VitisOmics

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August 24, 2015

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1 Overview

This document describes the "VitisOmics" project. This project aims at handling "omics" data in the genus Vitis (e.g. grapevine) in an open and reproducible way. Nevertheless, it requires some basic knowledge and skills about bioinformatics on GNU/Linux computers.

Such "omics" data are already available from various places, such as URGI, NCBI, EBI, etc; and several committee from the IGGP (International Grape Genome Program) strive at improving interoperability. But my attempt, via the usage of git and GitHub, could prove for the community to be a useful addition to these efforts.

The project directory is organized as advised by Noble (PLoS Computational Biology 2009). On any Unix-like system, it is easily done with the following commands:

```
mkdir -p VitisOmics; cd VitisOmics/
touch AUTHORS COPYING README; mkdir -p doc data src results
```

On any Unix-like system, the project directory can also be easily compressed and transferred (ignoring large data files):

```
cd ..; tar -czvf VitisOmics.tar.gz \
--exclude=VitisOmics/data --exclude=VitisOmics/results \
--exclude="*~" --exclude=".*" VitisOmics
```

In order to concretely promote collaborative editing in a distributed manner, the content of the "VitisOmics" repository should be based on "plain text" files. As a consequence, this document is written in the org format, and can thus be automatically exported into the pdf format, best by emacs, but also by pandoc.

1.1 Contributors

The person roles comply with R's guidelines (The R Journal Vol. 4/1, June 2012).

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2 Data

```
mkdir -p data; cd data/
```

TODO: retrieve genome data from other cultivars than PN40024, e.g. Sultanina

2.1 URGI

• https://urgi.versailles.inra.fr/Species/Vitis/Data-Sequences/Genome-sequences

```
mkdir -p urgi; cd urgi/
../../src/download_urgi.bash
```

When needed, the script decompresses zip files and compress them again but with gzip instead.

2.2 NCBI

- http://www.ncbi.nlm.nih.gov/genome/401
- ftp://ftp.ncbi.nlm.nih.gov/genomes/Vitis_vinifera/

```
mkdir -p ncbi; cd ncbi/
../../src/download_ncbi.bash
```

Note the important file scaffold_names which provides the correspondence between original scaffold names (i.e. from the sequencing center) and various NCBI identifiers (RefSeq, GenBank, etc).

2.3 EBI

12X.0 as well as soft-masking by RepeatMasker

```
mkdir -p ebi; cd ebi/
../../src/download_ebi.bash
```

3 Results

```
mkdir -p results; cd results/
```

TODO: compress fasta files with bgzip instead of gzip

On a computer cluster, indexed files could be copied for usage by everyone, e.g. in /Genomics/Vitis if on the CIRAD cluster of the SouthGreen platform.

We need to keep the information about the data source (e.g. URGI or NCBI) because there can be differences in terms of N spacers and additional scaffolds (from Aegilops?) at the NCBI.

3.1 URGI

```
mkdir -p urgi; cd urgi/
```

3.1.1 Reformat sequence headers for VITVI_PN40024_12x_v0_chroms_URGI

Launch script:

Check:

```
zcat VV_chr12x.fsa.gz | wc -1 # 8240706
zcat VV_chr12x.fsa.gz | grep -c ">" # 33
zcat VITVI_PN40024_12x_v0_chroms_URGI.fa.gz | wc -1 # 8240706
zcat VITVI_PN40024_12x_v0_chroms_URGI.fa.gz | grep -c ">" # 33
diff <(zcat VV_chr12x.fsa.gz) <(zcat VITVI_PN40024_12x_v0_chroms_URGI.fa.gz)</pre>
```

Only the headers differ, not the sequences, so everything is fine.

Basic stats:

```
zcat VITVI_PN40024_12x_v0_chroms_URGI.fa.gz | md5sum # eff315994fafe35333462b9595e10ce5
```

3.1.2 Reformat sequence headers for VITVI_PN40024_12x_v0_scaffolds_EMBL_r102

Launch script:

Check:

```
zcat VV_12X_embl_102_Scaffolds.fsa.gz | wc -1 # 8091565
zcat VV_12X_embl_102_Scaffolds.fsa.gz | grep -c ">" # 2059
zcat VITVI_PN40024_12x_v0_scaffolds_EMBL_r102.fa.gz | wc -1 # 8091565
```

```
zcat VITVI_PN40024_12x_v0_scaffolds_EMBL_r102.fa.gz | grep -c ">" # 2059
diff <(zcat VV_12X_embl_102_Scaffolds.fsa.gz) <(zcat
    VITVI_PN40024_12x_v0_scaffolds_EMBL_r102.fa.gz)</pre>
```

Only the headers differ, not the sequences, so everything is fine.

Basic stats:

```
zcat VITVI_PN40024_12x_v0_scaffolds_EMBL_r102.fa.gz | md5sum # 4
fa2432d7a66c019c7cb41ee4d0cb7bc
```

3.1.3 Reformat sequence headers for VITVI_PN40024_12x_v0_contigs_EMBL_r102

TODO

3.1.4 Reformat sequence headers for VITVI_PN40024_12x_v2_chroms_URGI

Launch script:

Check:

```
zcat 12Xv2_grapevine_genome_assembly.fa.gz | wc -l # 8103449
zcat 12Xv2_grapevine_genome_assembly.fa.gz | grep -c ">" # 20
zcat VITVI_PN40024_12x_v2_chroms_URGI.fa.gz | wc -l # 8103449
zcat VITVI_PN40024_12x_v2_chroms_URGI.fa.gz | grep -c ">" # 20
diff <(zcat 12Xv2_grapevine_genome_assembly.fa.gz) <(zcat
    VITVI_PN40024_12x_v2_chroms_URGI.fa.gz)</pre>
```

Only the headers differ, not the sequences, so everything is fine.

Basic stats:

```
zcat VITVI_PN40024_12x_v2_chroms_URGI.fa.gz | md5sum # 4
e487c28eaf19ef59b0b6128b73935af
```

Length of each sequence:

header	length (bp)
chr1 Vitis vinifera PN40024 assembly12x.2	24233538
chr2 Vitis vinifera PN40024 assembly12x.2	18891843
chr3 Vitis vinifera PN40024 assembly12x.2	20695524
chr4 Vitis vinifera PN40024 assembly12x.2	24711646
chr5 Vitis vinifera PN40024 assembly12x.2	25650743
chr6 Vitis vinifera PN40024 assembly12x.2	22645733
chr7 Vitis vinifera PN40024 assembly12x.2	27355740
chr8 Vitis vinifera PN40024 assembly12x.2	22550362
chr9 Vitis vinifera PN40024 assembly12x.2	23006712
chr10 Vitis vinifera PN40024 assembly12x.2	23503040
chr11 Vitis vinifera PN40024 assembly12x.2	20118820
chr12 Vitis vinifera PN40024 assembly12x.2	24269032
chr13 Vitis vinifera PN40024 assembly12x.2	29075116
chr14 Vitis vinifera PN40024 assembly12x.2	30274277
chr15 Vitis vinifera PN40024 assembly12x.2	20304914
chr16 Vitis vinifera PN40024 assembly12x.2	23572818
chr17 Vitis vinifera PN40024 assembly12x.2	18691847
chr18 Vitis vinifera PN40024 assembly12x.2	34568450
chr19 Vitis vinifera PN40024 assembly12x.2	24695667
chrUkn Vitis vinifera PN40024 assembly12x.2	27389308
total	486205130

$3.1.5 \quad Format \ \mathtt{VITVI_PN40024_12x_v0_chroms_URGI} \ for \ BLASTn$

TODO: change Vvin to VITVI

```
../../src/format_Vvin-PN40024-12x-chr_blastn.bash
```

3.1.6 Index $VITVI_PN40024_12x_v0_chroms_URGI$ for BWA

Launch:

3.1.7 Index VITVI_PN40024_12x_v2_chroms_URGI for BWA

Launch:

3.1.8 Prepare VITVI_PN40024_12x_v2_chroms_URGI for SAMtools and Picard

Make an index as well as a SAM header.

Launch:

3.1.9 Index VITVI_PN40024_12x_v0_chroms_URGI for Bowtie2

Launch:

3.1.10 Index VITVI_PN40024_12x_v2_chroms_URGI for Bowtie2

Launch:

3.1.11 Index VITVI_PN40024_12x_v2_chroms_URGI for Bowtie2 compatible with Tassel

Tassel requires numbers as chromosome identifiers.

Launch:

3.1.12 Translate CRIBI annotations from 12X.0 to 12X.2

Requirement: use or write a script taking as input the 12X.0 GFF3 file as well as the 12.0-12.2 AGP file, and returns as output the 12X.2 GFF3 file

The URGI provides the following AGP file: golden_path_V2_111113_allChr.csv. Unfortunately, after looking at the official specification of the AGP format, the URGI file doesn't seem to be valid, neither for version 1.1, nor 2.2.

TODO: contact URGI

3.2 R/Bioconductor

- http://www.bioconductor.org/
- Huber, W. et al. Orchestrating high-throughput genomic analysis with bioconductor. Nature Methods 12, 115-121 (2015). URL http://dx.doi.org/10.1038/nmeth.3252.

TODO: see AnnotationHub

3.2.1 BSgenome IGGP12Xv2 package

http://bioconductor.org/packages/release/bioc/html/BSgenome.html

Retrieve the sequence data from URGI:

```
cd results/
mkdir -p make_BSgenome_IGGP12Xv2
cd make_BSgenome_IGGP12Xv2/
ln -s ../../data/urgi/12Xv2_grapevine_genome_assembly.fa.gz .
```

Split into one chromosome per file (in the headers, discard everything after the first space):

```
zcat 12Xv2_grapevine_genome_assembly.fa.gz | awk 'BEGIN{RS=">"} {if(NF==0)next;
    split($0,a,"\n"); split(a[1],b," "); print b[1]; print ">"b[1] > b[1]".fa"; for(
    i=2;i<length(a);++i){print a[i] >> b[1]".fa"}}'
gzip chr*.fa
```

Prepare the seed file (IGGP12Xv2_seed.txt) by hand as indicated in the vignette as well as in the official R manual "Writing R extensions". Following this article, I chose the CC0 license (present in the R list of licenses in share/licenses/license.db).

Forge the target package from the seed file:

```
echo "date; echo \"library(BSgenome); forgeBSgenomeDataPkg(\\\"IGGP12Xv2_seed.txt\\\
")\" | R --vanilla; date" | qsub -cwd -j y -V -N forge_BSgenome -q normal.q
```

Build the package and check it:

```
echo "date; R CMD build BSgenome.Vvinifera.URGI.IGGP12Xv2; date" | qsub -cwd -j y -V
     -N build_BSgenome -q normal.q
echo "date; R CMD check BSgenome.Vvinifera.URGI.IGGP12Xv2_0.1.tar.gz; date" | qsub -
     cwd -j y -V -N check_BSgenome -q normal.q
```

The target package is now ready to be installed:

```
R CMD INSTALL BSgenome. Vvinifera. URGI. IGGP12Xv2_0.1.tar.gz
```

TODO: contact A.-F. Adam-Blondon (part of IGGP) and other colleagues from INRA to get their feedback

3.2.2 BSgenome IGGP12Xv0 package

http://bioconductor.org/packages/release/bioc/html/BSgenome.html

Retrieve the sequence data from NCBI (neglect unlocalized and unplaced sequences), and change headers into "chr"<id>:

```
cd results/
mkdir -p make_BSgenome_IGGP12Xv0
cd make_BSgenome_IGGP12Xv0/
ls ../../data/ncbi/Assembled_chromosomes/vvi_ref_12X_chr*.fa.gz | while read f; do
    name=$(basename $f | sed 's/.fa.gz//' | awk '{split($0,a,"_"); print a[4]}');
    echo ${name}; echo ">${name}" > ${name}.fa; zcat $f | grep -v ">" >> ${name}.fa;
    gzip ${name}.fa; done
```

TODO: Prepare the seed file (IGGP12Xv0_seed.txt) by editing the one used for IGGP12Xv2.

3.2.3 TxDb CRIBI12Xv0 package

 $\label{lem:http://www.bioconductor.org/packages/release/bioc/html/GenomicFeatures.html} \\ TODO$