTTTGAGTGGACCTCCGGGCCCCTCATAGGAGAGGAAGCTCGGGAGGTGGCCAGGCGGCAGGAAGCAGGCCAGTGCC
GCGAATCCGCGCGCGCGGGACAGAATCTCCTGCAAAGCCCTGCAGGAACTTCTTCTGGAAGACCTTCTCCCACCCCCCCAGC
TAAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAA
```

3.9 Program gto fasta find n pos

The gto_fasta_find_n_pos reports the "N" regions in a sequence or FASTA (seq) file. For help type:

```
./gto_fasta_find_n_pos -h
```

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

Input parameters

The gto_fasta_find_n_pos program needs two streams for the computation, namely the input and output standard. The input stream is a FASTA file or a sequence.

```
Usage: ./gto_fasta_find_n_pos [options] [[--] args]
  or: ./gto_fasta_find_n_pos [options]

It reports the 'N' regions in a sequence or FASTA (seq) file.
```

```
>ABO00264 |acc=ABO00264|descr=Homo sapiens mRNA
NCNNNACGGCCTCCTGCTGCTGCTCCTCCGGGGCCACGGCCCTGGAGGGTCCACCGCTGCCCTGCCATTGTCCCC
GNCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCAGGAAGCGGCAGGAA
GTNGTTTGAGTGGACCTCCGGGCCCCTCATAGGAGAGGGAAGCTCGGGAGGTGGCCAGGCGGCAGGAAGCAGCCAGTGCC
GCGAATCCGCGCGCGCGGGACAGATCTCCTGCAAAGCCCTGCAGGAACNTCTTCTGGAAGACCTTCTCCACCCCCCCAGC
TAAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAN
```

Output

The output of the gto_fasta_find_n_pos program is a structured report of "N" appearances in the sequence or FASTA file. The first column is the first position of the "N" appearance, the second is the position of the last "N" in the interval found, and the last column is the count of "N" in this interval.

Using the input above, an output example for this is the following:

```
1 1 1
3 5 3
82 82 1
163 163 1
289 289 1
```

3.10 Program gto_fasta_split_reads

The gto_fasta_split_reads splits a Multi-FASTA file to multiple FASTA files. For help type:

```
./gto_fasta_split_reads -h
```

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

Input parameters

The gto_fasta_split_reads program needs one stream for the computation, namely the input standard. This input stream is a Multi-FASTA file.

```
>ABOOO264 |acc=ABOOO264|descr=Homo sapiens mRNA
ACAAGACGGCCTCCTGCTGCTGCTCTCCCGGGGCCACGGCCCTGGAGGGTCCACCGCTGCCCTGCTGCCATTGTCCCC
GGCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCAGGAAGCGGCAGGAA
GTGGTTTGAGTGGACCTCCGGGCCCCTCATAGGAAGAGCACCTCGGGAGGTGGCCAGGCCAGGAAGCAGCCAGTGCC
GCGAATCCGCGCGCGCGGGACAGAATCTCCTGCAAAGCCCTGCAGGAACTTCTTCTGGAAGACCTTCTCCACCCCCCCAGC
TAAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAA
>ABOOO263 |acc=ABOOO263|descr=Homo sapiens mRNA
ACAAGATGCCATTGTCCCCCGGGCCTCCTGCTGCTGCTCCTCCGGGGCCACCGCCACCGCTGCCCTGGAGGGT
GGCCCCACCGGCCGAGACAGCGAGCATATGCAGGAAGCGGCAGGAATAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTG
GTGGTTTGAGTGGACCTCCCAGGCCAGTGCCCGGGCCCCTCATAGGAGAGCTCCGGGAGGTGGCCAGGCGCAGGAAG
GCGCACCCCCCCAGCAATCCGCGCGCCCGGGACAGAATGCCCTGCAGGAACTTCTTCTCGGAAGACCTTCTCCTCCTGCAAA
```

Output

The output of the gto_fasta_split_reads program is a report summary of the execution, and the files created in the defined location.

Using the input above, an output example for this is the following:

```
1 : Splitting to file:./out1.fasta
2 : Splitting to file:./out2.fasta
```

3.11 Program gto fasta rename human headers

The gto_fasta_rename_human_headers changes the headers of FASTA or Multi-FASTA file to simple chrX by order, where X is the number.

For help type:

```
./gto_fasta_rename_human_headers -h
```

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

Input parameters

The gto_fasta_rename_human_headers 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: ./gto_fasta_rename_human_headers [options] [[--] args]
  or: ./gto_fasta_rename_human_headers [options]

It changes the headers of FASTA or Multi-FASTA file to simple chr$1 by order.

  -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: ./gto_fasta_rename_human_headers < input.mfasta > output.mfasta
```

An example of such an input file is:

```
> ABOOO264 | acc = ABOOO264 | descr = Homo sapiens mRNA

ACAAGACGGCCTCCTGCTGCTGCTCTCCCGGGGCCACGGCCCTGAGGGTCCACCGCTGCCCTGCCATTGTCCCC

GGCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCAGGAAGCGGCAGGAA

GTGGTTTGAGTGGACCTCCGGGCCCCTCATAGGAGAGGAAGCTCGGGAGGTGGCCAGGCGGCAGGAAGCAGGCCAGTGCC

GCGAATCCGCGCGCGCGGGACAGAATCTCCTGCAAAGCCCTGCAGGAACTTCTTCTGGAAGACCTTCTCCACCCCCCAGC

TAAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAA

> ABOOO263 | acc = ABOOO263 | descr = Homo sapiens mRNA

ACAAGATGCCATTGTCCCCCGGGCCTCCTGCTGCTGCTGCTCCCCGGGGCCACCGCTGCCCTGCCCCTGGAGGGT

GGCCCCACCGGCCGAGACAGCATATGCAGGAAGCGGCAGGAATAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTG

GTGGTTTGAGTGGACCTCCCAGGCCAGTGCCGGGCCCCTCATAGGAGAGAGCTCCGGGAGGTGGCCAGGCGGCAGGAAG

GCGCACCCCCCCAGGAATCCGCGCGCCGGGACAGAATGCCCTGCAGAACTTCTTCTCGGAAGACCTTCTCCTCCTGCAAA

TAAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAA
```

Output

The output of the gto_fasta_rename_human_headers program is a FASTA or Multi-FASTA file. Using the input above, an output example for this is the following:

```
>chr1

ACAAGACGGCCTCCTGCTGCTGCTCCTCCGGGGCCACGGCCCTGGAGGGTCCACCGCTGCCCTGCTGCCATTGTCCCC

GGCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCAGGAAGCGGCAGGAA

GTGGTTTGAGTGGACCTCCGGGCCCCTCATAGGAGAGGAAGCTCGGGAGGTGGCCAGGCGGCAGGAAGCCAGTGCC

GCGAATCCGCGCGCCGGGACAGAATCTCCTGCAAAGCCCTGCAGGAACTTCTTCTGGAAGACCTTCTCCCACCCCCCCAGC

TAAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAA

>chr2

ACAAGATGCCATTGTCCCCCGGCCTCCTGCTGCTGCTCCCCGGGGCCACGGCCACCGCTGCCCTTGCCCCTGGAGGGT

GGCCCCACCGGCCGAGACAGCATATGCAGGAAGCGGCAGGAATAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTG

GTGGTTTGAGTGGACCTCCCAGGCCAGTGCCGGGCCCCTCATAGGAGAGAGCTCGGGAGGTGGCCAGGCGAGGAAG
```

3.12 Program gto fasta extract pattern coords

The gto_fasta_extract_pattern_coords extracts the header and coordinates from a Multi-FASTA file format given a pattern/motif in the sequence.

For help type:

```
./gto_fasta_extract_pattern_coords -h
```

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

Input parameters

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

The attribution is given according to:

An example of such an input file is:

```
>ABO00264 |acc=ABO00264|descr=Homo sapiens mRNA
ACAAGACGGCCTCCTGCTGCTGCTCCTCCGGGGCCACGGCCCTGGAGGGTCCACCGCTGCCCTGCCATTGTCCCC
GGCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCAGGAAGCGGCAGGAA
GTGGTTTGAGTGGACCTCCGGGCCCCTCATAGGAGAGGAAGCTCGGGAGGTGGCCAGGCGGCAGGAAGCAGCCAGTGCC
GCGAATCCGCGCGCCCGGGACAGATCTCCTGCAAAGCCCTGCAGGAACTTCTTCTGGAAGACCTTCTCCACCCCCCCAGC
TAAAACCTCACCCATGAATGCTCGCAACACGCAAGTTTAATTCGCAAGTTAGACCTGAACGGGAGGTGGCCACGCAAGTT
```

Output

The output of the gto_fasta_extract_pattern_coords program is a Multi-FASTA file. Using the input above, with the pattern ACA, an output example for this is the following:

```
1 3 >AB000264 |acc=AB000264|descr=Homo sapiens mRNA
131 133 >AB000264 |acc=AB000264|descr=Homo sapiens mRNA
259 261 >AB000264 |acc=AB000264|descr=Homo sapiens mRNA
347 349 >AB000264 |acc=AB000264|descr=Homo sapiens mRNA
```

3.13 Program gto_fasta complement

The gto_fasta_complement replaces the ACGT bases with their complements in FASTA or Multi-FASTA file format.

For help type:

```
./gto_fasta_complement -h
```

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

Input parameters

The gto_fasta_complement 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: ./gto_fasta_complement [options] [[--] args]
    or: ./gto_fasta_complement [options]

It replaces the ACGT bases with their complements in FASTA or Multi-FASTA file format.

-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: ./gto_fasta_complement < input.mfasta > output.mfasta
```

An example of such an input file is:

Output

The output of the gto_fasta_complement program is FASTA or Multi-FASTA file with the ACGT base complements.

Using the input above, an output example for this is the following:

3.14 Program gto fasta reverse

The gto_fasta_reverse reverses the ACGT bases order for each read in a FASTA or Multi-FASTA file format.

For help type:

```
./gto_fasta_reverse -h
```

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

Input parameters

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

```
>ABOOO264 |acc=ABOOO264|descr=Homo sapiens mRNA
ACAAGACGGCCTCCTGCTGCTGCTCTCCCGGGGCCACGGCCCTGGAGGGTCCACCGCTGCCCTGCTGCCATTGTCCC
CGGCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCAGGAAGCGGCAGG
AAGTGGTTTGAGTGGACCTCCGGGCCCCTCATAGGAGAGAGCTCGGGAGGTGGCCAGGCGGCAGGAAGCAGCCAGT
GCCGCGAATCCGCGCGCCGGGACAGAATCTCCTGCAAAGCCCTGCAGGAACCTTCTTCTGGAAGACCTTCTCCACCCCC
CAGCTAAAACCTCACCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAA
>ABOOO263 |acc=ABOOO263|descr=Homo sapiens mRNA
ACAAGATGCCATTGTCCCCCGGCCTCCTGCTGCTGCTCCTCCGGGGCCACCGCCACCGCTGCCCCTGGAGGG
TGGCCCCACCGGCCGAGACAGCGAGCATATGCAGGAAGCCGCCAGGAAATAAGGAAAAGCAGCCTCCTGACTTTCCTCGCT
TGGTGGTTTGAGTGGACCTCCCAGGCCAGTGCCGGGCCCCTCATAGGAGAAGCAGCTCCGGAAGACCTTCTCCTCCTC
CAAATAAAACCTCACCCCATGAATGCTCACGCAAGTTTAATTACAGACCTGAA
```

Output

The output of the gto_fasta_reverse program is FASTA or Multi-FASTA file with the bases reversed and the flag "(Reversed)" added in the header.

Using the input above, an output example for this is the following:

3.15 Program gto fasta split streams

The gto_fasta_split_streams splits and writes a FASTA file into three channels of information: headers, extra and DNA.

For help type:

```
./gto_fasta_split_streams -h
```

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

Input parameters

The gto_fasta_split_streams program needs the input standard stream for the computation. The input stream is a FASTA or Multi-FASTA file.

The attribution is given according to:

```
Usage: ./gto_fasta_split_streams [options] [[--] args]
or: ./gto_fasta_split_streams [options]

It splits and writes a FASTA file into three channels of information: headers, extra and DNA.

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

Basic options
-e, --extra=<str>
output file for the extra information
-d, --dna=<str>
Output file for the DNA information
-H, --headers=<str>
Output file for the headers information
<input.fastq Input FASTA file format (stdin)

Example: ./gto_fasta_split_streams -e <filename> -d <filename> -H <filename> < input.fasta
```

An example of such an input file is:

Output

The output of the gto_fasta_split_streams program are three files containing the headers, extra information and DNA. The name of those files can be passed in the tool's parameters. The default names are HEADERS.JV2, EXTRA.JV2 and DNA.JV2.

3.16 Program gto_fasta_merge_streams

The gto_fasta_merge_streams merges the three channels of information (headers, extra and DNA) and writes it into a FASTA file.

For help type:

```
./gto_fasta_merge_streams -h
```

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

Input parameters

The gto_fasta_merge_streams program needs the three files resulting from the execution of the gto_fasta_split_str tool, and the output standard stream for computation. The output stream is a FASTA or Multi-FASTA file.

The attribution is given according to:

```
Usage: ./gto_fasta_merge_streams [options] [[--] args]
or: ./gto_fasta_merge_streams [options]

It merges the three channels of information (headers, extra and DNA) and writes it into a FASTA file.

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

Basic options
-e, --extra=<str>
-e, --extra=<str>
Output file for the extra information
-d, --dna=<str>
Output file for the DNA information
-H, --headers=<str>
Output file for the headers information
> output Output FASTA file format (stdout)

Example: ./gto_fasta_merge_streams -e <filename> -d <filename> -H <filename> > output.fasta
```

Output

The output of the gto_fasta_merge_streams program is a FASTA or Multi-FASTA file.

Using the three output files of the gto_fasta_split_streams tool as input in this example, the output of this tool is the following:

```
>AB000264 |acc=AB000264|descr=Homo sapiens mRNA

ACAAGACGGCCTCCTGCTGCTGCTGCTCCCGGGGCCCACGGCCCTGGAGGGTCCACCGCTGCCCTGCCATTGTCCCCGGCCCCACCTAAGGAAAAGCAGCCTCCT

>AB000263 |acc=AB000263|descr=Homo sapiens mRNA

ACAAGATGCCATTGTCCCCCGGCCTCCTGCTGCTGCTGCTCCCCGGGGCCACCGCCCACCGCCCCCTGCCCCTGGAGGGTGGCCCCACCGGCCGAGACAGCGAGCATA
```

Chapter 4

Genomic sequence tools

The Genomic Sequence subset works directly with the DNA sequences, without any standard format. These tools allow the data extraction, summarising and some mathematical operations over those files. Usually, these are used in the pipeline as a complementary tool. The current available genomic sequence tools, for analysis and manipulation, are:

- 1. gto_genomic_gen_random_dna: it generates a synthetic DNA.
- 2. gto_genomic_rand_seq_extra_chars: it substitues in the DNA sequence the outside ACGT chars by random ACGT symbols.
- 3. gto_genomic_dna_mutate: it creates a synthetic mutation of a sequence file given specific rates of mutations, deletions and additions.
- 4. gto_genomic_extract: it extracts sequences from a sequence file, which the range is defined by the user in the parameters.
- 5. gto_genomic_period: it calculates the best order depth of a sequence, using FCMs.
- 6. gto_genomic_count_bases: it counts the number of bases in sequence, FASTA or FASTQ files.
- 7. gto_genomic_compressor: it compress and decompress genomic sequences for storage purposes (also under the alias gto geco).
- 8. gto_genomic_complement: it replaces the ACGT bases with their complements in a DNA sequence.
- 9. gto_genomic_reverse: it reverses the ACGT bases order for each read in a sequence file (also under the alias gto reverse).
- 10. gto_genomic_variation_map: this tool is an alias to gto_fastq_variation_map tool. Please check the documentation of this tool in the in the section of FASTQ tools.
- 11. gto_genomic_variation_filter: this tool is an alias to gto_fastq_variation_filter tool. Please check the documentation of this tool in the in the section of FASTQ tools.

12. gto_genomic_variation_visual: this tool is an alias to gto_fastq_variation_visual tool. Please check the documentation of this tool in the in the section of FASTQ tools.

4.1 Program gto_genomic_gen_random_dna

The gto_genomic_gen_random_dna generates a synthetic DNA.

For help type:

```
./gto_genomic_gen_random_dna -h
```

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

Input parameters

The gto_genomic_gen_random_dna program needs one stream for the computation, namely the output standard.

The attribution is given according to:

```
Usage: ./gto_genomic_gen_random_dna [options] [[--] args]
  or: ./gto_genomic_gen_random_dna [options]
It generates a synthetic DNA.
    -h, --help
                              show this help message and exit
Basic options
                              Output synthetic DNA sequence (stdout)
    > output.seq
Optional
    -s, --seed=<int>
                             Starting point to the random generator (Default 0)
    -n, --nSymbols=<int>
                             Number of symbols generated (Default 100)
    -f, --frequency=<str>
                             The frequency of each base. It should be represented
                              in the following format: <fa,fc,fg,ft>.
Example: ./gto_genomic_gen_random_dna -s <seed> -n <nsybomls> -f <fa,fc,fg,ft> > output.seq
```

Output

The output of the gto_genomic_gen_random_dna program is a sequence group file whith the synthetic DNA.

Using the input above with the seed value as 1 and the number of symbols as 400, an output example for this is the following:

4.2 Program gto genomic rand seq extra chars

The gto_genomic_rand_seq_extra_chars substitues in the DNA sequence the outside ACGT chars by random ACGT symbols. It works in sequence file formats.

For help type:

```
./gto_genomic_rand_seq_extra_chars -h
```

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

Input parameters

The gto_genomic_rand_seq_extra_chars 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:

```
Usage: ./gto_genomic_rand_seq_extra_chars [options] [[--] args]
    or: ./gto_genomic_rand_seq_extra_chars [options]

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

It works in sequence file formats

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

Basic options
    < input.seq Input sequence file (stdin)
    > output.seq Output sequence file (stdout)

Example: ./gto_genomic_rand_seq_extra_chars < input.seq > output.seq
```

An example of such an input file is:

Output

The output of the gto_genomic_rand_seq_extra_chars program is a sequence file. Using the input above, an output example for this is the following:

4.3 Program gto genomic dna mutate

The gto_genomic_dna_mutate creates a synthetic mutation of a sequence file given specific rates of mutations, deletions and additions. All these parameters are defined by the user, and their are optional. For help type:

```
./gto_genomic_dna_mutate -h
```

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

Input parameters

The gto_genomic_dna_mutate 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 sequence File.

```
Usage: ./gto_genomic_dna_mutate [options] [[--] args]
   or: ./gto_genomic_dna_mutate [options]
Creates a synthetic mutation of a sequence file given specific rates of mutations,
deletions and additions
    -h, --help
                                  show this help message and exit
Basic options
    < input.seq</pre>
                                  Input sequence file (stdin)
    > output.seq
                                  Output sequence file (stdout)
Optional
    -s, --seed=<int>
                                  Starting point to the random generator
    -m, --mutation-rate=<dbl>
                                 Defines the mutation 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: ./gto_genomic_dna_mutate -s <seed> -m <mutation rate> -d <deletion rate> -i <insertion rate> -a < input.seq > output.seq
```

Output

The output of the gto_genomic_dna_mutate program is a sequence file whith the synthetic mutation of input file.

Using the input above with the seed value as 1 and the mutation rate as 0.5, an output example for this is the following:

4.4 Program gto genomic extract

The gto_genomic_extract extracts sequences from a sequence file, which the range is defined by the user in the parameters.

For help type:

```
./gto_genomic_extract -h
```

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

Input parameters

The gto_genomic_extract 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 sequence file.

Output

The output of the gto_genomic_extract program is a group sequence.

Using the input above with the value 0 as the extraction starting point and the 50 as the ending, an output example for this is the following:

```
TCTTTACTCGCGCGTTGGAGAAATACAATAGTGCGGCTCTGTCTCCTTAT
```

4.5 Program gto_genomic_period

The gto_genomic_period calculates the best order depth of a sequence, using FCMs. It only works "ACGT", while the rest will be discarded.

This application has a dependency to represent the results. It requires the Gnuplot to show the execution result.

For help type:

```
./gto_genomic_period -h
```

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

Input parameters

The gto_genomic_period 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 of such an input file is:

Output

The output of the gto_genomic_period program is a execution report, followed by the plot with this information.

Using the input above, an report example for this is the following:

```
Running order: 1 ... Done!
Running order: 2 ... Done!
Running order: 3 ... Done!
Running order: 4 ... Done!
Running order: 5 ... Done!
Running order: 6 ... Done!
Running order: 7 ... Done!
Running order: 8 ... Done!
Running order: 9 ... Done!
Running order: 10 ... Done!
Running order: 11 ... Done!
Running order: 12 ... Done!
Running order: 13 ... Done!
Running order: 14 ... Done!
Running order: 15 ... Done!
Running order: 16 ... Done!
```

```
Running order: 17 ... Done!
Running order: 18 ... Done!
Running order: 19 ... Done!
Running order: 20 ... Done!
 1 2.246
   2.225
    2.237
    2.079
   1.821
   1.733
    1.717
    1.708
    1.717
    1.712
10
   1.717
11
12
   1.721
13
    1.725
14
   1.729
15
   1.733
16
   1.738
17
   1.742
18
   1.746
   1.75
19
20 1.754
```

In the Figure 4.1 is represented the plot for the execution above.

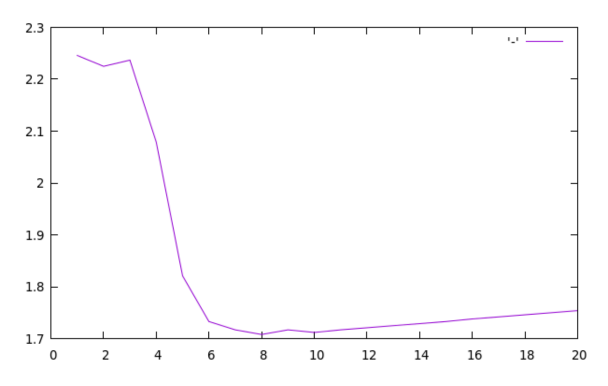


Figure 4.1: gto_genomic_period execution plot.

4.6 Program gto genomic count bases

The gto_genomic_count_bases counts the number of bases in sequence, FASTA or FASTQ files. For help type:

```
./gto_genomic_count_bases -h
```

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

Input parameters

The gto_genomic_count_bases program needs two streams for the computation, namely the input and output standard. The input stream is a sequence, FASTA or FASTQ file.

The attribution is given according to:

```
Usage: ./gto_genomic_count_bases [options] [[--] args]
   or: ./gto_genomic_count_bases [options]
It counts the number of bases in sequence, FASTA or FASTQ files.
    -h, --help
               Show this help message and exit
Basic options
   < input
                 Input sequence, FASTA or FASTQ file format (stdin)
   > output
                Output read information (stdout)
Example: ./gto_genomic_count_bases < input.seq > output
Output example :
File type
          : value
Number of bases : value
Number of a/A : value
Number of c/C : value
Number of g/G : value
Number of t/T : value
Number of n/N
              : value
Number of others : value
```

An example of such an input file is:

Output

The output of the gto_genomic_count_bases program is report which describes the number of each base in the file, and the file type.

Using the input above, an output example for this is the following:

```
File type : DNA
Number of bases : 480
Number of a/A : 114
Number of c/C : 116
Number of g/G : 120
Number of t/T : 130
Number of n/N : 0
Number of others : 0
```

4.7 Program gto genomic compressor

The gto_genomic_compressor is able to provide additional compression gains over several top specific tools, while as an analysis tool, it is able to determine absolute measures, namely for many distance computations, and local measures, such as the information content contained in each element, providing a way to quantify and locate specific genomic events.

For help type:

```
./gto_genomic_compressor -h
```

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

Input parameters

The gto_genomic_compressor program needs a sequence to compress.

```
SYNOPSIS
      ./gto_genomic_compressor [OPTION]... -r [FILE] [FILE]:[FILE]:[FILE]:[...]
SAMPLE
      {\tt Run~Compression} \qquad \qquad : \quad ./{\tt gto\_genomic\_compressor} \ \hbox{-v~-l~3 sequence.txt}
                           : ./gto_genomic_decompressor -v sequence.txt.co
      Run Decompression
      Run Information Profile : ./gto_genomic_compressor -v -l 3 -e sequence.txt
DESCRIPTION
      Compress and decompress genomic sequences for storage purposes.
      Measure an upper bound of the sequences entropy.
      Compute information profiles of genomic sequences.
      -h, --help
           usage guide (help menu).
      -V, --version
           Display program and version information.
      -F, --force
           force mode. Overwrites old files.
```

- -v, --verbose
 verbose mode (more information).
- -x, --examples
 show several running examples (parameter examples).
- -s, --show-levels show pre-computed compression levels (configured parameters).
- -e, --estimate it creates a file with the extension ".iae" with the respective information content. If the file is FASTA or FASTQ it will only use the "ACGT" (genomic) sequence.
- -1 [NUMBER], --level [NUMBER]

 Compression level (integer).

 Default level: 5.

 It defines compressibility in balance with computational resources (RAM & time). Use -s for levels perception.
- -tm [NB_C]:[NB_D]:[NB_I]:[NB_H]:[NB_G]/[NB_S]:[NB_E]:[NB_A]

 Template of a target context model.

 Parameters:
 - [NB_C]: (integer [1;20]) order size of the regular context model. Higher values use more RAM but, usually, are related to a better compression score.
 - [NB_D]: (integer [1;5000]) denominator to build alpha, which
 is a parameter estimator. Alpha is given by 1/[NB_D].
 Higher values are usually used with higher [NB_C],
 and related to confiant bets. When [NB_D] is one,
 the probabilities assume a Laplacian distribution.
 - [NB_I]: (integer {0,1,2}) number to define if a sub-program which addresses the specific properties of DNA sequences (Inverted repeats) is used or not. The number 2 turns ON this sub-program without the regular context model (only inverted repeats). The number 1 turns ON the sub-program using at the same time the regular context model. The number O does not contemple its use (Inverted repeats OFF). The use of this sub-program increases the necessary time to compress but it does not affect the RAM.
 - [NB_H]: (integer [1;254]) size of the cache-hash for deeper context models, namely for [NB_C] > 14. When the [NB_C] < 14 use, for example, 1 as a default. The RAM is highly dependent of this value (higher value stand for higher RAM).
 - [NB_G]: (real [0;1)) real number to define gamma. This value represents the decayment forgetting factor of the regular context model in definition.
 - [NB_S]: (integer [0;20]) maximum number of editions allowed to use a substitutional tolerant model with the same memory model of the regular context model with order size equal to [NB_C]. The value 0 stands for

```
turning the tolerant context model off. When the
             model is on, it pauses when the number of editions
             is higher that [NB_C], while it is turned on when
             a complete match of size [NB_C] is seen again. This
             is probabilistic-algorithmic model very usefull to
             handle the high substitutional nature of genomic
             sequences. When [NB_S] > 0, the compressor used more
             processing time, but uses the same RAM and, usually,
             achieves a substantial higher compression ratio. The
             impact of this model is usually only noticed for
             [NB_C] >= 14.
     [NB\_E]: (integer [1;5000]) denominator to build alpha for
             substitutional tolerant context model. It is
             analogous to [NB_D], however to be only used in the
             probabilistic model for computing the statistics of
             the substitutional tolerant context model.
     [NB_A]: (real [0;1)) real number to define gamma. This value
             represents the decayment forgetting factor of the
             substitutional tolerant context model in definition.
             Its definition and use is analogus to [NB_G].
... (you may use several target models with custom parameters)
-rm [NB_C]:[NB_D]:[NB_I]:[NB_H]:[NB_G]/[NB_S]:[NB_E]:[NB_A]
     Template of a reference context model.
     Use only when -r [FILE] is set (referential compression).
     Parameters: the same as in -tm.
... (you may use several reference models with custom parameters)
-r [FILE], --reference [FILE]
     Reference sequence filename ("-rm" are trainned here).
     Example: -r file1.txt.
[FILE]
     Input sequence filename (to compress) -- MANDATORY.
    File(s) to compress (last argument).
     For more files use splitting ":" characters.
     Example: file1.txt:file2.txt:file3.txt.
```

In the following example, it will be downloaded seventeen DNA sequences, and compress and decompress one of the smallest (BuEb). Finally, it compares if the uncompressed sequence is equal to the original.

```
wget http://sweet.ua.pt/pratas/datasets/DNACorpus.zip
unzip DNACorpus.zip
cp DNACorpus/BuEb .
../../bin/gto_genomic_compressor -v -1 2 BuEb
../../bin/gto_genomic_decompressor -v BuEb.co
cmp BuEb BuEb.de -1
```

4.8 Program gto genomic complement

The gto_genomic_complement replaces the ACGT bases with their complements in a DNA sequence. It works in sequence file formats.

For help type:

```
./gto_genomic_complement -h
```

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

Input parameters

The gto_genomic_complement program needs two parameters, namely the input and output standard. The input stream is a sequence file.

The attribution is given according to:

```
Usage: ./gto_genomic_complement [options] [[--] args]
or: ./gto_genomic_complement [options]

It replaces the ACGT bases with their complements in a DNA sequence.

It works in sequence file formats

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

Basic options
<input.seq Input sequence file (stdin)
> output.seq Output sequence file (stdout)

Example: ./gto_genomic_complement < input.seq > output.seq
```

An example of such an input file is:

Output

The output of the gto_genomic_complement program is a group sequence with the ACGT base complements.

Using the input above, an output example for this is the following:

4.9 Program gto genomic reverse

The gto_genomic_reverse reverses the ACGT bases order for each read in a sequence file. For help type:

```
./gto_genomic_reverse -h
```

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

Input parameters

The gto_genomic_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:

```
Usage: ./gto_genomic_reverse [options] [[--] args]
  or: ./gto_genomic_reverse [options]

It reverses the ACGT bases order for each read in a sequence file.

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

Basic options
  < input.seq Input sequence file (stdin)
  > output.seq Output sequence file (stdout)

Example: ./gto_genomic_reverse < input.seq > output.seq
```

An example of such an input file is:

Output

The output of the gto_genomic_reverse program is a group sequence. Using the input above, an output example for this is the following:

Chapter 5

Amino acid sequence tools

A more specific subset of tools is the Amino Acid Sequence tools, designed to manipulate amino acid sequences. The main features of those tools are grouping sequences, for instance by their properties, such as electric charge (positive and negative), uncharged side chains, hydrophobic side chains and special cases. It is also possible generating pseudo-DNA with characteristics passed by amino acid sequences, or for data compression, using cooperation between multiple contexts and substitutional tolerant context models. The current available amino acid sequence tools, for analysis and manipulation, are:

- 1. gto_amino_acid_to_group: it converts an amino acid sequence to a group sequence.
- 2. gto_amino_acid_to_pseudo_dna: it converts an amino acid (protein) sequence to a pseudo DNA sequence.
- 3. gto_amino_acid_compressor: it is a new lossless compressor to compress efficiently amino acid sequences (proteins).
- 4. gto_amino_acid_from_fasta: it converts DNA sequences in FASTA or Multi-FASTA file format to an amino acid sequence.
- 5. gto_amino_acid_from_fastq: it converts DNA sequences in the FASTQ file format to an amino acid sequence.
- 6. gto_amino_acid_from_seq: it converts DNA sequences to an amino acid sequence.

5.1 Program gto_amino_acid_to_group

The gto_amino_acid_to_group converts an amino acid sequence to a group sequence. For help type:

```
./gto_amino_acid_to_group -h
```

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

Input parameters

The gto_amino_acid_to_group 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: ./gto_amino_acid_to_group [options] [[--] args]
   or: ./gto_amino_acid_to_group [options]
It converts a amino acid sequence to a group sequence.
    -h, --help
                         show this help message and exit
Basic options
   < input.prot</pre>
                          Input amino acid sequence file (stdin)
   > output.group
                        Output group sequence file (stdout)
Example: ./gto_amino_acid_to_group < input.prot > output.group
Table:
       Group
Prot
       P
       P Amino acids with electric charged side chains: POSITIVE
K
       Р
D
       N
       N Amino acids with electric charged side chains: NEGATIVE
Ε
S
        U
Т
       U
N
       U
          Amino acids with electric UNCHARGED side chains
Q
       IJ
С
        S
IJ
       S
G
       S Special cases
Р
       S
       Н
Α
V
        Н
Ι
        Н
L
       Η
M
       H Amino acids with hydrophobic side chains
       Н
F
Y
       Н
W
        Η
        * Others
X
        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 of such an input file is:

```
IPFLLKKQFALADKLVLSKLRQLLGGRIKMMPCGGAKLEPAIGLFFHAIGINIKLGYGMTETTATVSCWHDFQFNPNSIG\\ TLMPKAEVKIGENNEILVRGGMVMKGYYKKPEETAQAFTEDGFLKTGDAGEFDEQGNLFITDRIKELMKTSNGKYIAPQY
```

Output

The output of the gto_amino_acid_to_group program is a group sequence.

Using the input above, an output example for this is the following:

HSHHHPPUHHHHNPHHHUPHPUHHSSPHPHHSSSSHPHNSHHSHHPHHSHUHPHSHSHUNUUHUHUSHPNHUHUSUUHS UHHSPHNHPHSNUUNHHHPSSHHHPSHHPPSNNUHUHHUNNSHHPUSNHSNHNNUSUHHHUNPHPNHHPUUUSPHHHSUH HNUPHSPNPHHNUHHHHHNHPPHUHHHHSSHNUHNNHHPUHUHPHPNPHNHHPUUNHHPHHN

5.2 Program gto amino acid to pseudo dna

The gto_amino_acid_to_pseudo_dna converts an amino acid (protein) sequence to a pseudo DNA sequence.

For help type:

```
./gto_amino_acid_to_pseudo_dna -h
```

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

Input parameters

The gto_amino_acid_to_pseudo_dna 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: ./gto_amino_acid_to_pseudo_dna [options] [[--] args]
   or: ./gto_amino_acid_to_pseudo_dna [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: ./gto_amino_acid_to_pseudo_dna < input.prot > output.dna
Table:
Prot
        DNA
        GCA
С
        TGC
D
        GAC
        {\tt GAG}
F
        TTT
G
        GGC
Н
        CAT
Т
        ATC
```

```
K
           A\,A\,A
L
           CTG
Μ
           ATG
N
           AAC
Р
           CCG
Q
           CAG
R
           CGT
S
           TCT
T
           ACG
٧
           GTA
W
           TGG
Y
           TAC
           TAG
           \tt GGG
```

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

```
IPFLLKKQFALADKLVLSKLRQLLGGRIKMMPCGGAKLEPAIGLFFHAIGINIKLGYGMTETTATVSCWHDFQFNPNSIG
TLMPKAEVKIGENNEILVRGGMVMKGYYKKPEETAQAFTEDGFLKTGDAGEFDEQGNLFITDRIKELMKTSNGKYIAPQY
IESKIGKDKFIEQIAIIADAKKYVSALIVPCFDSLEEYAKQLNIKYHDRLELLKNSDILKMFE
```

Output

The output of the gto_amino_acid_to_pseudo_dna program is a DNA sequence. Using the input above, an output example for this is the following:

5.3 Program gto amino acid compressor

The gto_amino_acid_compressor is a new lossless compressor to compress efficiently amino acid sequences (proteins). It uses a cooperation between multiple context and substitutional tolerant context models. The cooperation between models is balanced with weights that benefit the models with better performance according to a forgetting function specific for each model.

For help type:

```
./gto_amino_acid_compressor -h
```

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

Input parameters

The gto_amino_acid_compressor program needs a file with amino acid sequences to compress. The attribution is given according to:

```
Usage: ./gto_amino_acid_compressor [OPTION]... -r [FILE] [FILE]:[...]
Compression of amino acid sequences.
{\tt Non-mandatory\ arguments:}
                                                            give this help,
                                                           show AC compression levels,
     _ 0
                                                           verbose mode (more information),
     - v
     – V
                                                            display version number,
                                                          force overwrite of output,
    -f
    -1 <level>
                                                          level of compression [1;7] (lazy -tm setup),
     -t <threshold>
                                                            threshold frequency to discard from alphabet,
                                                            it creates a file with the extension ".iae"
                                                            with the respective information content.
    -rm < c < d < g / m < c < d < g / m >: < e < d >: < g / m >: < e >: < d >: < f / m >: 
    -tm <c>:<d>:<g>/<m>:<e>:<a> target model (-tm 1:1:0.8/0:0:0),
    -tm <c>:<d>:<g>/<m>:<e>:<a> target model (-tm 7:100:0.9/2:10:0.85),
                                                             target and reference templates use <c> for
                                                            context-order size, \langle d \rangle for alpha (1/\langle d \rangle), \langle g \rangle
                                                            for gamma (decayment forgetting factor) [0;1),
                                                            \mbox{\ensuremath{\mbox{cm}}>} to the maximum sets the allowed mutations,
                                                            on the context without being discarded (for
                                                             deep contexts), under the estimator <e>, using
                                                            <a>> for gamma (decayment forgetting factor)</a>
                                                            [0;1) (tolerant model),
     -r <FILE>
                                                           reference file ("-rm" are loaded here),
Mandatory arguments:
     <FILE>:<...>:<...>
                                                         file to compress (last argument). For more
                                                            files use splitting ":" characters.
Example:
     [Compress]
                                  ./gto_amino_acid_compressor -v -tm 1:1:0.8/0:0:0 -tm 5:20:0.9/3:20:0.9 seq.txt
     [Decompress] ./gto_amino_acid_decompressor -v seq.txt.co
```

In the following example, it will be downloaded nine amino acid sequences and compress and decompress one of the smallest (HI). Finally, it compares if the uncompressed sequence is equal to the original.

```
wget http://sweet.ua.pt/pratas/datasets/AminoAcidsCorpus.zip
unzip AminoAcidsCorpus.zip
cp AminoAcidsCorpus/HI .
```

```
./gto_amino_acid_compressor -v -l 2 HI
./gto_amino_acid_decompressor -v HI.co
cmp HI HI.de
```

5.4 Program gto amino acid from fasta

The gto_amino_acid_from_fasta converts DNA sequences in FASTA or Multi-FASTA file format to an amino acid sequence.

For help type:

```
./gto_amino_acid_from_fasta -h
```

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

Input parameters

The gto_amino_acid_from_fasta 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 of such an input file is:

Output

The output of the gto_amino_acid_from_fasta program is an amino acid sequence.

Using the input above, an output example for this is the following:

 $\label{thmu} TRRPPAAAALRGHGPGGSTAALLPLSPAPPKEKQPPDFPRLGRDSEHMQEAAGSGLSGPPGPS-ERKLGRWPGGRKQAS\\ AANPRAGTESPAKPCRNFFWKTFSTPPAKTSPMNAHASLITDLTRCHCPPASCCCCSPGPRPPLPCPWRVAPPAETASI\\ CRKRQE-GKAAS-LSSLGGLSGPPRPVPGPS-ERKLGRWPGGRKAHPPSNPRAGTECPAGTSSGRPSPPANKTSPMNAH\\ ASLITDL$

5.5 Program gto amino acid from fastq

The gto_amino_acid_from_fastq converts DNA sequences in the FASTQ file format to an amino acid sequence.

For help type:

```
./gto_amino_acid_from_fastq -h
```

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

Input parameters

The gto_amino_acid_from_fastq 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 of such an input file is:

Output

The output of the gto_amino_acid_from_fastq program is an amino acid sequence. Using the input above, an output example for this is the following:

```
G-WPLPMASNPTKLPLTT-GFSNRVQGYDVCILRI-SRSR--YASFYH
```

5.6 Program gto amino acid from seq

The gto_amino_acid_from_seq converts DNA sequence to an amino acid sequence. For help type:

```
./gto_amino_acid_from_seq -h
```

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

Input parameters

The gto_amino_acid_from_seq program needs two streams for the computation, namely the input and output standard. The input stream is a DNA sequence.

```
Usage: ../../bin/gto_amino_acid_from_seq [options] [[--] args]
    or: ../../bin/gto_amino_acid_from_seq [options]

It converts DNA sequence to an amino acid sequence (translation).

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

Basic options
    < input.seq Input sequence file (stdin)
    > output.prot Output amino acid sequence file (stdout)

Optional
    -f, --frame=<int> Translation codon frame (1, 2 or 3)

Example: ../../bin/gto_amino_acid_from_seq < input.seq > output.prot
```

Output

The output of the gto_amino_acid_from_seq program is an amino acid sequence. Using the input above, an output example for this is the following:

 $\label{thm:constraint} TRRPPAAAALRGHGPGGSTAALLPLSPAPPKEKQPPDFPRLGRDSEHMQEAAGSGLSGPPGPS-ERKLGRWPGGRKQAS\\ AANPRAGTESPAKPCRNFFWKTFSTPPAKTSPMNAHASLITDLKQDAIVPRPPAAAALRGHGHRCPAPGGWPHRPRQRA\\ YAGSGRNKEKQPPDFPRLVV-VDLPGQCRAPHRRGSSGGGQAAGRRTPPAIRAPGQNALQELLLEDLLLLQIKPHP-ML <math display="block">TQV-LQT-$

Chapter 6

General purpose tools

The toolkit also has a set of tools with a more general-purpose, which were not designed to work with a specific data format. Instead, it was developed as an auxiliary component to help the construction of pipelines combining all the described subsets. This contains tools for char manipulations, such as reversing, segmentation and permutation, for manipulating numerical scores, such sum, filter, calculate the min and the max of a numeric matrix mainly originated from the tools' outputs. The current available tools for general purposes are:

- 1. gto_char_to_line: it splits a sequence into lines, creating an output sequence which has a char for each line.
- 2. gto_new_line_on_new_x: it splits different rows with a new empty row.
- 3. gto_upper_bound: it sets an upper bound in a file with a value per line.
- 4. gto_lower_bound: it sets an lower bound in a file with a value per line.
- 5. gto_brute_force_string: it generates all combinations, line by line, for an inputted alphabet and specific size.
- 6. gto_real_to_binary_with_threshold: it converts a sequence of real numbers into a binary sequence, given a threshold.
- 7. gto_sum: it adds decimal values in file, line by line, splitted by spaces or tabs.
- 8. gto_filter: it filters numerical sequences.
- 9. gto_word_search: it search for a word in a file.
- 10. gto_permute_by_blocks: it permutates by block sequence, FASTA and Multi-FASTA files.
- 11. gto_info: it gives the basic properties of the file, namely size, cardinality, distribution percentage of the symbols, among others.
- 12. gto_segment: it segments a filtered sequence based on a threshold.

- 13. gto_comparative_map: it creates a visualization for comparative maps.
- 14. gto_max: it computes the maximum value in each row between two files.
- 15. gto_min: it computes the minimum value in each row between two files.

6.1 Program gto char to line

The gto_char_to_line splits a sequence into lines, creating an output sequence which has a char for each line.

For help type:

```
./gto_char_to_line -h
```

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

Input parameters

The gto_char_to_line 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 of such an input file is:

Output

The output of the gto_char_to_line program is a group sequence splited by \n foreach character. Using the input above, an output example for this is the following:

```
С
Α
G
Α
С
G
G
С
С
Т
С
С
T
G
С
Т
G
С
Т
```

6.2 Program gto_new_line_on_new_x

The gto_new_line_on_new_x splits different rows with a new empty row.

For help type:

```
./gto_new_line_on_new_x -h
```

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

Input parameters

The gto_new_line_on_new_x program needs two streams for the computation, namely the input and output standard. The input stream is a matrix file format with 3 columns.

```
Usage: ./gto_new_line_on_new_x [options] [[--] args]
  or: ./gto_new_line_on_new_x [options]

It splits different rows with a new empty row.
  -h, --help show this help message and exit
```

```
      1
      2
      2

      1
      2
      2

      4
      4
      1

      10
      12
      2

      15
      15
      1

      45
      47
      3

      45
      47
      3

      45
      47
      3

      55
      55
      1
```

Output

The output of the gto_new_line_on_new_x program is a 3 column matrix, with an empty line between different rows.

Using the input above, an output example for this is the following:

```
1.000000
          2.000000
                     2.000000
1.000000
          2.000000
                     2.000000
4.000000
          4.000000 1.000000
10.000000
         12.000000 2.000000
15.000000
         15.000000
                    1.000000
45.000000 47.000000 3.000000
45.000000 47.000000 3.000000
45.000000 47.000000 3.000000
45.000000 47.000000
                    3.000000
55.000000 55.000000
                    1.000000
```

6.3 Program gto upper bound

The gto_upper_bound sets an upper bound in a file with a value per line.

For help type:

```
./gto_upper_bound -h
```

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

Input parameters

The gto_upper_bound program needs two streams for the computation, namely the input and output standard. The input stream is a numeric file.

The attribution is given according to:

An example of such an input file is:

```
0.123
3.432
2.341
1.323
7.538
4.122
0.242
0.654
5.633
```

Output

The output of the gto_upper_bound program is a set of numbers truncated at the a defined upper bound. Using the input above, an output example for this is the following:

```
Using upper bound: 4
0.123000
3.432000
2.341000
1.323000
4.000000
4.000000
0.242000
0.654000
4.000000
```

6.4 Program gto lower bound

The gto_lower_bound sets an lower bound in a file with a value per line.

For help type:

```
./gto_lower_bound -h
```

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

Input parameters

The gto_lower_bound program needs two streams for the computation, namely the input and output standard. The input stream is a numeric file.

The attribution is given according to:

An example of such an input file is:

```
0.123
3.432
2.341
1.323
7.538
4.122
0.242
0.654
5.633
```

Output

The output of the gto_lower_bound program is a set of numbers truncated at the a defined lower bound. Using the input above, an output example for this is the following:

```
Using lower bound: 2
2.000000
3.432000
2.341000
```

```
2.000000
7.538000
4.122000
2.000000
2.000000
5.633000
```

6.5 Program gto brute force string

The gto_brute_force_string generates all combinations, line by line, for an inputted alphabet and specific size.

For help type:

```
./gto_brute_force_string -h
```

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

Input parameters

The gto_brute_force_string program needs some parameters for the computation, namely the alphabet and the key size.

The attribution is given according to:

Output

The output of the gto_brute_force_string program is a set of all possible word combinations with a defined size, using the input alphabet.

Using the input above with the alphabet "abAB" with the word size of 3, an output example for this is the following:

```
aaa
aab
aaA
aaB
```

```
aba
...
BBb
BBA
BBB
```

6.6 Program gto_real_to_binary_with_threshold

The gto_real_to_binary_with_threshold converts a sequence of real numbers into a binary sequence, given a threshold. The numbers below to the threshold will be 0.

For help type:

```
./gto_real_to_binary_with_threshold -h
```

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

Input parameters

The gto_real_to_binary_with_threshold program needs two streams for the computation, namely the real sequence as input. These numbers should be splitted by lines.

The attribution is given according to:

```
Usage: ./gto_real_to_binary_with_threshold [options] [[--] args]
  or: ./gto_real_to_binary_with_threshold [options]

It converts a sequence of real numbers into a binary sequence given a threshold.

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

Basic options
  -t, --threshold=<dbl> The threshold in real format
  < input.num Input numeric file (stdin)
  > output.bin Output binary file (stdout)

Example: ./gto_real_to_binary_with_threshold -t <threshold> < input.num > output.bin
```

An example of such an input file is:

```
12.25

1.2

5.44

5.51

7.97

2.34

8.123
```

Output

The output of the gto_real_to_binary_with_threshold program is a binary sequence.

Using the input above with the threshold of 5.5, an output example for this is the following:

```
1
0
0
1
1
1
0
```

6.7 Program gto sum

The gto_sum adds decimal values in file, line by line, splitted by spaces or tabs.

For help type:

```
./gto_sum -h
```

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

Input parameters

The gto_sum program needs two streams for the computation, namely the input, which is a decimal file. The attribution is given according to:

```
Usage: ./gto_sum [options] [[--] args]
   or: ./gto_sum [options]
It adds decimal values in file, line by line, splitted by spaces or tabs.
   -h, --help
                    show this help message and exit
Basic options
                Input numeric file (stdin)
   < input.num
   > output.num
                   Output numeric file (stdout)
Optional
                   When active, the application adds all the values line by line
   -r, --sumrows
   -a. --sumall
                   When active, the application adds all values
Example: ./gto_sum -a < input.num > output.num
```

An example of such an input file is:

```
      0.123
      5
      5

      3.432
      2

      2.341
      3
      2

      1.323
      7.538
      5

      4.122
      0.242

      0.654
      0.654
```

```
5.633 10
```

Output

The output of the gto_sum program is a sum of the elements in the input file.

Executing the application with the provided input and with the flag to add only the elements in each row, the output of this execution is:

```
10.123000
3.432000
7.341000
1.323000
12.538000
4.122000
0.242000
0.654000
15.633000
```

6.8 Program gto_filter

The gto_filter filters numerical sequences using a low-pass filter.

For help type:

```
./gto_filter -h
```

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

Input parameters

The gto_filter program needs two streams for the computation, namely the input and output standard. The input stream is a numeric file.

```
Usage: ./gto_filter [options] [[--] args]
  or: ./gto_filter [options]
It filters numerical sequences using a low-pass filter.
    -h, --help
                              show this help message and exit
Basic options
    < input.num
                              Input numeric file (stdin)
    > output.num
                              Output numeric file (stdout)
Optional
    -w, --windowsize=<int>
                             Window size (defaut 0)
    -d, --drop=<int>
                             Discard elements (default 0.0)
    -t, --windowtype <<int> Window type (0=Hamm, 1=Hann, 2=Black, 3=rec) (default 0 (Hamm))
    -c, --onecolumn
                              Read from one column
```

```
-p, --printone Print one column
-r, --reverse Reverse mode

Example: ./gto_filter -w <windowsize> -d <drop> -t <windowtype> -c -p -r < input.num > output.num
```

```
1 1.77
5 2.18
10 2.32
15 3.15
20 2.52
25 4.43
30 1.23
```

Output

The output of the gto_filter program is a numeric file, identical of the input.

Using the input above with the window size of 3, an output example for this is the following:

```
Got 7 entries from file

1    2.085

5    2.256

10    2.507

15    2.757

20    2.905

25    2.860

30    2.674
```

$6.9 \quad {\bf Program~gto_word_search}$

The gto_word_search search for a word in a file. It is case sensitive.

For help type:

```
./gto_word_search -h
```

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

Input parameters

The gto_word_search program needs two streams for the computation, namely the input and output standard. The input stream is a text file.

```
Usage: ./gto_word_search [options] [[--] args]
  or: ./gto_word_search [options]

Searching for a word in a text file. It is case sensitive.
```

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

Basic options
-w, --word=<str> Word to search in the file
< input.txt Input text file (stdin)
> output.txt Output text file (stdout)

Example: ./gto_word_search -w <word> < input.txt > output.txt
```

```
No guts, no story. Chris Brady
My life is my message. Mahatma Gandhi
Screw it, letâĂŹs do it. Richard Branson
Boldness be my friend. William Shakespeare
Keep going. Be all in. Bryan Hutchinson
My life is my argument. Albert Schweitzer
Fight till the last gasp. William Shakespeare
Leave no stone unturned. Euripides
```

Output

The output of the gto_word_search program is a text file with the matching paragraphs and the location of the word found.

Using the input above with the word "Shakespeare", an output example for this is the following:

```
Found match in range [ 1536 : 2048 ]
Boldness be my friend. William Shakespeare

Found match in range [ 3072 : 3584 ]
Fight till the last gasp. William Shakespeare
```

6.10 Program gto permute by blocks

The gto_permute_by_blocks permutates by block sequence, FASTA and Multi-FASTA files. For help type:

```
./gto_ -h
```

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

Input parameters

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

```
>ABOOO264 |acc=ABOOO264|descr=Homo sapiens mRNA
ACAAGACGGCCTCCTGCTGCTGCTCCTCCGGGGCCACGGCCCTGGAGGGTCCACCGCTGCCCTGCCGTTGCCCATTGTCCCC
GGCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGAGCATATGCAGGAAGCGGCAGGAA
GTGGTTTGAGTGGACCTCCGGGCCCCTCATAGGAGAGGAAGCTCGGGAAGCTGGCCAGGCGAGAGCAGGCCAGTGCC
GCGAATCCGCGCGCGCGGGACAGATCTCCTGCAAAGCCCTGCAGGAACTTCTTCTTGGAAGACCTTCTCCACCCCCCAGC
TAAAACCTCACCCATGAATGCTCGCAACACGCAAGTTTAATTCGCAAGTTAGACCTGAACGGGAGGTGGCCACGCAAGTT
```

Output

The output of the gto_permute_by_blocks program is a sequence, FASTA or Multi-FASTA file permuted following some parameters.

Using the input above with the base number as 80, an output example for this is the following:

6.11 Program gto_info

The gto_info gives the basic properties of the file, namely size, cardinality, distribution percentage of the symbols, among others.

For help type:

```
./gto_info -h
```

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

Input parameters

The gto_info program needs two streams for the computation, namely the input and output standard. The input stream is a file withou any specific format.

The attribution is given according to:

```
Usage: ./gto_info [options] [[--] args]
   or: ./gto_info [options]
It gives the basic properties of the file, namely size, cardinality, distribution
percentage of the symbols, among others.
    -h, --help show this help message and exit
Basic options
   < input Input file (stdin)
> output Output read information (stdout)
Optional
   -a, --ascii When active, the application shows the ASCII codes
Example: ./gto_info < input > output
Output example :
Number of symbols : value
                 : value
Alphabet size
Alphabet
           : value
Symbol distribution:
<Symbol/Code ASCII> <Symbol count> <Distribution percentage>
```

An example of such an input file is:

```
>ABO00264 |acc=AB000264|descr=Homo sapiens mRNA
ACAAGACGGCCTCCTGCTGCTGCTCTCCCGGGGCCCACGGCCCTGGAGGGTCCACCGCTGCCCTGCCATTGTCCCC
GGCCCCACCTAAGGAAAAGCAGCCTCCTGACTTTCCTCGCTTGGGCCGAGACAGCGGAGCATATGCAGGAAGCGGCAGGAA
GTGGTTTGAGTGGACCTCCGGGCCCCTCATAGGAGAGGAAGCTCGGGAGGTGGCCAGGCGGCAGGAAGCAGGCCAGTGCC
GCGAATCCGCGCGCCCGGGACAGATCTCCTGCAAAGCCCTGCAGGAACTTCTTCTGGAAGACCTTCTCCACCCCCCCAGC
TAAAACCTCACCCATGAATGCTCGCAACACGCAAGTTTAATTCGCAAGTTAGACCTGAACGGGAGGTGGCCACGCAAGTT
```

Output

The output of the gto_info program is a set of information related to the file read.

Using the input above, an output example for this is the following:

```
0.22075055
        1
o : 2 0.4415011
n : 1 0.22075058
m : 2 0.4415011
i : 1 0.22075058
e : 2 0.4415011
                   0.22075055
                   0.22075055
d: 1
                 0.22075055
0.66225166
                   0.22075055
        1
N :
                 0.22075055
                 0.22075055
H: 1
G : 117 25.827815
C : 131 28.918322
B : 2 0.4415011
A : 89 19.646799
> : 1 0.22075055
        1
                 0.4415011
= : 2
6 : 2 0.4415011

4 : 2 0.4415011

2 : 2 0.4415011

0 : 6 1.3245033

: 4 0.88300221
\n : 5
                  1.1037528
```

6.12 Program gto segment

The gto_segment segments a filtered sequence based on a threshold.

For help type:

```
./gto_segment -h
```

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

Input parameters

The gto_segment program needs two streams for the computation, namely the input and output standard. The input stream is a numeric file.

```
1 1.77

5 2.18

10 2.32

15 3.15

20 2.52

25 4.43

30 1.23
```

Output

The output of the gto_segment program is the interval of values below the threshold.

Using the input above with a threshold of 3, an output example for this is the following:

```
0:10
```

6.13 Program gto comparative map

The gto_comparative_map creates a visualization for comparative maps.

For help type:

```
./gto_comparative_map -h
```

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

Input parameters

The gto_comparative_map program needs an input file with the plot positions, respecting a defined structure.

```
Usage: ./gto_comparative_map [options] [[--] args]
or: ./gto_comparative_map [options]

It creates a visualization for comparative maps.

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

Basic options
<FILE> Contigs filename with positions (.pos),

Optional

-h Give this help,
```

```
– V
                         Display version number,
    - v
                         Verbose mode (more information),
    -1 <link>
                         Link type between maps [0;4],
    -w <width>
                        Chromosome width,
    -s <space>
                        Space between chromosomes,
   -m <mult>
                        Color id multiplication factor,
   -b <begin>
                        Color id beggining,
    -c <minimum>
                         Minimum block size to consider,
                        Do NOT show inversion maps,
   - i
                        Do NOT show regular maps,
   -r
    -o <FILE>
                         Output image filename with map,
Example: ./gto_comparative_map -o map.svg map.config
```

```
#SCF 5000000 5000000

aaa 1 1000000 1 1000000 bbbb 3000000 4000000 3000000 4000000

bbb 1500000 2000000 1500000 2000000 cccc 1500000 2000000 1500000 2000000

aaa 2000000 3000000 2000000 3000000 bbbb 3000000 2000000 3000000 2000000
```

Output

The output of the gto_comparative_map program is a executing report, and a svg plot with the maps. Using the input above, an output example for this is the following:

In the Figure 6.1 is represented the plot for the execution above.

6.14 Program gto max

The gto_max computes the maximum value in each row between two files. For help type:

```
./gto_max -h
```

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

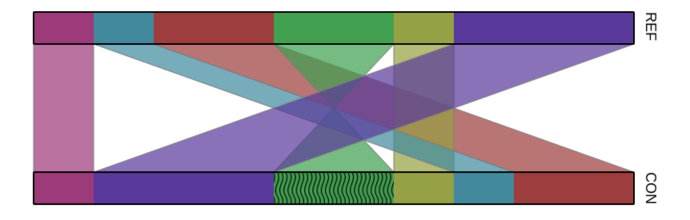


Figure 6.1: gto_comparative_map execution plot.

Input parameters

The gto_max program needs two streams for the computation, namely the input, which are two decimal files.

The attribution is given according to:

An example of such an input files are:

File 1:

```
0.123
3.432
2.341
1.323
7.538
4.122
0.242
0.654
5.633
```

File 2:

```
2.123
5.312
2.355
0.124
1.785
3.521
0.532
7.324
2.312
```

Output

The output of the gto_max program is the numeric file with the maximum value for each row between both input files.

Executing the application with the provided input, the output of this execution is:

```
2.123000
5.312000
2.355000
1.323000
7.538000
4.122000
0.532000
7.324000
5.633000
```

6.15 Program gto_min

The gto_min computes the minium value in each row between two files.

For help type:

```
./gto_min -h
```

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

Input parameters

The gto_min program needs two streams for the computation, namely the input, which are two decimal files.

File 1:

```
0.123
3.432
2.341
1.323
7.538
4.122
0.242
0.654
5.633
```

File 2:

```
2.123
5.312
2.355
0.124
1.785
3.521
0.532
7.324
2.312
```

Output

The output of the gto_min program is the numeric file with the minimum value for each row between both input files.

Executing the application with the provided input, the output of this execution is:

```
0.123000
3.432000
2.341000
0.124000
1.785000
3.521000
0.242000
0.654000
2.312000
```

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