# COMPUTEL VO.4

# **User Manual**

# **Contents**

troductionto	2
evelopment	2
tation	
cense	
equirements	
ownload and installation	
Setting up samtools	2
Running Computel from command line	
gorithm description	3
put and advanced options	
4Q	

#### Introduction

Computel is R-based software for computation of mean telomere length from whole genome next generation sequencing (NGS) data.

# Development

Computel has been developed by the members of the Bioinformatics Group at the Institute of Molecular Biology of the National Academy of Sciences of the Republic of Armenia (IMB NAS RA). You can visit the group's webpage at the following link: http://big.sci.am.

#### Citation

When using Computel in your research, please refer to the GitHub repository at <a href="https://github.com/lilit-nersisyan/computel">https://github.com/lilit-nersisyan/computel</a>.

### License

Copyright (C) 2014 Lilit Nersisyan & Arsen Arakelyan BIG IMB NAS RA.

This program is free software: you can redistribute and/or modify it under the terms of the GNU General Public License version 3. The license can be found at http://www.gnu.org/licenses/gpl.html.

## Requirements

**Computel v0.4** works for **Unix** systems (tested for Linux Ubuntu), refer to Computl v02 for Windows and Mac OS tested versions. It uses the following dependencies, which should be installed on your system: **R 3.0.3** or higher, **Samtools 1.3** or higher.

The following binaries are included in the Computel package, and should work without worries for installation (there is a plan B in case these binaries do not work for your system):

bowtie2-align and bowtie2-build programs from Bowtie2-2.1.0 (not 2.2.0!), http://sourceforge.net/projects/bowtie-bio/files/bowtie2/2.1.0/

Computel has been tested only on provided versions of software. It does not work with Bowtie2-2.2.0, and is not guaranteed to work with higher versions of Samtools and Picard.

#### Download and installation

Computel can be downloaded from GitHub at <a href="https://github.com/lilit-nersisyan/computel">https://github.com/lilit-nersisyan/computel</a>.

Download and extract the package into a local directory. The folder contains the needed scripts, binaries and files for checking proper setup. Do not change the relative location of the folders within the package.

#### **Setting up samtools**

If samtools 1.3 or higher is not installed on your system, download it from <a href="https://github.com/samtools/samtools">https://github.com/samtools/samtools</a>. You may install it by running "make install" or you may run "make" and specify the samtools executable with -sam option of Computel.

#### **Running Computel from command line**

To make computel.sh executable, navigate to the Computel folder and run:

chmod +x computel.sh

The following lines demonstrate the input-output arguments and how to run Computel, the next sections will provide more detailed explanations on arguments and the algorithm.

#### Basic usage:

./computel.sh [options] {-1 <fq1> -2 <fq2> -3 <fq3> -o <o>}

#### Input:

<fq1> fastq file (the first pair or the only fastq file (for single end reads)

<fq2> fastq file (optional: the second pair of fastq files, if exists)

<fq3> fastq file (optional: the third pair of fastq files, if exists)

<o> output directory (optional: the default is computel\_out)

#### Options (advanced):

<-proc> number of processors to be used (default: 4)

<-sam> samtools path (optional: if not supplied, Computel will use the samtools installed on the

system)

<-bowal> bowtie2-align path (optional: the bowtie2-align is located at computel's bin directory by

default.)

<-bowb> bowtie2-build path (optional: the bowtie2-build is located at computel's bin directory by

default.)

<-nchr> number of chromosomes in a haploid set (the default is 23)

<-lgenome> whole genome length (the default is 3244610000)

<-pattern> telomere repeat pattern (the default is 'TTAGGG'; change this if you're using Computel

for a non-human organism)

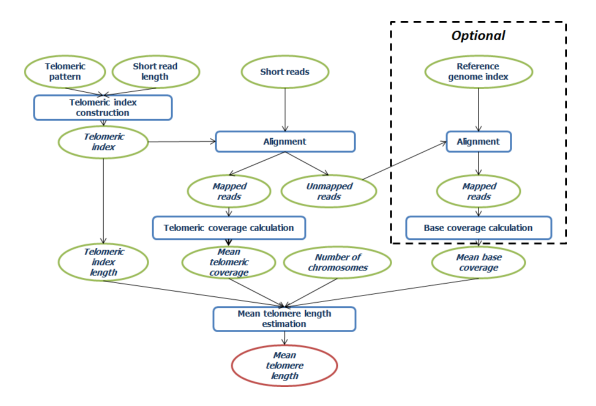
<-minseed> the min seed length (read length minus the number of flanking N's in the telomeric

index; should be in the range [12-read.length]; This is a tested and carefully set

parameter (defualt = 12); Change this only if you REALLY KNOW what you're doing!)

# Algorithm description

The algorithm workflow is presented in the figure. For its detailed description, refer to the main paper [Nersisyan L, Arakelyan A (2015) Computel: Computation of Mean Telomere Length from Whole-Genome Next-Generation Sequencing Data. PLoS ONE 10(4): e0125201. doi:10.1371/journal.pone.0125201].



# Input and advanced options

#### Fastq files:

Computel uses whole genome fastq files as input. If the reads are single end, usually only one fastq file is available (use only -1 <fq1> option and skip the rest). If the reads are paired-end there will be two or three fastq files (the third one contains unpaired reads), which can be supplied with -1 <fq1>, -2 <fq2> and -3 <fq3> options.

The output directory will contain the index and alignment files generated by Computel and the final output file (tel.length.xls). The directory may be specified to replace the default (/computel\_out) with the option -o <outputdir>.

#### **Options (advanced):**

<-proc>: number of processors to be used for parallelization of the alignment process (default: 4)

<-sam>: samtools path. Specifying Samtools directory rather then using the default binary available in the Computel package should only be done if the latter

doesn't work. Please, refer to the "Setting up samtools" section for details.

<-bowal>: bowtie2-align path. This is also optional: the bowtie2-align is located at computel's bin

directory by default. Replace it only if the binary does not work.

<-bowb>: bowtie2-build path. This is also optional: the bowtie2-build is located at computel's bin

directory by default. Replace it only if the binary does not work.

<-nchr>\*: number of chromosomes in a haploid set (the default is 23 for human genomes). Use

this for organisms other than humans or for special cases where the number of chromosomes is not a multiple of 23. Note, you might need to change the genome

length accordingly.

<-lgenome>: whole genome length (the default is 3244610000 for humans)

<-pattern>: telomere repeat pattern (the default is 'TTAGGG'; change this if you're using Computel

for a non-human organism). Note, this is the repeat pattern present at the 3' end (on

the 5'-3' strand).

<-minseed>\*\*: the min seed length (read length minus the number of flanking N's in the telomeric

index; should be in the range [12-read.length]; This is a tested and carefully set

parameter (defualt = 12); Change this only if you REALLY KNOW what you're doing!)

\*Notes on number of haploid chromosomes

Telomere length is calculated with the formula:

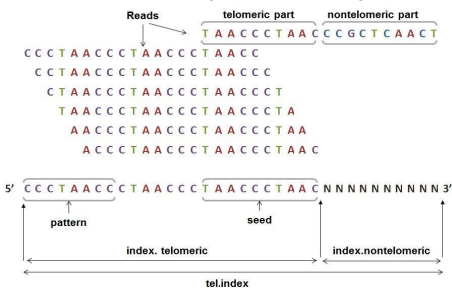
```
(mean(tel.cov/base.cov))*(rl+pl-1) / (2*num.haploid.chr),
```

where *tel.cov* is coverage at telomeric index and *base.cov* is coverage at reference genome; *rl* is read length and *pl* is telomeric pattern length, *num.haploid.chr* is the number of haploid chromosomes.

The division by *num.haploid.chr* is for deriving at a mean value of telomere length for each chromosome end. The number 2 is to account for the two ends of each chromosome. Note that the ploidy of the genome will have no influence on the results, meaning that, adjusted for the rest of the parameters, the results will not differ for haploid, diploid and polyploid genomes.

#### \*\*Min.seed

This is an advanced option and specifies the minimum number of telomeric repeat bases. More specifically, this value is used for building the telomeric index with trailing N bases of length (*read.length - min.seed*), which is the "index.nontelomeric" region of the index in the figure below.



If this option is omitted, the default value of 12 will be used. Increasing **min.seed** will increase specificity but decrease sensitivity of capturing telomeric reads, and vice-versa.

We believe that for the majority of biological cases, the default value is optimal.

## Notes on some configuration options

#### Pattern

Telomeric repeat pattern is specified for the 3'-end of the chromosome. E.g. in case of human telomeres, the sequence at 3'-end it is TTAGGG, while the 5'-end has pattern CCCTAA.

If the pattern is omitted in the configuration file, the default value TTAGGG for human telomere repeats will be used. In case of a different study organism another pattern should be specified.

Number of haploid chromosomes

Telomere length is calculated with the formula:

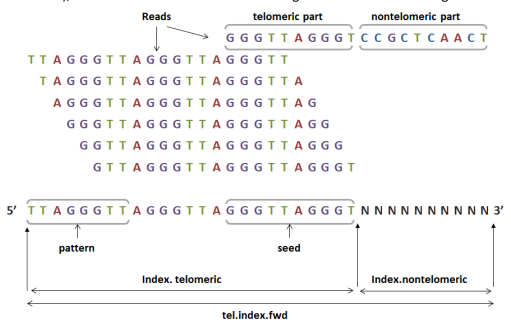
(mean(tel.cov/base.cov))\*(rl+pl-1) / (2\*num.haploid.chr),

where *tel.cov* is coverage at telomeric index and *base.cov* is coverage at reference genome; *rl* is read length and *pl* is telomeric pattern length, *num.haploid.chr* is the number of haploid chromosomes.

The division by *num.haploid.chr* is for deriving at a mean value of telomere length for each chromosome end. The number 2 is to account for the two ends of each chromosome. Note that the ploidy of the genome will have no influence on the results, meaning that, adjusted for the rest of the parameters, the results will not differ for haploid, diploid and polyploid genomes.

#### Min.seed

This is an advanced option and specifies the minimum number of telomeric repeat bases. More specifically, this value is used for building the telomeric index with trailing N bases of length (*read.length - min.seed*), which is the "index.nontelomeric" region of the index in the figure below.



If this option is omitted, the default value of 12 will be used. Increasing **min.seed** will increase specificity but decrease sensitivity of capturing telomeric reads, and vice-versa.

We believe that for the majority of biological cases, the default value is optimal.

### FAQ

The FAQ section will be generated once a question arises.

Feel free to ask questions and make suggestions by mailing to Lilit Nersisyan at <u>Inersisyan@mb.sci.am</u>.