IFB cloud:

Integration of snakemake workflows in an appliance designed for ChIP-seq analysis

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French Institute of Bioinformatics (IFB)



- National service infrastructure in bioinformatics
- 32 platforms grouped into regional centers
- French node of the European research infrastructure ELIXIR



 Provides services and infrastructures in bioinformatics for scientists and engineers working in the field of life sciences



http://www.france-bioinformatique.fr

IFB cloud

- 200 computing cores
- 2 To of RAM
- 50 To of storage

By 2016

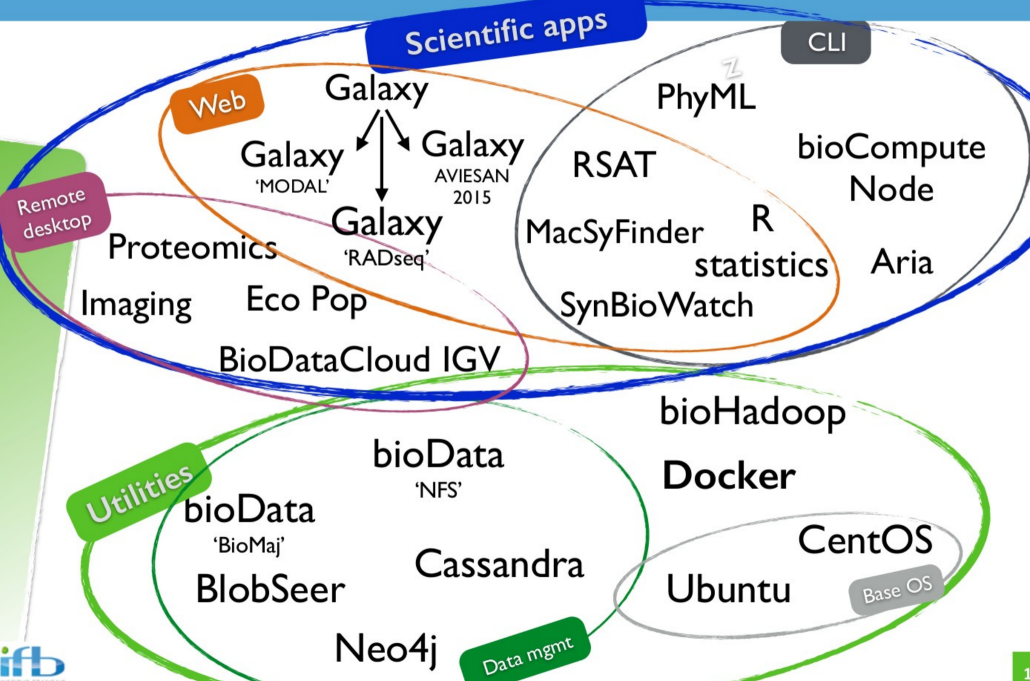
- 3,000 cores
- 500 To of storage

By 2017

- 10,000 cores
- 2 Po of storage



IFB's bioinformatics appliances



A lot of bioinformatics tools



ABYSS 1.3.4
ARIA 2.3
Bioconductor 2.11
biomaj
BLAST+ 2.2.27
Blat 35
Bowtie 0.12.8
Bowtie2 2.0.0beta7

BWA 0.6.2

BWA 0.7.10 CAP3 **CD-HIT 4.6.1** Clustal Omega 1.0.3 **CLUSTALW 2.1** Cufflinks 2.0.2 Cutadapt 1.2.1 **E-SURGE 1.9.0** Exonerate 2.2.0 eXpress 1.5.1 FastA 3.6 FastQC 0.10.1 Galaxy portal **GATK 2.3.4** HMMer 3.0 Imagel 1.48 khmer I.I. **M-SURGE 1.8.5 MEME 4.7** MMSEQ 0.11.2a

Mobyle MODAL MultAlin 5.4.1 **MUSCLE 3.8.31** neo4i Oases 0.2.08 **OMSSA 2.1.9 PeptideShaker** 0.18.3 phyml 3.1 PREDATOR 2.1.2 proline python 2.7 R 2.13 R 3.1.1 R 3.1.2 R-studio Ray 1.3 **RSAT** samtools 0.1.18

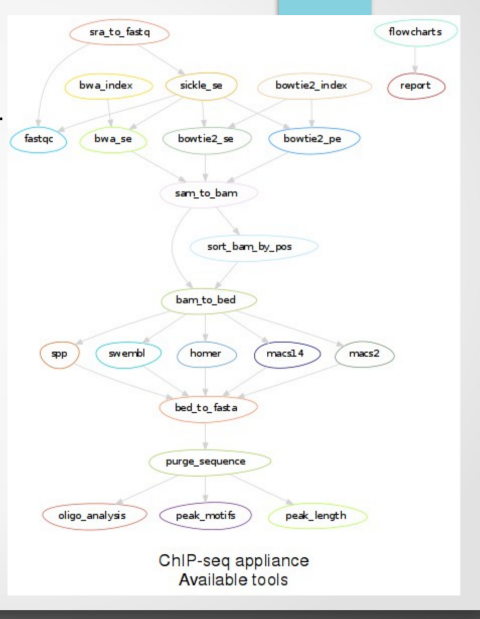
Samtools I.I.

SearchGUI 1.10.4 SegClean Shiny Stacks **STAR 2.4.0fl** SuMo vI **TGICL** TopHat 2.0.6 trim_galore 0.3.7 Trinity 2.0.4 **U-CARE 2.3.2** VCFtools 0.1.11 Velvet 1.2.10 X!tandem 12-10-01-1 XPLOR-NIH 2.30



ChIP-seq appliance

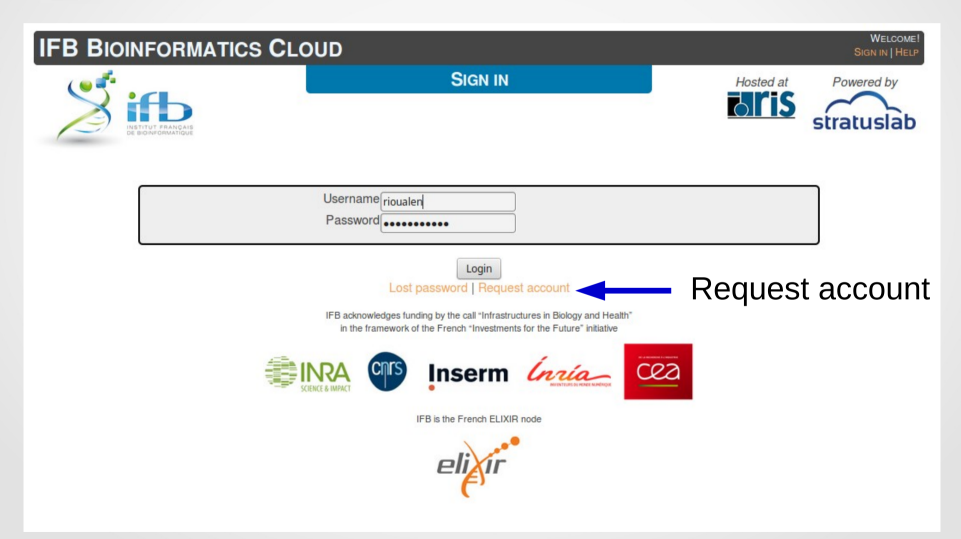
- Base Ubuntu 14.04
 Including ssh, rsync, git, python 2, python 3...
- Programming tools
 snakemake, graphviz, pandas library...
- NGS tools
- fg-chip-seq git repository
 Including workflow examples (py + json)



Using the IFB cloud: request account



cloud.france-bioinformatique.fr



Using the IFB cloud: dashboard



m1 medium

cloud.france-bioinformatique.fr

Add ssh key YOU ARE SIGNED IN AS RIOUALEN IFB BIOINFORMATICS CLOUD News | Dashboard | Monitor | Settings | Help | Sign out **DASHBOARD** Hosted at Powered by



Create a virtual disk



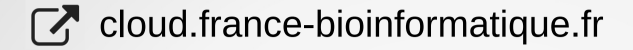
CPU

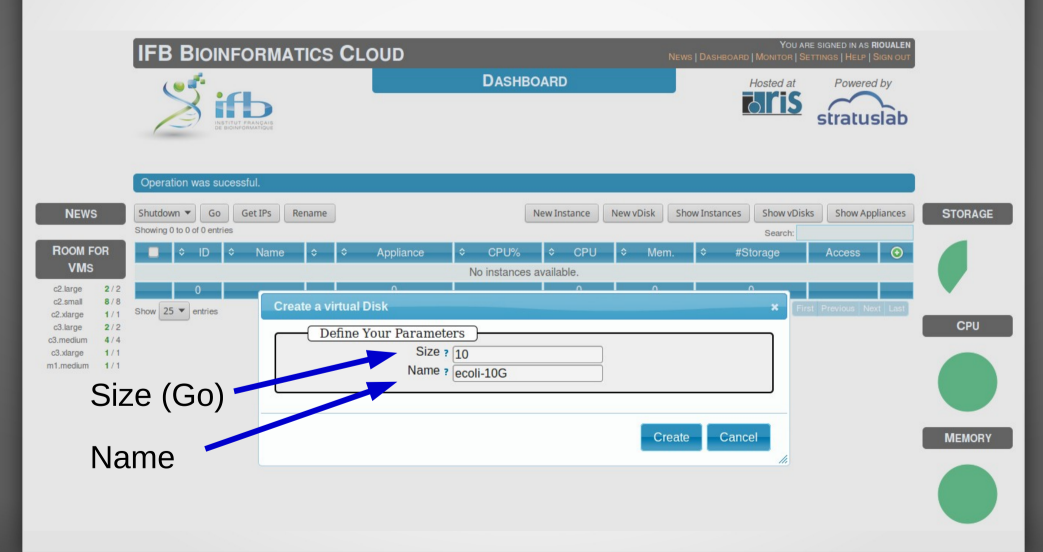
free (100,00%)

STORAGE

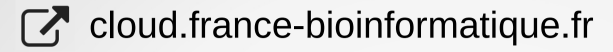


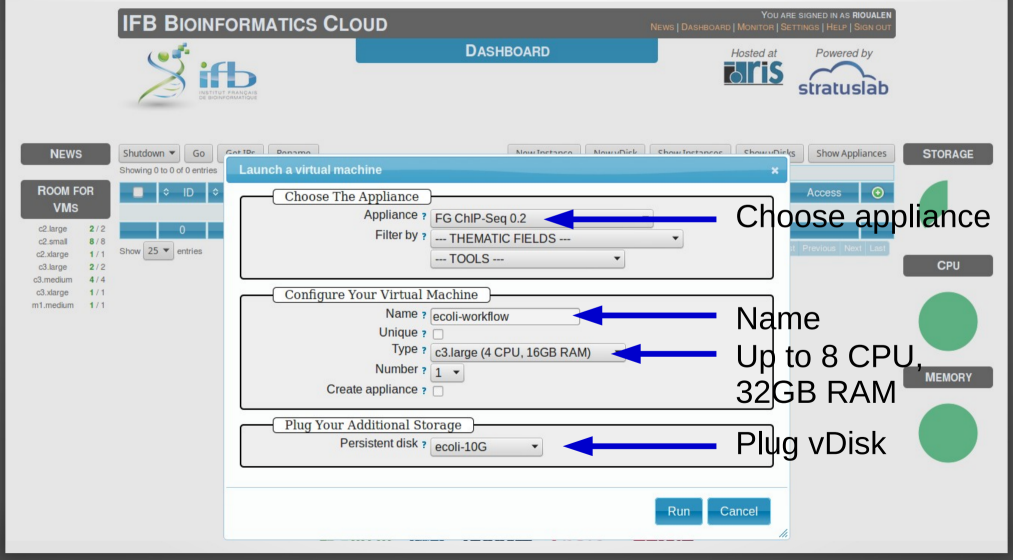
Using the IFB cloud: vDisk



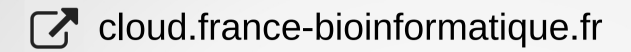


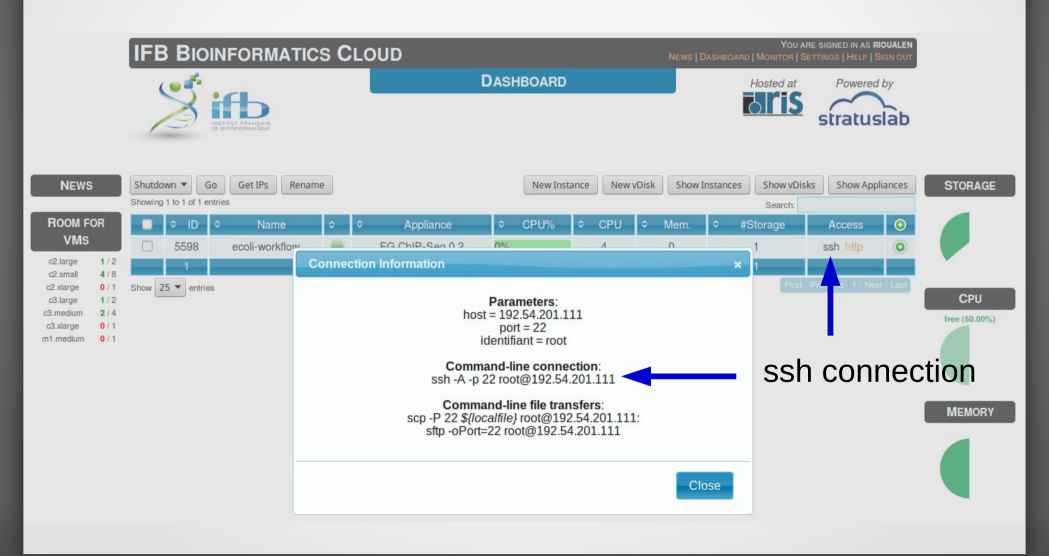
Using the IFB cloud: create instance





Using the IFB cloud: ssh connection





Using the IFB cloud: data

```
root@vm0080:~# ls -R /root/mydisk
/root/mydisk:
data genomes results
/root/mydisk/data:
design.tab GSM000001 GSM000002
                                  samples.tab
/root/mydisk/data/GSM000001:
SRR12345.fastq
/root/mydisk/data/GSM000002:
SRR12345.fastq
/root/mydisk/genomes:
ecoli
/root/mydisk/genomes/ecoli:
ecoli K12.fasta
/root/mydisk/results:
root@vm0080:~#
```

Using the IFB cloud: execute workflow

- Several example workflows already available & adaptable in the git repository
- 1 config file (json) + 1 workflow file (py)
 - Paths to data, genome & results directories
 - Parameters
- Results are saved in the vDisk

```
raw data
bwa index
                sickle se
                       fastac
        bwa se
      sam to bam
       bam to bed
                                flowcharts
```

```
root@vm0080:~# su rg
rg@vm0080:/root$ cd ~/workspace/fg-chip-seq/
rg@vm0080:~/workspace/fg-chip-seq$ ls
doc README.md scripts
rg@vm0080:~/workspace/fg-chip-seq$ snakemake -s scripts/snakefiles/workflows/ecoli.py
```

Thank you!