



IFB cloud:

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

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2015/10/28

# 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

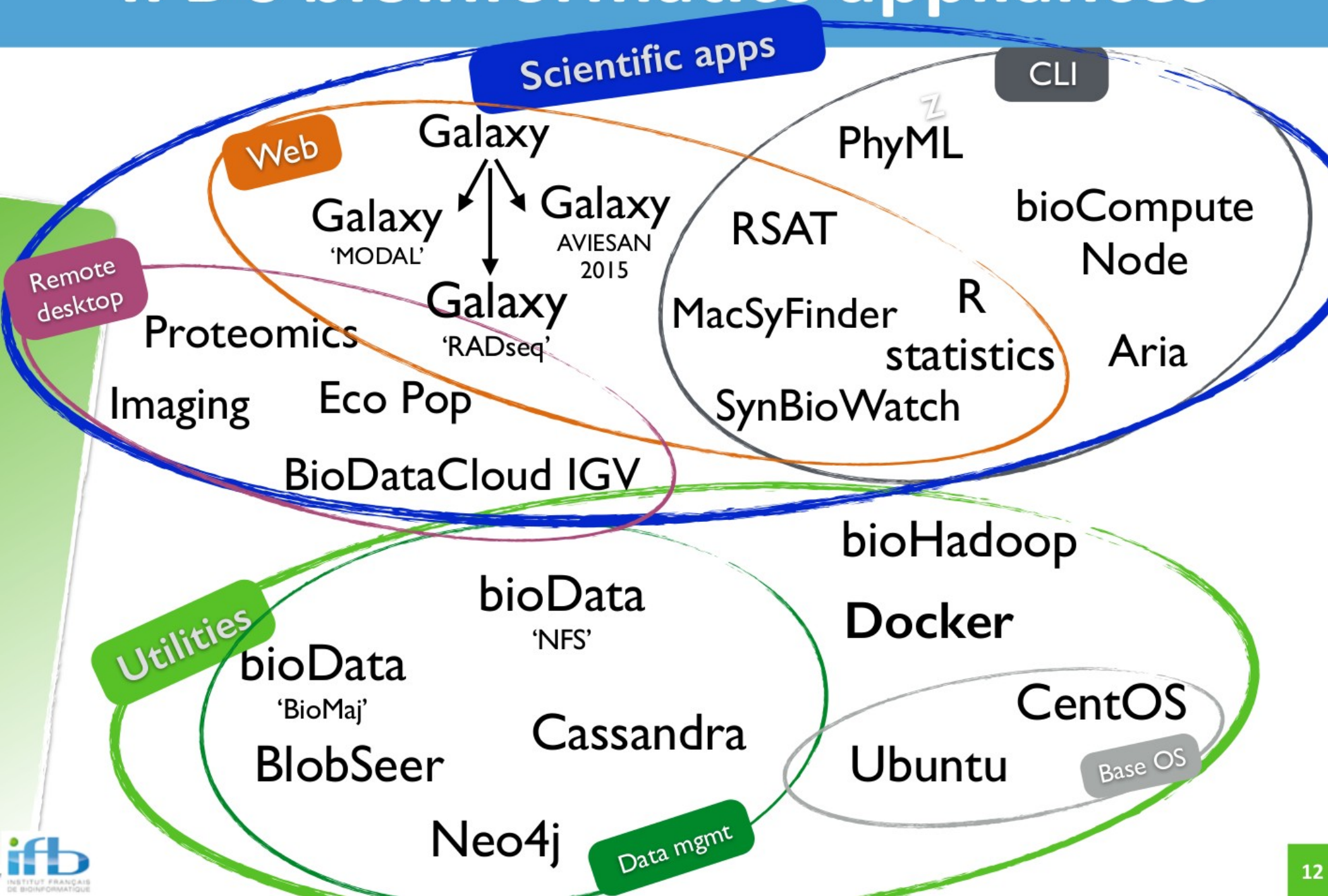
- 3,000 cores
- 500 To of storage

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 2.0.0-beta7  
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  
ImageJ 1.48  
khmer 1.1  
M-SURGE 1.8.5  
MEME 4.7  
MMSEQ 0.11.2a

Mobyle  
MODAL  
MultAlin 5.4.1  
MUSCLE 3.8.31  
neo4j  
Oases 0.2.08  
OMSSA 2.1.9  
PeptideShaker 0.18.3  
phym1 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 1.1


SearchGUI 1.10.4  
SeqClean  
Shiny  
Stacks  
STAR 2.4.0f1  
SuMo v1  
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

IFB BIOINFORMATICS CLOUD

WELCOME!  
SIGN IN | HELP



SIGN IN

Hosted at  Powered by 






Username

Password


Login

[Lost password](#) | [Request account](#)

IFB acknowledges funding by the call "Infrastructures in Biology and Health"  
in the framework of the French "Investments for the Future" initiative




IFB is the French ELIXIR node




Request account

# Using the IFB cloud: dashboard

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Add ssh key



**IFB BIOINFORMATICS CLOUD**

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NEWS | DASHBOARD | MONITOR | SETTINGS | HELP | SIGN OUT

**DASHBOARD**

Hosted at **oris** Powered by **stratuslab**

**NEWS**

Shutdown Go Get IPs Rename

**ROOM FOR VMS**

- c2.large 2 / 2
- c2.small 8 / 8
- c2.xlarge 1 / 1
- c3.large 2 / 2
- c3.medium 4 / 4
- c3.xlarge 1 / 1
- m1.medium 1 / 1

Showing 0 to 0 of 0 entries

New Instance New vDisk Show Instances Show vDisks Show Appliances

| ID                      | Name | Appliance | CPU% | CPU | Mem. | #Storage | Access |
|-------------------------|------|-----------|------|-----|------|----------|--------|
| No instances available. |      |           |      |     |      |          |        |
| 0                       |      | 0         | 0    | 0   | 0    | 0        |        |

Show 25 entries

First Previous Next Last

**STORAGE**

**CPU**


free (100.00%)

**MEMORY**




Create a virtual disk



# Using the IFB cloud: vDisk

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NEWS | **DASHBOARD** | MONITOR | SETTINGS | HELP | SIGN OUT

 **DASHBOARD**  

Operation was successful.

**NEWS** Shutdown Go Get IPs Rename New Instance New vDisk Show Instances Show vDisks Show Appliances **STORAGE**

Showing 0 to 0 of 0 entries

|                         | ID | Name | Appliance | CPU% | CPU | Mem. | #Storage | Access |
|-------------------------|----|------|-----------|------|-----|------|----------|--------|
| No instances available. |    |      |           |      |     |      |          |        |

Show 25 entries

**ROOM FOR VMS**

- c2.large 2 / 2
- c2.small 8 / 8
- c2.xlarge 1 / 1
- c3.large 2 / 2
- c3.medium 4 / 4
- c3.xlarge 1 / 1
- m1.medium 1 / 1

**Create a virtual Disk**

Define Your Parameters

Size ? 10

Name ? ecoli-10G

Create Cancel


Size (Go)

Name

**CPU**



**MEMORY**

# Using the IFB cloud: create instance

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**DASHBOARD**

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**NEWS** **ROOM FOR VMS**

Showing 0 to 0 of 0 entries

| ID              | 0 |
|-----------------|---|
| Show 25 entries |   |

c2.large 2 / 2  
c2.small 8 / 8  
c2.xlarge 1 / 1  
c3.large 2 / 2  
c3.medium 4 / 4  
c3.xlarge 1 / 1  
m1.medium 1 / 1

**Launch a virtual machine**

**Choose The Appliance**

Appliance ? **FG ChIP-Seq 0.2** **Choose appliance**  
Filter by ? --- THEMATIC FIELDS ---  
--- TOOLS ---

**Configure Your Virtual Machine**

Name ? **ecoli-workflow** **Name**  
Unique ? ☐  
Type ? **c3.large (4 CPU, 16GB RAM)** **Up to 8 CPU, 32GB RAM**  
Number ? 1  
Create appliance ? ☐


**Plug Your Additional Storage**

Persistent disk ? **ecoli-10G** **Plug vDisk**

**Run** **Cancel**



**STORAGE** **CPU** **MEMORY**

# Using the IFB cloud: ssh connection

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**DASHBOARD**

Hosted at  Powered by 

**NEWS** Shutdown Go Get IPs Rename

Showing 1 to 1 of 1 entries

**ROOM FOR VMS**

- c2.large 1 / 2
- c2.small 4 / 8
- c2.xlarge 0 / 1
- c3.large 1 / 2
- c3.medium 2 / 4
- c3.xlarge 0 / 1
- m1.medium 0 / 1

| ID   | Name           | Appliance       | CPU% | CPU | Mem. | #Storage | Access   |
|------|----------------|-----------------|------|-----|------|----------|----------|
| 5598 | ecoli-workflow | EG-ChIP-Seq-0.2 | 0%   | 4   | 0    | 1        | ssh http |

Search:

First | Prev | 1 | Next | Last

**STORAGE**

**CPU** free (50.00%)

**MEMORY**

**Connection Information**

**Parameters:**  
host = 192.54.201.111  
port = 22  
identifiant = root

**Command-line connection:**  
ssh -A -p 22 root@192.54.201.111

**Command-line file transfers:**  
scp -P 22 \${localfile} root@192.54.201.111:  
sftp -oPort=22 root@192.54.201.111

Close

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

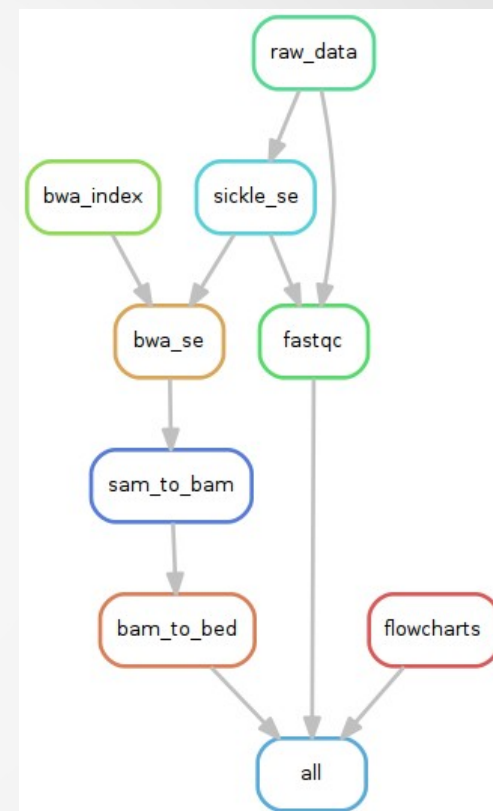
/root/mydisk/genomes/ecoli:
ecoli_K12.fasta

/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



```
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!