Fundamentals of computational biology

Lecture notes

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Preface

We started this book with the aim of compiling the lectures of the course Fundamentals of Computational Biology offered at Universidad EAFIT for undergrad students in Biology. The course has been taught from different perspectives from its creation, yet the last iteration was divided into three modules. i) introduction to Unix (4 lectures) ii) introduction to sequence analysis and genomics (7 lectures) and iii) principles of structural biology (4 lectures).

Lectures are focused on a theoretical-practical approach were basic concepts from biology, bioinformatics and computer science and interleave with the practice to solve challenges.

Introduction

Here we present a course centered book of the Fundamentals of Computational Biology. We will cover several topics, from using the unix tools, the importance of package manager systems (such as homebrew and conda), sequencing technologies, sequence alignments, molecular phylogenetics, genome assembly and annotation, and variant calling analysis.

Part I

Unix

1 Welcome to the command line

In this chapter we will explore the fundamentals of the command line. That is the concepts of Unix based systems the command line (CLI) and how we can use it to access information programmatically.

Part II Sequence analysis

2 Introduction to sequence analysis

In this chapter we will discuss several about several points of view about bioinformatics and computational biology and how to get started with the command line being a biologist, we will further consider several biological concepts that appear central to understand the manipulation of biological data.

Endless debate: bioinformatics vs. computational biology

Getting started with the command line

The duality of DNA

The central dogma theory of molecular biology extended

Sequencing strategies

Sequencing over time

Some insights from sequencing genomes

3 Sanger analysis

This is a section about the first gen sequencing tech

Databases exploration

Sanger sequencing methods

The chain termination method

Sanger with capillary electrophoresis

Strengths and limitations of Sanger methods

Files from Sanger

Sanger processing workflow

The 16S rRNA and its relevance for sequencing

Part III Challenges demostrations

4 Genome searching

Challenge

Your profesor is interested on knowing how many complete genomes of Bacillus subtilis are there in the NCBI databases. He ask you later to count the number of features (genes, CDS, ncRNA, rRNA, etc.) in the genome of Bacillus subtilis NCIB 3610 (GCF_002055965.1). And tell you to document each of the steps and how did you end up with the answer. Saving the file with your initials (e.g., CG-activity01.{md,txt,docx})

Downloading a genome

```
ncbi-genome-download
```

Downloading from NCBI

The first step in this journey is to download a bunch of sequences programatically. To do so, we will use the program ncbi-genome-download.

You could inspect all the options it provides, now we will set our command as the following:

```
ngd --genera "Bacillus subtilis"\
    -s refseq\
    -l complete\
    -o Data\
    --flat-output\
    --format features\
    -n bacteria\
    | head -n 10
```

```
Considering the following 193 assemblies for download: GCF_000772125.1 Bacillus subtilis ATCC 13952 GCF_000772165.1 Bacillus subtilis ATCC 19217
```

```
GCF_000772205.1 Bacillus subtilis Bs-916

GCF_000782835.1 Bacillus subtilis SG6

GCF_000789295.1 Bacillus subtilis PS832

GCF_000952895.1 Bacillus subtilis BS34A

GCF_000953615.1 Bacillus subtilis BS49

GCF_001015095.1 Bacillus subtilis UD1022

GCF_001037985.1 Bacillus subtilis TO-A JPC
```

Listing files

. . .

```
ls Data | head -n 10
```

Decompressing using gzip

```
gzip -d *
```

Some files in our data dir

```
ls Data | head
```

Importing the files into R

```
library(tidyverse)
library(fs)

all_features <- dir_ls("Data/") %>%
   map_df(read_tsv)

all_features %>%
   head()
```

. . .

```
library(tidyverse)
-- Attaching packages ----- tidyverse 1.3.1 --
v ggplot2 3.3.5
                v purrr
                         0.3.4
v tibble 3.1.6
                v dplyr 1.0.8
       1.2.0
v tidyr
                v stringr 1.4.0
v readr
        2.1.2
                v forcats 0.5.1
-- Conflicts ----- tidyverse conflicts() --
x dplyr::filter() masks stats::filter()
x dplyr::lag()
              masks stats::lag()
library(fs)
all_features <- dir_ls("Data/") %>%
 map_df(read_tsv)
all_features %>%
head()
```

A tibble: 0 x 0

Data processing

```
all_features_grouped <- all_features %>%
  rename(feature = `# feature`) %>%
  select(assembly, feature) %>%
  group_by(assembly, feature) %>% operations
  count() %>%
  pivot_wider(names_from = feature, values_from = n) %>%
  arrange(desc(CDS))

all_features_grouped %>%
  head()
```

create a new dataset that will group by features per accession. get read of the weird name of the column. Select these two columns. Group by these two columns to perform. count the numbers of rows based on the applied group. generate a wide dataset sending row names as columns. Arrange descending by the number of CDSs.

5 Sanger processing

Challenge

Your professor gives you a couple of .ab1 files of a 16S rRNA gene from an old project a student conducted. She tells you to process and analyse them using the sanger sequence pipeline analysis. And as she doesn't know from which species they belong, she ask you to identify the organism to whom it belongs by using the resulting consensus sequence. She finally reminds you to document each step of the process including the identification step.

Procesing a single .ab1 pair

Processing a bulk of .ab1 files

```
library(fs)
library(purrr)
dirs <- fs::dir_ls("~/Projects/Bacillus/Data/Sanger/Inter")</pre>
sanger_bulk <- function(dir) {</pre>
  SangerAlignment(
   ABIF_Directory = dir,
    REGEX_SuffixForward = "_1_F.ab1",
    REGEX_SuffixReverse = "_2_R.ab1"
  )
}
genes <- dirs %>%
  map(sanger_bulk)
launchApp(genes$`/Users/camilogarcia/Projects/Bacillus/Data/Sanger/Inter/gyrA`)
writeFasta(
  outputDir = "~/Documents/Teaching/BiologyCourses/BI0487/Demos/02-demo-sangeranalysis",
  selection = "contigs_unalignment"
```

6 Sequence alignment demo

Challenge

Your professor is working with species from genus Bacillus and want to align an orthologous gene from 10 genomes of different isolates. He gives you the GenBank accession number of these isolates and ask you to select one orthologus gene (Nucleotide seq) that you consider might be useful to differentiate the bacterial isolates and ask you to align those genes as you better consider. He finally ask you to document each step and send him the sequence alignment file in FASTA format along with the sequence alignment general stats in a TXT file (length, number of each nucletides and other stats you consider important).

```
Accessions: GCA_012225885.1, GCA_000196735.1, GCA_000742895.1, GCA_001584335.1, GCA_000007825.1, GCA_000832905.1, GCA_000008425.1, GCA_000507105.1, GCA_000832605.1, GCA_900186955.1
```

Download sequences

Make sure to use the --flat-output avoiding download of multiple metadata

```
ngd --flat-output -p 4 -s genbank -A genome-accessions.txt -F cds-fasta bacteria
```

In this case cds-fasta parameter will download the nucleotide sequences of the gene. Other alternatives could be useful such as blast search on a genome database or searching through the GENBANK annotation files (both files also could be downloaded using ngd).

Unwrapping FASTA records

NCBI registries came with an undesirable wrapping around the lines of sequencing which basically is inserting a return character after some established number of characters. Then a way to get rid of them is to use a command line utility from AstrobioMike (Mike Lee) which will give a line per sequence after the FASTA header. We can later assume the the first line after the header will be the entire sequence

```
for i in GCA_*; do
   N=$(basename $i .fna);
   bash bit-remove-wraps.sh ${i} > ${N}_unwrapped.fasta;
done
```

Gene search

A possible way to search throughout the file registries is by using the grep command, that recursively will search each file. Fine tuned it allow to search for the first match, but also for the "after-context" in terms of lines desired to be printed:

```
grep -h\
    -m 1\
    -A 1\
    -E "DNA gyrase, A| gyrase subunit A | gyrase alpha| gyrase \(subunit A\)| gyrA" *.fast
    sed "s/--//g" |\
    sed "/^$/d"
```

After finding the genes we could exclude some lines using sed avoid the "–" characters and the empty blank line using the appropriate regular expression (^\$/d). We are now with an almost clean multi sequence file, because header names are still and will be problematic. How do we programatically change the FASTA headers? We will see in the next step.

Renaming fasta headers

A simple but powerful script to do this is bit-dedup-fasta-heades it was developed by AstrobioMike (Mike Lee) and it simply parses the headers and substitutes by a simple encoder found en each of them:

```
python bit-dedupe-fasta-headers.py -i all_gyrA.fasta -o all_gyrA_renamed.fasta
```

Now the files has files names that are simply to work with. Which will enable to asses better out sequence alignment matrix.

Sequence alignment

There are many programs that are suited for performed multiple sequence alignments. Perhaps the two most used are MAFFT and MUSCLE both specialized in multiple sequence alignment (that is: when having two or more than two sequences). The second tends to be more accurate when having large data-sets, but the first on is more versatile, fast and accurate on different kind of data-sets.

Both programs take as input a single file containing all the sequences concatenated horizontally (that is a multi-fasta file) careless of the extension but (MFA, FA, FASTA, FNA, etc). And generate a simple output (whether with the -o in MUSCLE or to the std output in MAFFT)

```
ginsi --preservecase --reorder all_gyrA_renamed.fasta > all_gyrA_renamed_ginsi.fasta # globaleinsi --preservecase --reorder all_gyrA_renamed.fasta > all_gyrA_renamed_einsi.fasta # generinsi --preservecase --reorder all_gyrA_renamed.fasta > all_gyrA_renamed_linsi.fasta # localemuscle -i all_gyrA_renamed.fasta -o all_gyrA_renamed_muscle.fasta

famsa -t 8 all_gyrA_renamed.fasta > all_gyrA_renamed_famsa.fasta

kalign -i all_gyrA_renamed.fasta -o all_gyrA_renamed_kalign.fasta
```

Assesment of the alignment

Inspection of the alignment is there very first step for assesing its quality. A CDS tends to genereate a codon-like alignment starting with the methione codon (ATG,GTG) and finishing with a stop (TAA, TAG, etc.). Therefore finding this structure when aligning a complete genes is expected. If a middle fraction of the gene is being aligned ORF might not display any stop codon. Verifying a codon-like alignment shows a biological order on the sequences other that mere artifact of the alignment, that is an evolutionary behavior of the sequence. We can do it usin seqfu from the CLI or interactively with AliView.

A second step is to find the variability of the alignment. A simple way to find that is to calculate simpl stats from the alignment (sites, variable sites, As, Ts, etc.). A powerful cli program to do so is goalign

```
goalign stats -i all_gyrA_renamed_linsi.fasta
```

An alternative approach using BLAST

```
ngd --flat-output -p 4 -s genbank -A genome-accessions.txt -F fasta --parallel 8 bacteria

for i in GCA_*; do cat ${i} >> all_genomes.fasta; done

makeblastdb -in all_genomes.fasta -parse_seqids -blastdb_version 5 -title "demo" -dbtype nucl
blastn -db all_genomes.fasta -query gyrA.fasta -outfmt "6 sseqid sseq" -word_size 5 -evalue
```

The alternative using the GCF

```
for i in *fna; do; goalign subset -e "gyrA" -i ${i} --unaligned;done | grep ">"
```

7 Phylogenetic reconstruction

Challenge

Your professor has been working with some mammal species and want to know the relationships of some sampled individuals. To do so he extracted the DNA and amplified the mitochondrial CYTB gene of those individuals. He gives a folder with multiple sequences and ask you to align them and to reconstruct two trees one using maximum likelihood (ML) and other using a Bayesian inference (BI). Then ask you to explain if both trees are congruent with each other.

Sequence alignment cytb

```
linsi --preservecase --reorder cytB.fasta > cytB-aligned.fasta
```

Evolutionary substitution model

```
modeltest-ng -i cytB-aligned.fasta -d nt -o model-cytB.txt

04-demo-phylogenetics/model-cytB.txt.log
```

Maximum likelihood reconstruction

```
raxml-ng --msa cytB-aligned.fasta --model GTR+I+G4
```

Tree building

```
raxml_data <- read.tree("cytB-aligned.fasta.raxml.support")

raxml_data$tip.label <- str_replace_all(raxml_data$tip.label, "_", " ")

(
    raxml_tree <- ggtree(raxml_data) +
        geom_tiplab() +
        # geom_point2(aes(subset = !isTip, fill = as.integer(label)), shape = 24, size = 3) +
        geom_text(aes(label = as.integer(label))) +
        theme_tree(legend.position = c(0.8, 0.7))
)</pre>
```

Bayes inference reconstruction

Tree building

```
tail -n 18 ../Data/cytB-mb.nex

mb -i ../../Data

sumtrees.py -s mcct -o=cytB-mb-mcct.tre cytB-aligned.fasta-out.nex.run1.t
```

8 Sequence reads assesment

Challenge

Your professor challenges you to assembly a bacterial genome. He wants to know if an assembly using Illumina reads or Nanopore reads is better. The raw sequences for Illumina could be downloaded with the code SRR15634574. And the Nanopore raw reads with SRR15634573 here: https://sra-explorer.info. Choose at least on set and follow the instruction from the lecture to assemble the genome. Document each step and send to the professor the assembly-{your-initials}.fasta. You could form groups of max. 4 students.

General stats from fastq files

A simple but fast cli to display the general stats from fastq is seqfu

```
seqfu stats -n *.gz
```

A graphical assessment of reads

Often its important to assess reads graphically and nanoplot offers a complete graphical summary (and general stats as well) of the reads

```
nanoplot --fastq reads.fastq.gz --output reads-report
```

9 Genome assembly

9.1 Challenge

Your professor has sequenced a bacterial isolate using PacBio and Nanopore sequencing methods and has got the FASTQ files from both technologies. Now he needs to know the quality and quantity of these data before start any other analysis and ask you to assess the data. He needs to know how many sequences there are, how many base pairs (in GB) are there and the N50. He is also interested in see a visualization of the i) number of bases vs. sequence lengths (log transformed) and ii) the read length vs. read quality vs. read number.

He ask you to document every step and to conclude what data should be used.

Download the reads

wget

Assess read qualities

When using illumina fastqc is a very fast alternative. For nanopore nanplot will do the job.

fastqc

Exploring assemblers

One of the most popular genome assemblers for NGS is spades whereas for TGS flye has been widely used

Shovill: spades under the hood

shovill is a pipeline that enables pre and post processing of genomic data. It can be tunned to several tools for the processing steps and also to select different standalone assemblers

```
shovill --outdir MxanthusIllumina\
    --R1 R1.fq.gz\
    --R2 R2.fq.gz\
    --trim\
    --cpus 32
```

Dragonflye: flye under the hood

Similar to shovill (and inspired by it) dragonflye is a pipeline that enables several processing steps of genomic data be

```
dragonflye --outdir MxanthusNanopore\
    --gsize 9Mb\
    --trim\
    --reads ont-readsfastq.gz\
    --racon 5
```

```
wget
```

Since we are trying to assemble a bacterial genome, computer memory appears to be a limiting features of a local machine. Then, a computer cluster with high performance turns out to be an important need.

First we need environment installations, therefore its important to have conda environments with the assemblers and other programs (conda create -c bioconda dragonflye dragonflye and conda create -c bioconda shovill shovill). That way both assemblers pipelines will lie in separate environment avoiding possible dependenies problems

We will use Apolo computer cluster which uses Slurm as the computer system workload manager (i.e a program that manages the time and resources of the computer).

References