



Session de formation 2023



bioinformatics platform dedicated to the genetics and genomics of tropical and Mediterranean plants and their pathogens

génomique formations ressources montpelliérain
Infrastructure internationale
plantes orientée développement
Ressources sud service développement
compétences plateforme d'analyses
végétale communauté outils multi-instituts
s'appuie mutualisation partage



SNP detection genome assembly
phylogeny transcriptome assembly structural variation
comparative genomics differential expression
GWAS pangenomics
population genetics megabases
polyploidy

Mutualisation



Cacao



Banana



Coffee



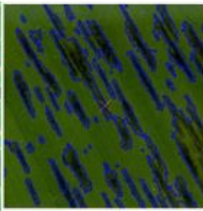
Rice



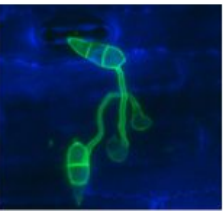
Palm



Cassava



Pseudocercospora



Magnaporthe

South Green

bioinformatics platform



4 institutes



3 research units

Tools

Storage and computing
resources



Trainings





Meso@LR au CINES

1090 threads :

35 standard nodes

2 bigmem nodes

1 GPU node

500 To of replicated storage



CINES

1130 threads:

30 standard node

1 supermem node

1 GPU node

150 To on 3 NAS + 210 To scratch

slurm
workload manager
400+



600+ tools

Resources mutualised at Meso@LR through the
Mudis4Ls project (purchase/storage/data)

Collaborative development of tools

Genomics

Pangenomic

Gene families

Comparative

Phylogeny

Assemblies

Annotation

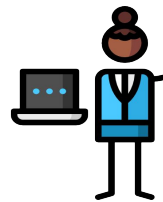
Data mining

Diversity exploration

genotype manipulation

mosaic manipulation

Metagenomic



+20
tools



web applications (16)



visualisation (8)



workflows(5)



packages (4)



<https://github.com/SouthGreenPlatform/>

I-Trop

Plant & Health Bioinformatics Platform



<https://bioinfo.ird.fr/>



AURORE
COMTE



JACQUES
DAINAT



ALEXIS
DEREEPER



BRUNO
GRANOUILLAGC



JULIE
ORJUELA-



NDOMASSI
TANDO



CHRISTINE
TRANCHANT



bioinfo@ird.fr



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South Green

bioinformatics platform



Florian Charriat
Antoni Exbrayat



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Bruno Granouillac
Jacques Dainat



Nicolas Fernandez

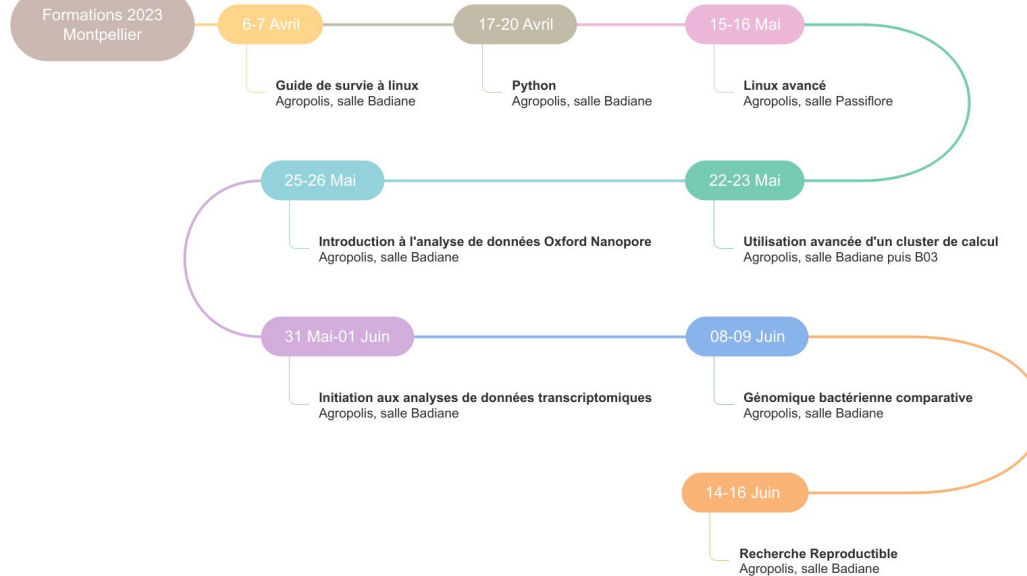


Thomas Denecker

And more collaborators !

South Green

bioinformatics platform



Modules de formation 2023

- Toutes nos formations :
<https://southgreenplatform.github.io/trainings/>
- Topo & TP :
https://github.com/SouthGreenPlatform/training_ONT_teaching/tree/2023_MTP
- Environnement de travail : [Logiciels à installer](#)

Initiation à l'analyse de données Oxford Nanopore



Alliance

Bioinformatics resources

On va travailler sous Linux !

- 2 façons d'utiliser linux :

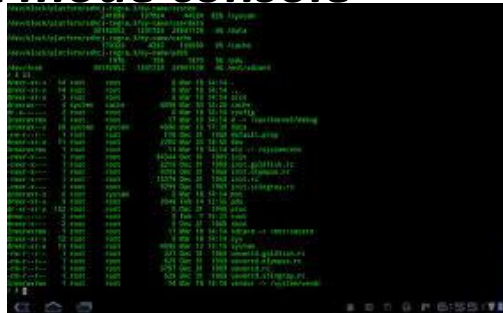
en *mode graphique*



En mode terminal

- 2 façons d'utiliser linux :

en *mode console*



avant tout !

Bases de Linux

https://github.com/SouthGreenPlatform/training_NT_teaching/blob/main/slides/GuideDeSurvieLinux-french2022.pdf

En mode jupyter book

- Une troisième façon d'utiliser linux :

en *mode jupyter book*



Sur le cloud IFB!



Let's discover Jupyter !

Working environment

What is jupyter book ?

- One of the most popular tool among data scientists to perform data analysis
- Provides a complete environment in which numerous programming languages can be used through a simple web browser

ex : Bash (Linux), Python, Java, R, Julia, Matlab, Octave, Scheme, Processing, Scala

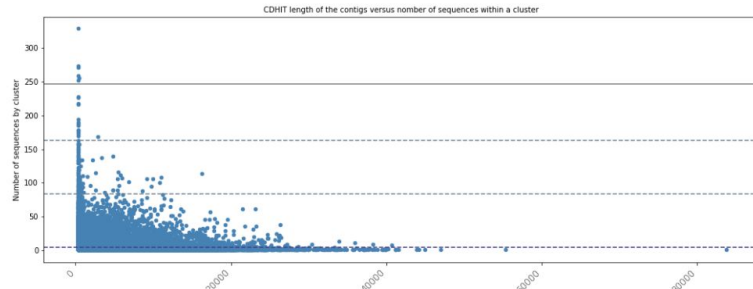


Why use jupyter book ?

An unique interface/file where text,code and output codes can be mixed :

- code can be executed inside each cell of the notebook
- code output is directly displayed in the notebook

```
All ctgs
Entrée [30]: 1 plt.figure(figsize=(17, 6))
2 ax = plt.gca()
3 df_cdhit[(df_cdhit.sp == '0b')].plot(x='pb', y='ln', kind="scatter", color='steelblue', ax=ax, linewidth=1)
4 df_cdhit[(df_cdhit.sp == '0g')].plot(x='pb', y='ln', kind="scatter", color='steelblue', ax=ax, linewidth=1)
5 plt.axhline(y=5, color='darkslateblue', linestyle='--')
6 plt.axhline(y=84, color='slategrey', linestyle='--')
7 plt.axhline(y=163, color='slategrey', linestyle='--')
8 plt.axhline(y=247, color='grey', linestyle='--')
9
10 plt.title("CDHIT length of the contigs versus number of sequences within a cluster", fontsize=10)
11 plt.xlabel('Cluster length')
12 plt.ylabel('Number of sequences by cluster')
13 plt.xticks(
14     rotation=45,
15     horizontalalignment='right',
16     fontweight='light',
17     fontsize=12,
18 )
19 plt.show()
20
```



Why use jupyter book ?

An unique interface/file where text,code and output codes can be mixed :

- code can be executed inside each cell of the notebook
- code output is directly displayed in the notebook
- explanations, formulas, charts can be added

The screenshot shows a Jupyter Notebook interface with the following details:

- Header:** jupyter parseCistr-Copy1 Dernière Sauvegarde : Il y a 8 minutes (auto-sauvegarde) Se déconnecter Python 3 O
- Toolbar:** Fichier Édition Affichage Insérer Cellule Noyau Widgets Aide
- Cell Content:**
 - Section Header:** Anchoring data analysis
 - Section 1:** 1 - CDHIT data analysis *before anchoring on genome*
 - Section 1.1:** 1.1 Removing redundancy with CDHIT
 - CDHIT Input : 1,306,676 contigs assembled from no mapped reads
 - Tests & results
 - Table:** A table showing cluster statistics:

	0.9	0.95
0.80	378,615	484,394
0.85	418,136	531,326
0.90	473,270	588,983
0.95	544,441	659,658
 - Text:** clusters generated after cdhit analysis : 484,394
 - Section 2:** 1.2 Converting cdhit file into a csv loaded as a dataframe with pandas
 - Text:** The script cdhitVsAnchoring.py creates the csv file allCtgtsIRIGIN_TOG5681.dedup8095.PANDAS.csv
 - Section 3:** Load csv file into a pandasframe
 - Code Cell [1]:**

```
Entrée [1]: 1 import pandas as pd
2 import matplotlib.pyplot as plt
3 import numpy as np
4
5 csv_cdhit_file = "/home/christine/Documents/These/Data/CDHIT/ALL_CGTGS_MERGE/allCtgtsIRIGIN_TOG5681.dedup8095.PANDAS.csv"
6 df_cdhit= pd.read_csv(csv_cdhit_file,names=['ctg','sp','ctg-list','sp_list'], header=0)
7 #print(df_cdhit)
8
```

Lab notebook for science data ?

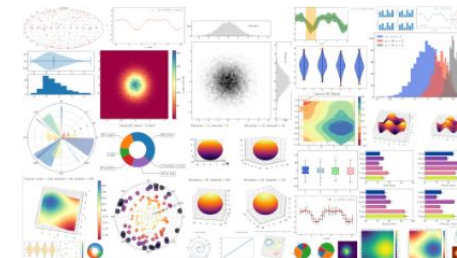
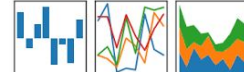


- One file to analyze data and generate reports
- Can be exported to many formats, including PDF and HTML, which makes it easy to share your project with anyone.
- Analysis are more transparent, repeatable and shareable

How to become a super datascientist ?

- easily import/export tabular files into/from dataframes (similar to R dataframe).
- manipulate these data tables / DataFrames
- easily draw beautiful graphs from these DataFrames with matplotlib

pandas
 $y_{it} = \beta' x_{it} + \mu_i + \epsilon_{it}$



How will you use Jupyter Notebook ?

- Launch our analyses through a jupyter book within a virtual machine launched via the IFB cloud “BIOSPHERE”



How will you use Jupyter Notebook ?

- Launch our analyses through a jupyter book within a virtual machine launched via the IFB cloud “BIOSPHERE”
- Through this virtual machine, we will create jupyter books and execute all our analysis

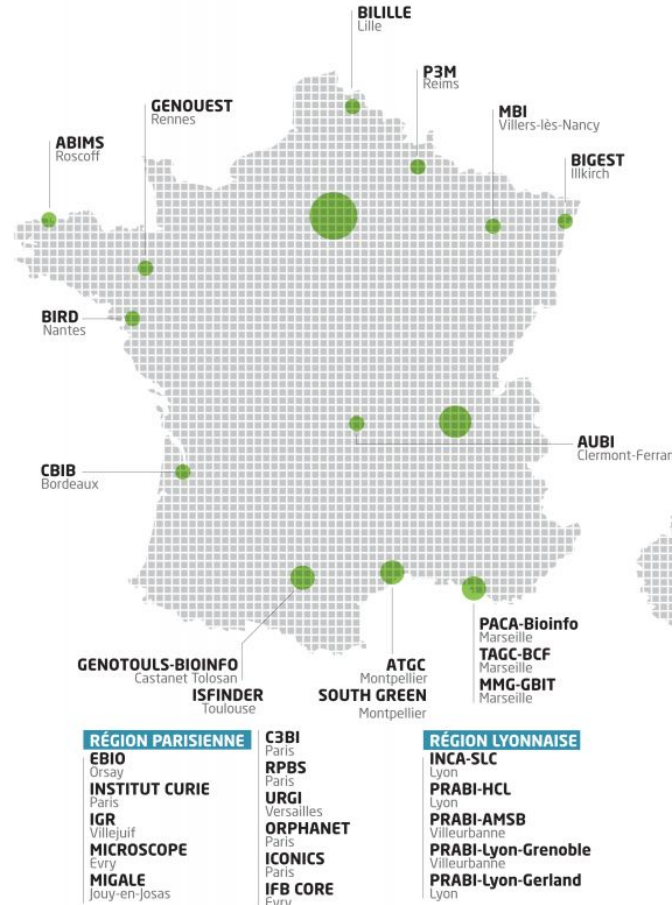


A screenshot of a web browser window titled "IFB Cloud". The address bar shows "mydatalocal/" and the URL "https://134.158.247.8/tree/mydatalocal". The main content area displays a "jupyter" interface with tabs for "Files", "Running", and "Clusters". A message says "Select items to perform actions on them." Below it, there's a file list for "mydatalocal" with a single item "jil ya q". A message at the bottom says "La liste des notebooks est vide." To the right, there's a "Notebook" dropdown menu with options like "Bash", "Julia 1.5.3", "Python 3", and "R". A "New" button is also visible. On the far right, there's a sidebar with sections for "Other:", "Text File", "Folder", and "Terminal".

IFB ?



22 plateformes-membres
7 plateformes contributrices
8 équipes associées
>400 experts (~200 FTE)

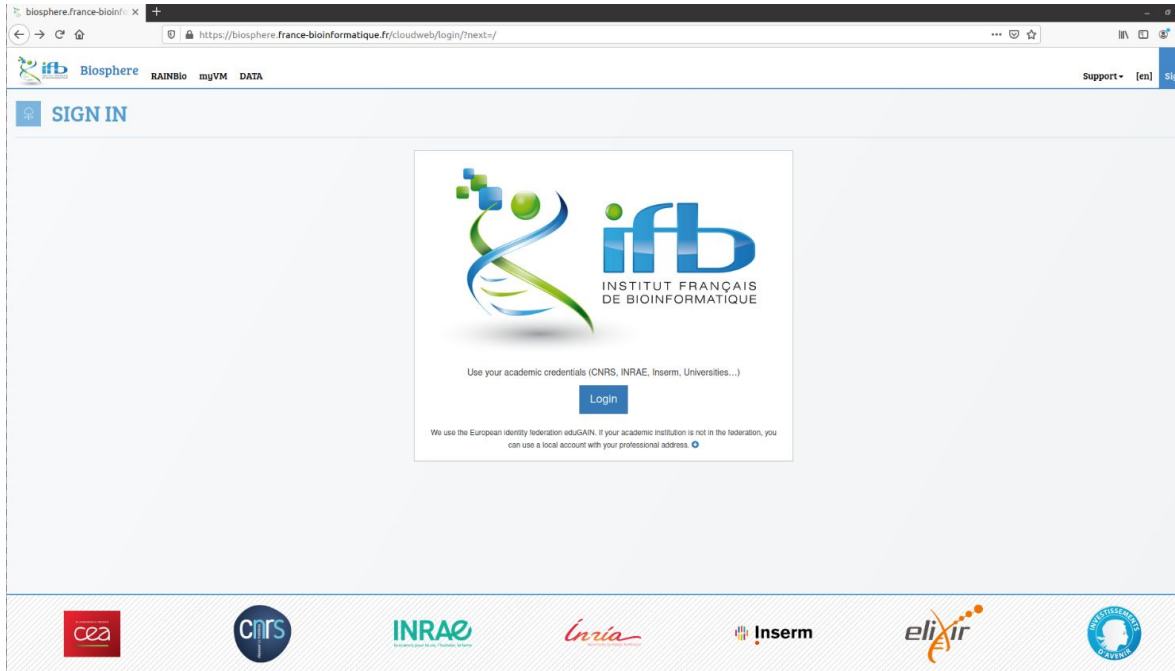


Biosphere, IFB CLOUD FOR LIFE SCIENCES

- A federation of clouds, which relies on interconnected IFB's infrastructures, providing distributed services to analyze life science data
- Access to a large set of virtual machines (computing resources, bioinformatics tool)
- Used for scientific production in the life sciences, developments, and also to support events like cloud and scientific training sessions, hackathons or workshops.

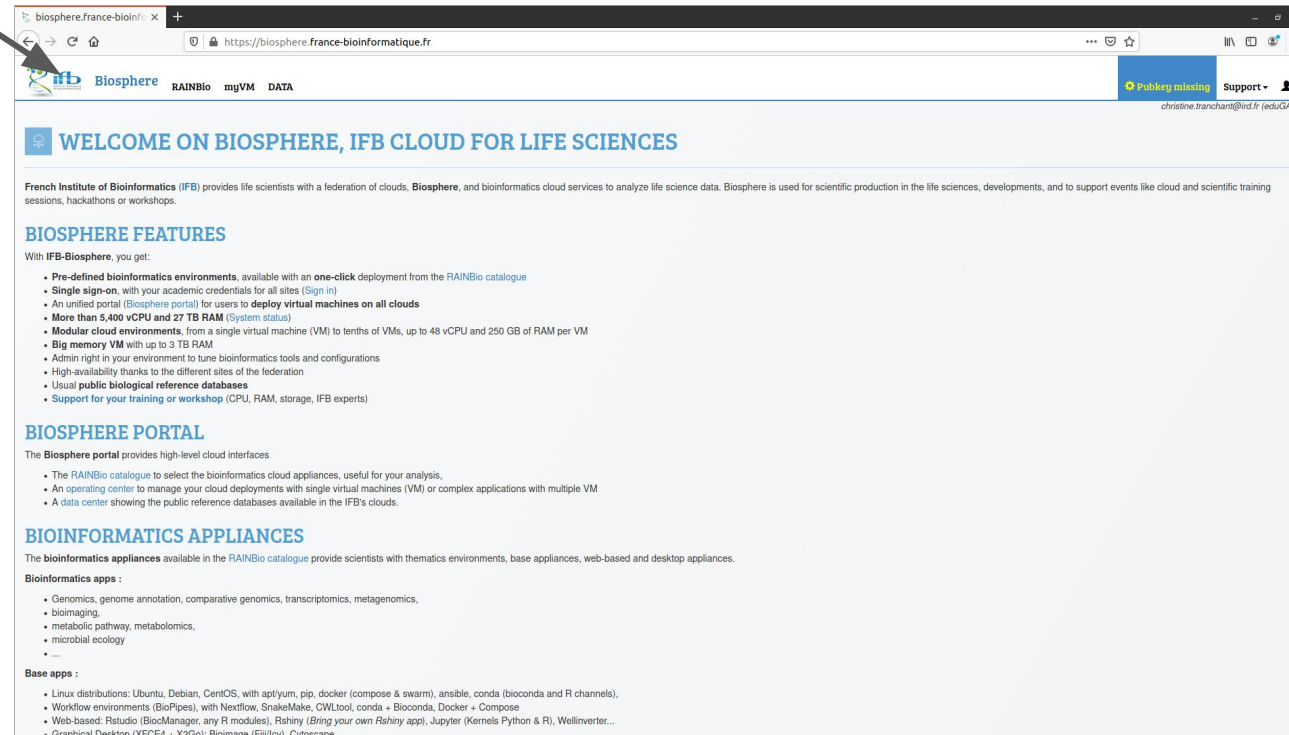
Let's start with biosphere

- Open the biosphere website : <https://biosphere.france-bioinformatique.fr/cloud/> and sign in



Connected / here we are

RAINBIO catalog to access our Virtual Machine (VM)



The screenshot shows a web browser window with the URL <https://biosphere.france-bioinformatique.fr>. The page title is "WELCOME ON BIOSPHERE, IFB CLOUD FOR LIFE SCIENCES". The content includes sections on BIOSPHERE FEATURES, BIOSPHERE PORTAL, and BIOINFORMATICS APPLIANCES, each listing various services and tools available through the RAINBio catalogue.

BIOSPHERE FEATURES

With IFB-Biosphere, you get:

- Pre-defined bioinformatics environments, available with an one-click deployment from the RAINBio catalogue
- Single sign-on, with your academic credentials for all sites ([Sign in](#))
- An unified portal (Biosphere portal) for users to [deploy virtual machines on all clouds](#)
- More than 5.400 vCPU and 27 TB RAM ([System status](#))
- Modular cloud environments, from a single virtual machine (VM) to tenths of VMs, up to 48 vCPU and 250 GB of RAM per VM
- Big memory VM with up to 3 TB RAM
- Admin right in your environment to tune bioinformatics tools and configurations
- High-availability thanks to the different sites of the federation
- Usual public biological reference databases
- Support for your training or workshop (CPU, RAM, storage, IFB experts)

BIOSPHERE PORTAL

The Biosphere portal provides high-level cloud interfaces

- The RAINBio catalogue to select the bioinformatics cloud appliances, useful for your analysis.
- An operating center to manage your cloud deployments with single virtual machines (VM) or complex applications with multiple VM
- A data center showing the public reference databases available in the IFB's clouds.

BIOINFORMATICS APPLIANCES

The **bioinformatics appliances** available in the RAINBio catalogue provide scientists with thematic environments, base appliances, web-based and desktop appliances.

Bioinformatics apps :

- Genomics, genome annotation, comparative genomics, transcriptomics, metagenomics,
- biomining,
- metabolic pathway, metabolomics,
- microbial ecology
- ...

Base apps :

- Linux distributions: Ubuntu, Debian, CentOS, with apt/yum, pip, docker (compose & swarm), ansible, conda (bioconda and R channels),
- Workflow environments (BioPipes), with Nextflow, SnakeMake, CWLtool, conda + Bioconda, Docker + Compose
- Web-based: Rstudio (BioManager, any R modules), Rshiny (Bring your own Rshiny app), Jupyter (Kernels Python & R), Wellinverter...
- Graphical Desktop (XFCE4, Xfce, Bioimagine, Fiji/lov, Cytoscape)

Searching for the vm we will use

vm's name :

CoursAnalysesNanoporeSG

 **RAINBIO - APPLIANCES BIOINFORMATIQUES DANS LE CLOUD**

Catalogue des appliances bioinformatiques dans le cloud, filtrez-les en utilisant les termes présents dans l'ontologie EDAM, ou en langage naturel.

App Store (58) Appliances Outils Topics Appliance éditable Ajouter ⚙️

AnalysesSV ★ bcftools, BEDTools, BWA, Jupyter, Matplotlib, pandas, SAMtools ⚡ DNA polymorphism, Genetic variation, Genotyping experiment, GWAS study	CoursAnalysesNanoporeSG ★ bandage, Jupyter ⚡ Data architecture, analysis and design, Mathematics, Statistics and probability 🔧	virus_ONT ★ Jupyter ⚡ Data architecture, analysis and design, Mathematics, Statistics and probability 🔧	ANF MetaBioDiv ★ DESeq2, ggplot2, phyloseq, RStudio ⚡ Transcriptomics, Microbiology, Metagenomics, Sequence analysis
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Let's run your vm through the cloud

The screenshot shows a web-based interface for managing virtual machines. At the top, there is a navigation bar with the IFB Biosphere logo, RAINBio, myVM, and DATA links. On the right side, there is a user profile for 'julie.orjuela@ird.fr (eduGAIN)'.

The main content area displays a virtual machine named 'CoursAnalysesNanoporeSG'. It includes sections for 'Description' (VM used for train scientists and students from Burkina Faso and West Africa in bioinformatics analysis of data from Oxford nanopore sequencing technology with main of study viral métagenome.) and 'Domaines associés' (Computational biology, Sequence analysis).

In the 'Outils' section, there is a table with the following data:

Outils	
Jupyter	
OS	Debian 11
Recette de l'app (git)	https://github.com/SouthGreenPlatform/training_ONT_VM/tree/2022
App de base	Jupyter

A large green button labeled 'DÉPLOIEMENT AVANCÉ' is highlighted with a black arrow pointing towards it from the top right. Below it is another button labeled 'LANCEMENT'.

Let's run your vm through the cloud

The screenshot shows the IFB Biosphere interface for deploying a virtual machine (VM). The main title is "CoursAnalysesNanoporeSG". The deployment configuration window is open, titled "Configurer le déploiement d'une appliance". The sub-titile is "Déploiement de l'appliance 'virus_ONT'". The "Name" field is set to "Julie_ONT". The "Groupe à utiliser" dropdown shows "virus_ont (Initiation à l'analyse de la séquençage de virus)" and "tagé nome viraux) 828.01". The "Cloud" dropdown is set to "ifb-core-cloudbis". The "Gabarit d'image cloud" dropdown is expanded, showing various options. An arrow points to the "ifb.m4.2xlarge (8 vCPU, 32Go GB RAM, 200Go GB local disk)" option, which is highlighted with a blue selection bar. A tooltip above the dropdown asks "Quelle gabarit d'image doit être utilisé sur ce cloud ?". The background shows the IFB Biosphere dashboard with tabs like RAINBio, myVM, and DATA, and a user profile for julie.orjuela@ird.fr.

Description

VM used for train scientists and students from Burkina Faso and West Africa sequencing technology with main of study viral métagenome.

Configurer le déploiement d'une appliance

Déploiement de l'appliance "virus_ONT"

Name: Julie_ONT

Groupe à utiliser: virus_ont (Initiation à l'analyse de la séquençage de virus) tagé nome viraux) 828.01

Cloud: ifb-core-cloudbis

Gabarit d'image cloud:

- ifb.m4.large (2 vCPU, 8Go GB RAM, 50Go GB local disk)
- ifb.m4.xlarge (4 vCPU, 16Go GB RAM, 100Go GB local disk)
- ifb.m4.2xlarge (8 vCPU, 32Go GB RAM, 200Go GB local disk)**
- ifb.m4.4xlarge (16 vCPU, 64Go GB RAM, 400Go GB local disk)
- ifb.xt.e.4xlarge (BigMem) (16 vCPU, 384Go GB RAM, 600Go GB local disk)
- ifb.m4.6xlarge (24 vCPU, 96Go GB RAM, 600Go GB local disk)
- ifb.m4.8xlarge (32 vCPU, 128Go GB RAM, 800Go GB local disk)
- ifb.xt.e.8xlarge (BigMem) (32 vCPU, 768Go GB RAM, 600Go GB local disk)
- ifb.m4.12xlarge (48 vCPU, 192Go GB RAM, 1.2To GB local disk)
- ifb.xt.e.12xlarge (BigMem) (48 vCPU, 1.1To GB RAM, 50Go GB local disk)
- ifb.m4.14xlarge (56 vCPU, 240Go GB RAM, 1.4To GB local disk)
- ifb.xt.e.16xlarge (BigMem) (62 vCPU, 1.5To GB RAM, 1.5To GB local disk)
- ifb.xt.e.32xlarge (BigMem) (124 vCPU, 2.9To GB RAM, 2.9To GB local disk)

Computational biology

Annuler

VM/tree/2022

Support julie.orjuela@ird.fr (eduGAIN)

EDITER LANCER DÉPLOIEMENT AVANCÉ

Let's run your vm through the cloud

Loading...

The screenshot shows a web-based interface for managing cloud deployments. At the top, there are navigation tabs: IFB Biosphere, RAINBio, myVM, and DATA. On the right, there are links for Support (with an email address: julie.orjuela@ird.fr) and a user icon.

The main area is titled "CLOUD". It displays a table of "Déploiements" (Deployments). The columns include:

- ID: 19804, 19759
- Nom: virus_ONT (1.0), virus_ONT (1.0)
- Début: Sep 05 2022, 17h00; Sep 05 2022, 10h25
- Groupes: virus_ont, DIADE
- Spécification: 8 cores, 32 GB RAM, 200 GB disk (for ID 19804); 1 core, 4 GB RAM, 25 GB disk (for ID 19759)
- Broker: da98, b680
- Cloud: ifb-core-cloudbis (highlighted with a red arrow)
- Accès: (empty)

At the bottom left, there is a button labeled "Arrêter les déploiements" (Stop deployments).

Let's run your vm through the cloud

ready !

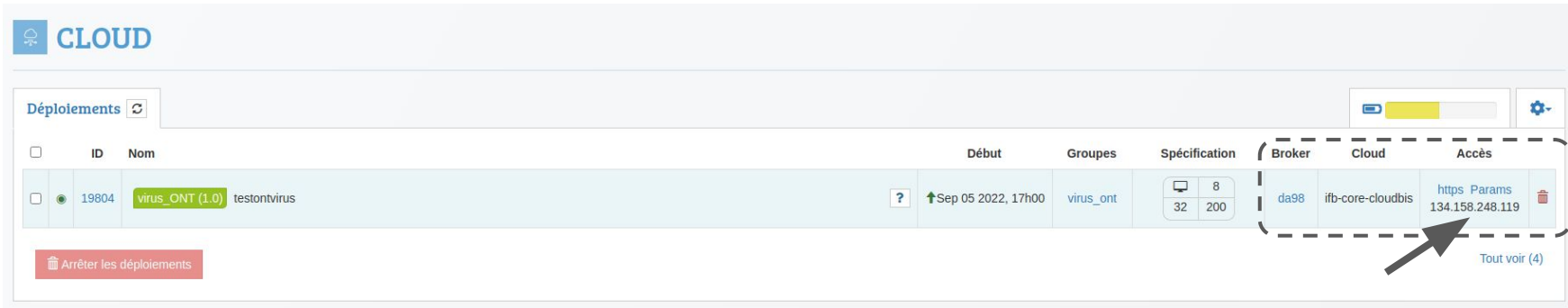
CLOUD

Déploiements

ID	Nom	Début	Groupes	Spécification	Broker	Cloud	Accès
19804	virus_ONT (1.0) testontvirus	Sep 05 2022, 17h00	virus_ont	8 32 200	da98	ifb-core-cloudbis	https Params 134.158.248.119

Arrêter les déploiements

Tout voir (4)



Let's run your vm through the cloud

get the url... link “https”

The screenshot shows a web-based interface for managing cloud deployments. At the top, there is a header with a cloud icon and the word "CLOUD". Below the header, a navigation bar includes "Déploiements" and a refresh icon. The main area displays a table of deployed VMs. The columns are labeled "ID", "Nom", "Début", "Groupes", "Spécification", "Broker", "Cloud", and "Accès". A single row is visible, representing a deployment named "virus_ONT (1.0)" with ID 19804. The "Début" column shows "Sep 05 2022, 17h00". The "Groupes" column contains "virus_ont". The "Spécification" column shows memory settings: "8" (with "32" and "200" below it). The "Broker" column lists "da98" and "ifb-core-cloudbis". The "Cloud" column lists "https Params" and the IP address "134.158.248.119". The "Accès" column has a trash bin icon. A red arrow points from the text "get the url... link “https”" to the "https Params" entry in the "Cloud" column. A red button at the bottom left says "Arrêter les déploiements".

Let's run our vm through the cloud

Get the token identifiant... link “Params”

The screenshot shows a cloud management interface with a modal dialog titled "Paramètres". Inside the dialog, there is a table with two columns: "nom" and "valeur". A single row is visible, containing "JUPYTER_TOKEN" in the "nom" column and "28f9a32ae92eaecbc816880489c9217e3263f9fd4614352" in the "valeur" column. The background of the interface displays a list of tasks or jobs. One job, named "virus", is shown in detail. The "Accès" (Access) section for this job includes a URL: "https://134.248.119.134". A yellow arrow points from the text "link ‘Params’" in the previous slide to this URL in the screenshot.

nom	valeur
JUPYTER_TOKEN	28f9a32ae92eaecbc816880489c9217e3263f9fd4614352

Début	Groupes	Spécification	Broker	Cloud	Accès
Sep 05 2022, 17h00	virus_ont	8 32 200	da98	ifb-core-cloudb1s	https://134.248.119.134

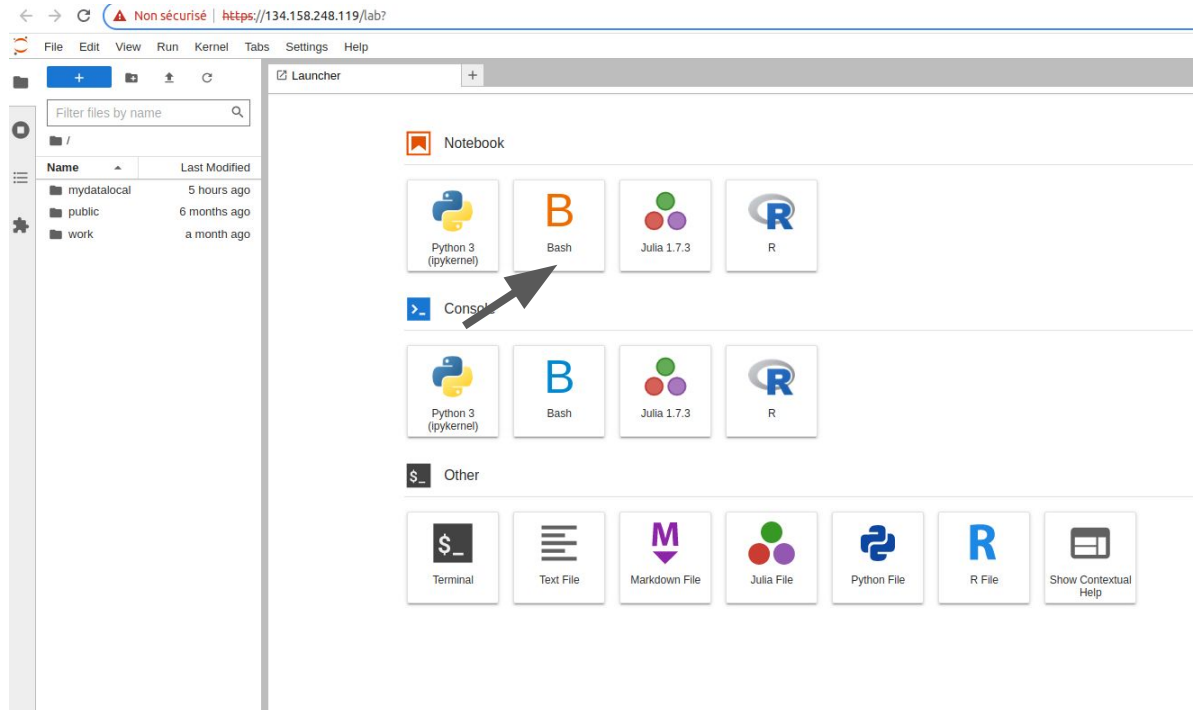
Let's run our vm through the cloud

Open your vm ([https link](https://134.158.247.8/tree)) to access to your own jupyter lab

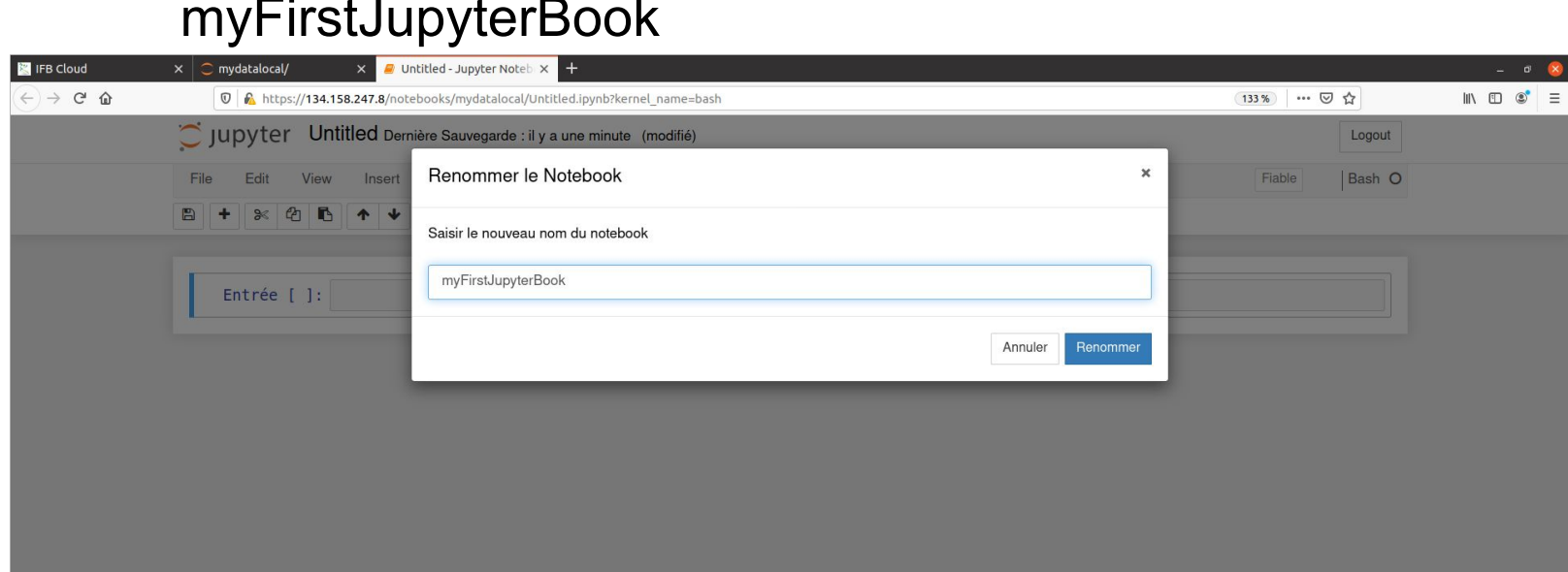


Create your first jupyter book

Go into the directory “work” and create a new jupyter book
-> kernel : bash



Rename your first jupyter book



Run your first bash command - *git clone*

All jupyterbook used for practice are here :

https://github.com/SouthGreenPlatform/training_ONT_teaching

Download all the jupyter books with the command *git clone*

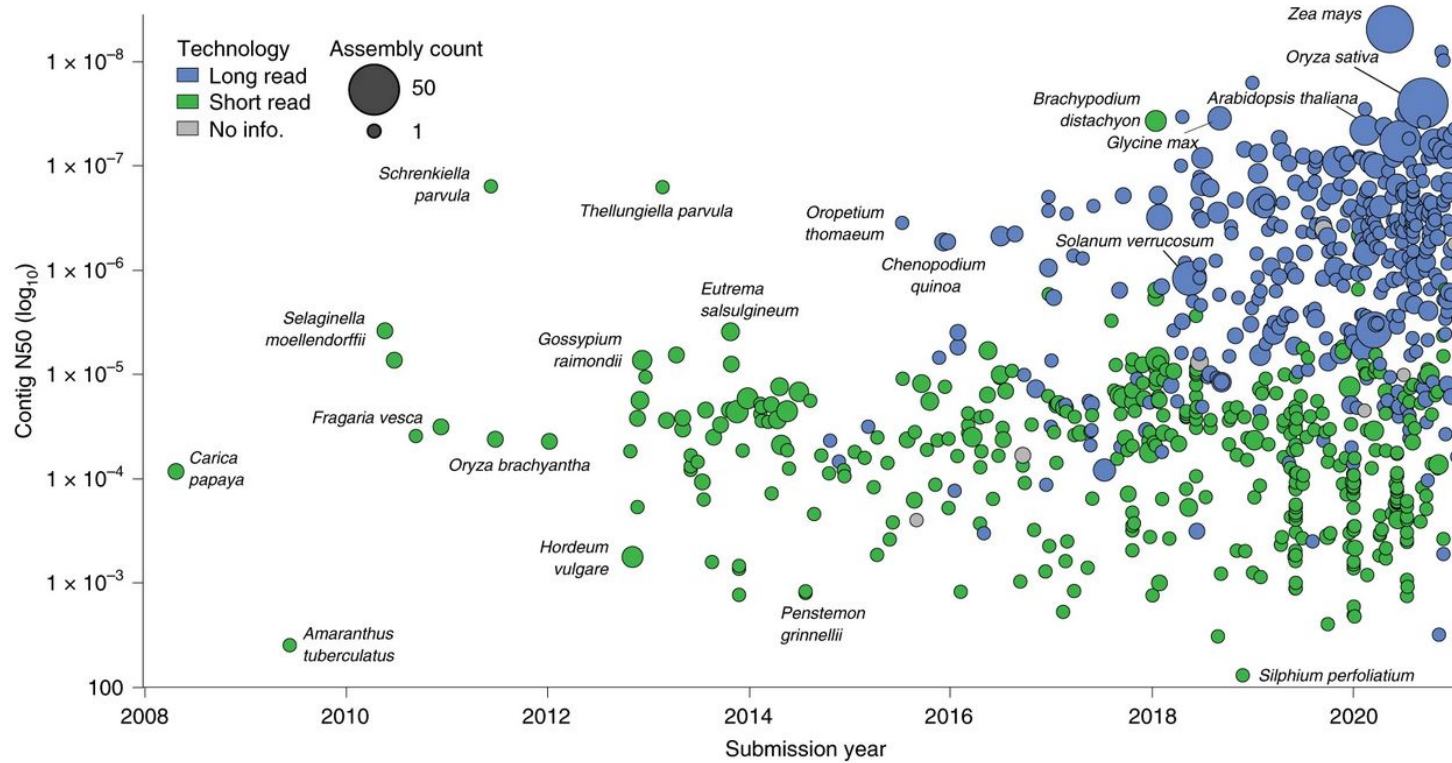
`git clone https://github.com/SouthGreenPlatform/training_ONT_teaching.git`

`git checkout 2023_MTP`

The screenshot shows a Jupyter Notebook interface. On the left, there's a file browser window titled 'work' showing two files: 'training_SV_teaching' (modified 6 minutes ago) and 'MyFirstJupyterBook.ipynb' (modified seconds ago). The main area has a title 'My first Juptyper book - Training SG SV'. Below it, a code cell displays the command 'My first linux command - pwd' followed by the output '[4]: /home/jovyan/work'. Another code cell below it shows the command 'Download all jupyter book we will use for this week - git clone' followed by the URL 'url https://github.com/SouthGreenPlatform/training_SV_teaching/tree/2022'. The final code cell at the bottom shows the command 'git clone --branch 2022 https://github.com/SouthGreenPlatform/training_SV_teaching.git' with its output, which includes progress messages like 'Cloning into 'training_SV_teaching'' and 'Unpacking objects: 100% (70/70), 134.35 KiB | 1.62 MiB/s, done.'

```
git clone --branch 2022 https://github.com/SouthGreenPlatform/training_SV_teaching.git
Cloning into 'training_SV_teaching'...
remote: Enumerating objects: 70, done.
remote: Counting objects: 100% (70/70), done.
remote: Compressing objects: 100% (48/48), done.
remote: Total 70 (delta 35), reused 49 (delta 20), pack-reused 0
Unpacking objects: 100% (70/70), 134.35 KiB | 1.62 MiB/s, done.
```

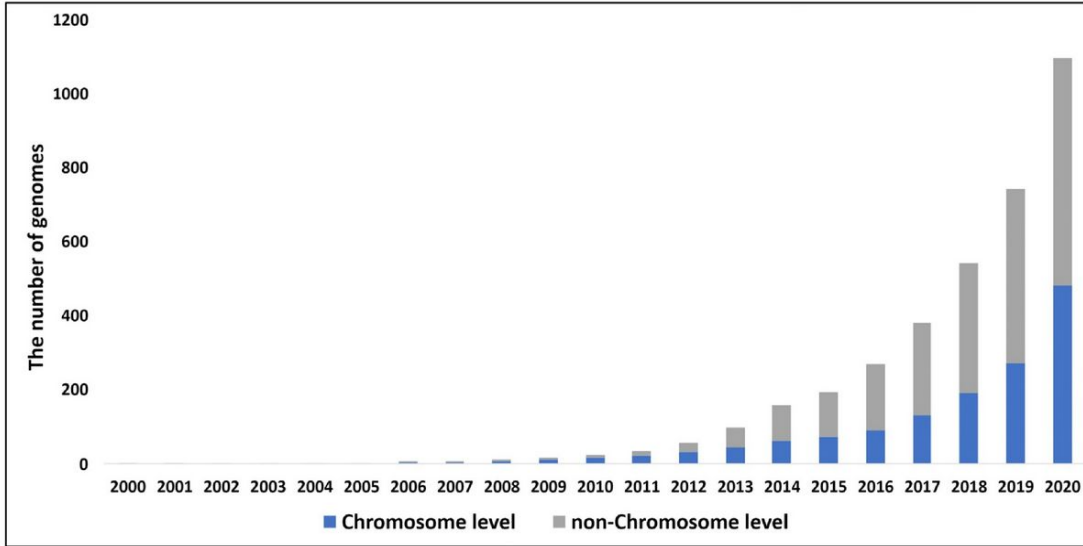
Let's start !



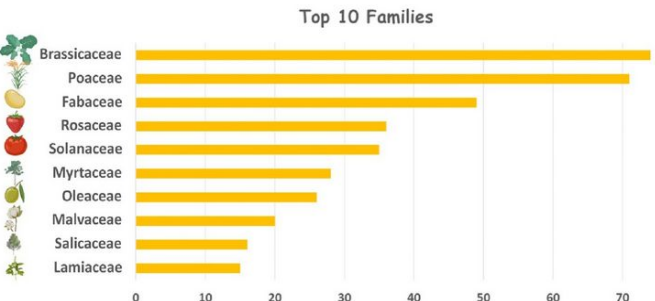
Assembly contiguity by submission date for 798 land plant species with publicly available genome assemblies. Points are coloured by the type of sequencing technology used and scaled by the number of assemblies available for that species. There is an improvement in contiguity associated with the advent of long-read sequencing technology, and a noticeable increase in the number of genome assemblies generated annually. All assemblies generated before 2008 have since been updated and are therefore not included.

Published plant genomes from 2000

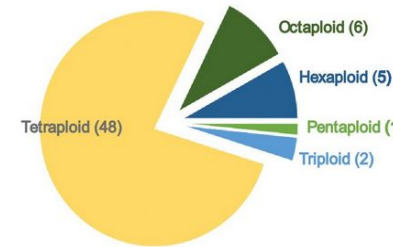
(A)



(B)



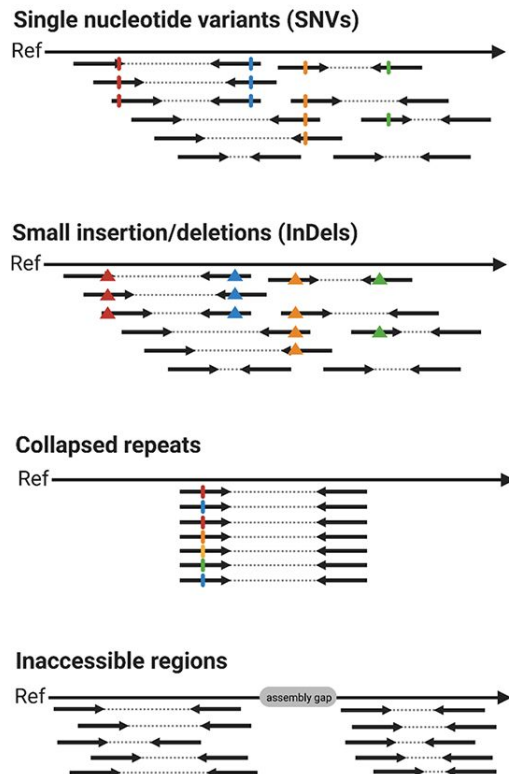
(C)



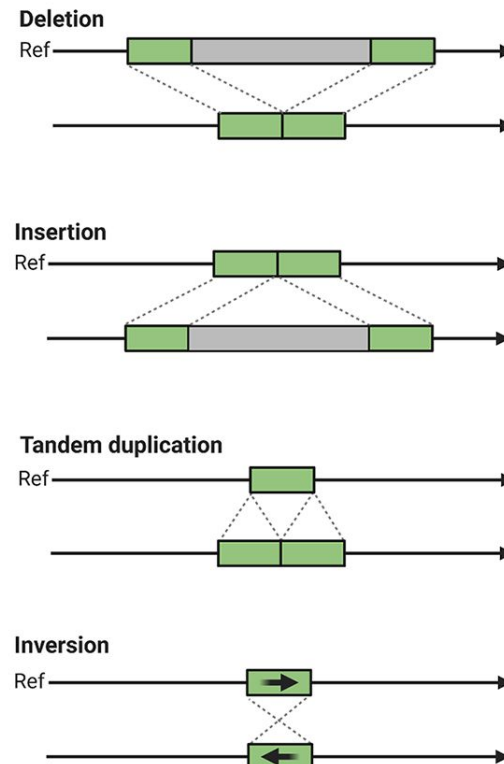
Sun et al, 2022

Long Reads - What you can do with it ?

A NGS variant calling

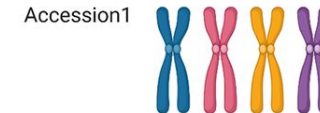


B long read variant calling

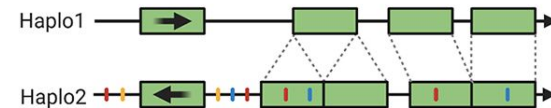


C *de novo* assembly

Chromosomal rearrangements



Separated haplophases

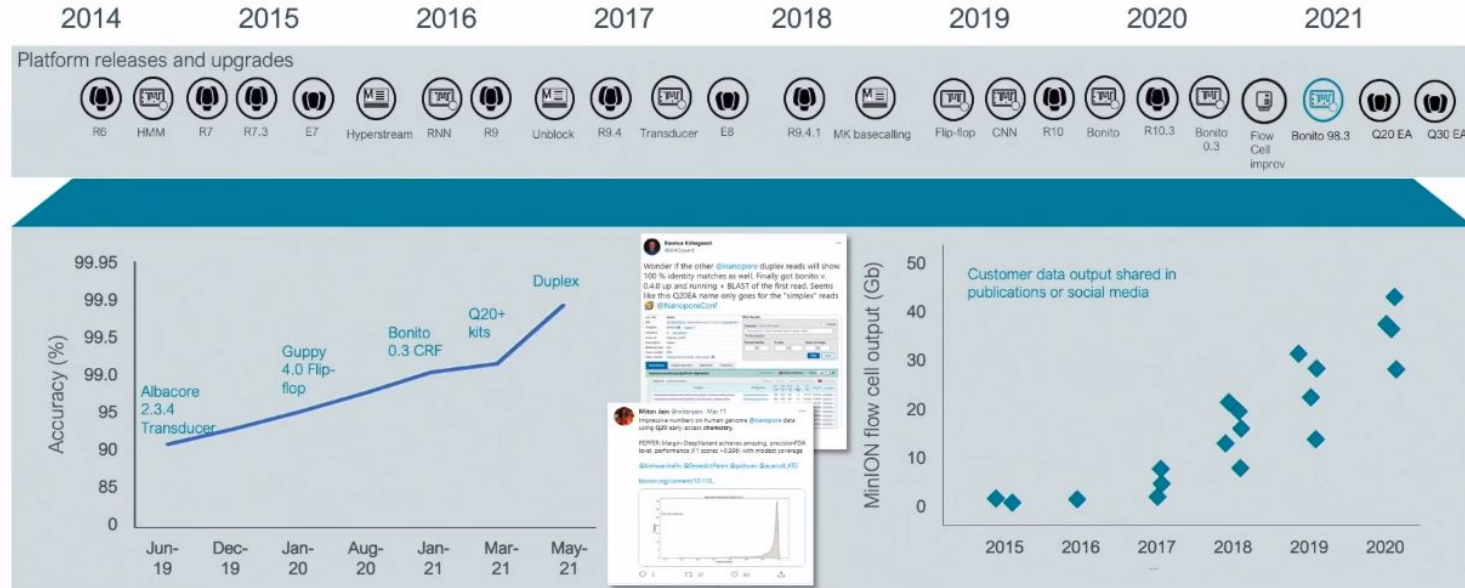


(Plant) genome project workflow from DNA extraction over ONT sequencing to data submission

	task	consumed time	hands-on time	equipment	estimated costs of consumables	estimated costs of lab equipment
A	 plant incubation in darkness	2-3d	1h			
B	 non-destructive sampling	-	1h			
C	 DNA extraction	1d	8h	waterbath, centrifuge	\$50	\$1000 \$8000
D	 quality control	1h	1h	NanoDrop, Qubit	\$20	
E	 short fragment depletion	2h	1h	centrifuge	\$50	
F	 quality control	1h	1h	NanoDrop, Qubit	\$20	\$5000 \$5000
G	 library preparation & sequencing	1-5d	4-16h	centrifuge, magnetic rack, sequencer	\$3000	\$250 \$1000
H	 basecalling	1d	1h	computer with GPU		\$3000
I	 assembly	1-15d	1h			
J	 polishing	1-5d	1h	compute cluster / cloud		
K	 annotation	1-5d	1h			
L	 data submission	2h	2h	fast internet connection		

Upgrades drive performance enhancements

...and core ones ship in consumables and software



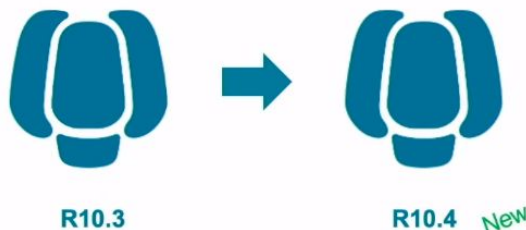
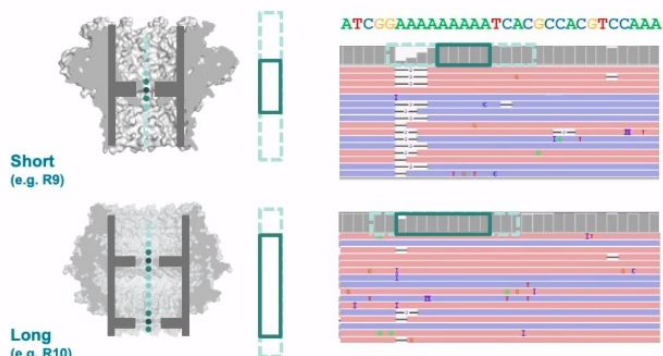
Last upgrades !

Oxford's Nanopores

R9 and R10

Short and long "reader heads"

- Length of the main discrimination site ("read head") affects accuracy
- Short read heads allow easier decoding of individual bases (R9 series)
- Longer read heads see more range and are more information rich (R10 series)



Improving the R10 series

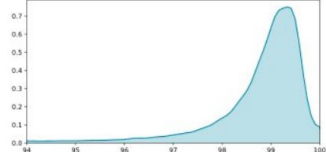
- We are continuously seeking to improve our nanopores
- R10 series of pores are still being iterated on – new R10.4 version
 - Extended discrimination profile, more sensing range
 - Higher flow cell yield of nanopores

← Tweet

 Oxford Nanopore @nanopore ...

Flow cells using our latest pore — R10.4 — can now be trialled through the expanding Q20+ Early Access Programme, which is now open to all applicants. Find out more about Q20+ and R10.4, and register to take part in the programme, here: bit.ly/3CEIJl9

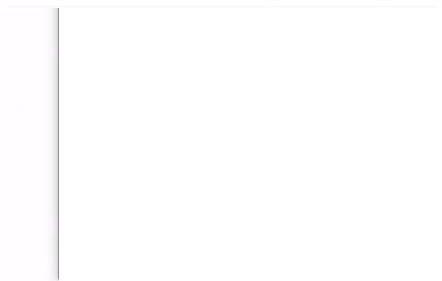
Raw read modal 99.3%, >Q20



A histogram showing the distribution of raw read accuracy percentages. The x-axis ranges from 94 to 100, and the y-axis ranges from 0.0 to 0.7. The distribution is highly skewed to the right, with the highest frequency (around 0.7) occurring at approximately 99.3% accuracy. The area under the curve is shaded light blue.

8:30 AM · Sep 23, 2021 · HubSpot

33 Retweets 1 Quote Tweet 62 Likes



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 OXFORD
NANOPORE
Technologies

<https://community.nanoporetech.com/posts/q20-early-access-group-br>

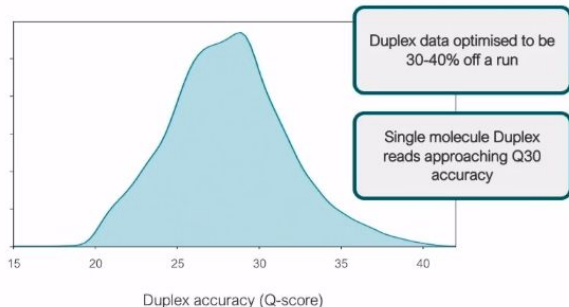
Last upgrades !

Nanopore accuracy

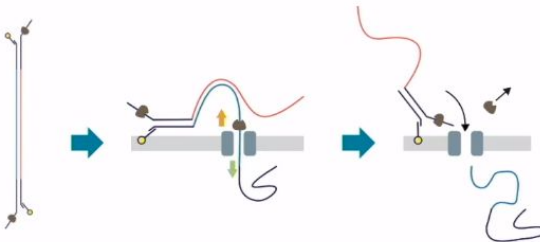
When we last spoke...

Duplex reads

- Possible when complement strand is sequenced immediately after template
- High duplex accuracy delivered by combining data template and complement
- New algorithms have been developed specifically for data combination
- Recent chemistries have optimised the amount of duplex data generated



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Generating duplex data

- Chances of seeing the complement follow template increased with Q20+ chemistry
- Early protocols available in EA community
- Longest Duplex Q30 read to date: 156 kbase

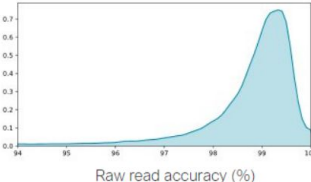


← Tweet

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Raw read modal 99.3%, >Q20



8:30 AM · Sep 23, 2021 · HubSpot

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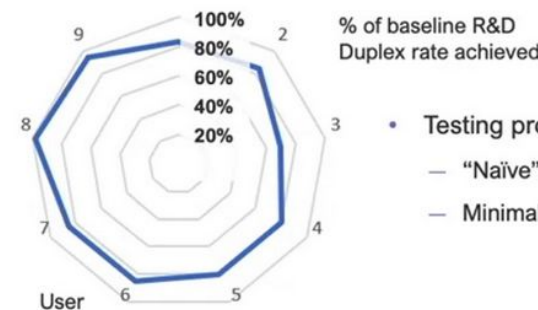
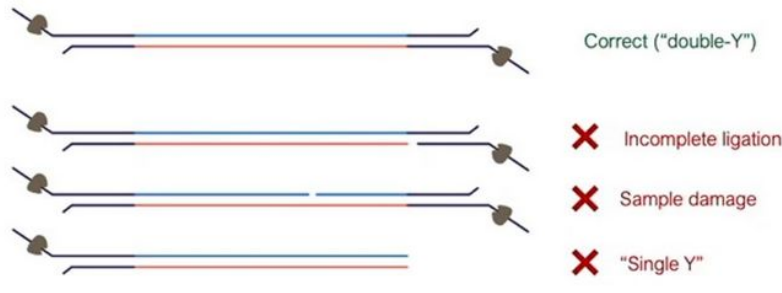
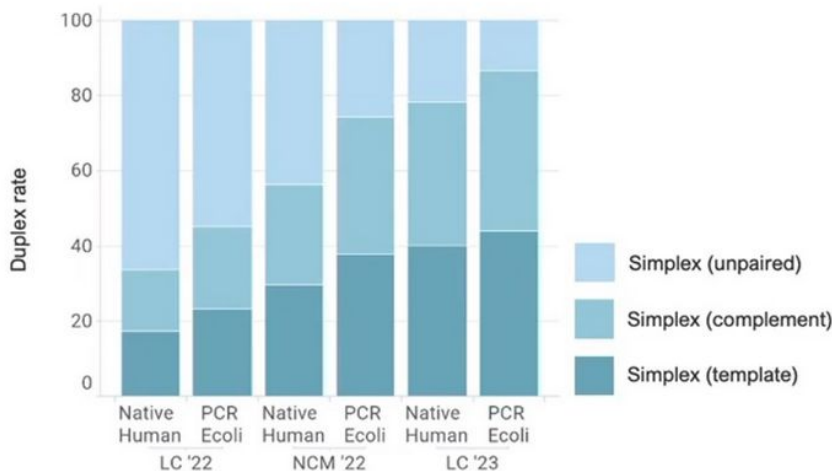


Duplex

Improved Duplex rates

Duplex rate

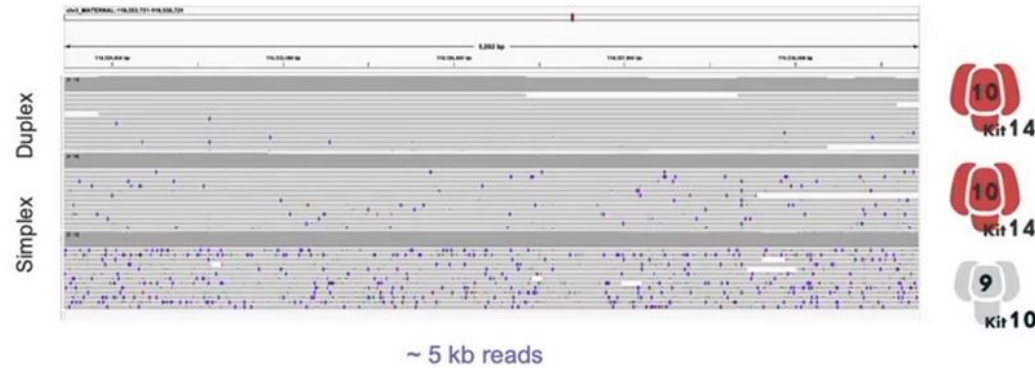
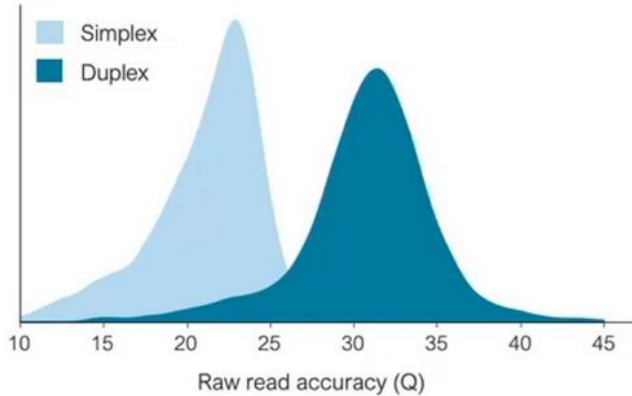
- Developer release of High Duplex flow cells in progress
- Expect majority of data in Duplex pairs (over 80%)
- Two-sided ligation achieved with LSK114



- Testing protocol robustness
 - “Naïve” nanopore users
 - Minimal prior library prep experience

Duplex Accuracy

Duplex accuracy



- Accuracy of Duplex raw reads ~ Q32 @ 5 kHz
- Runtime of new Stereo base caller similar to Simplex calls
- Duplex accuracy independent of read length
- Clearly visible improvements to errors

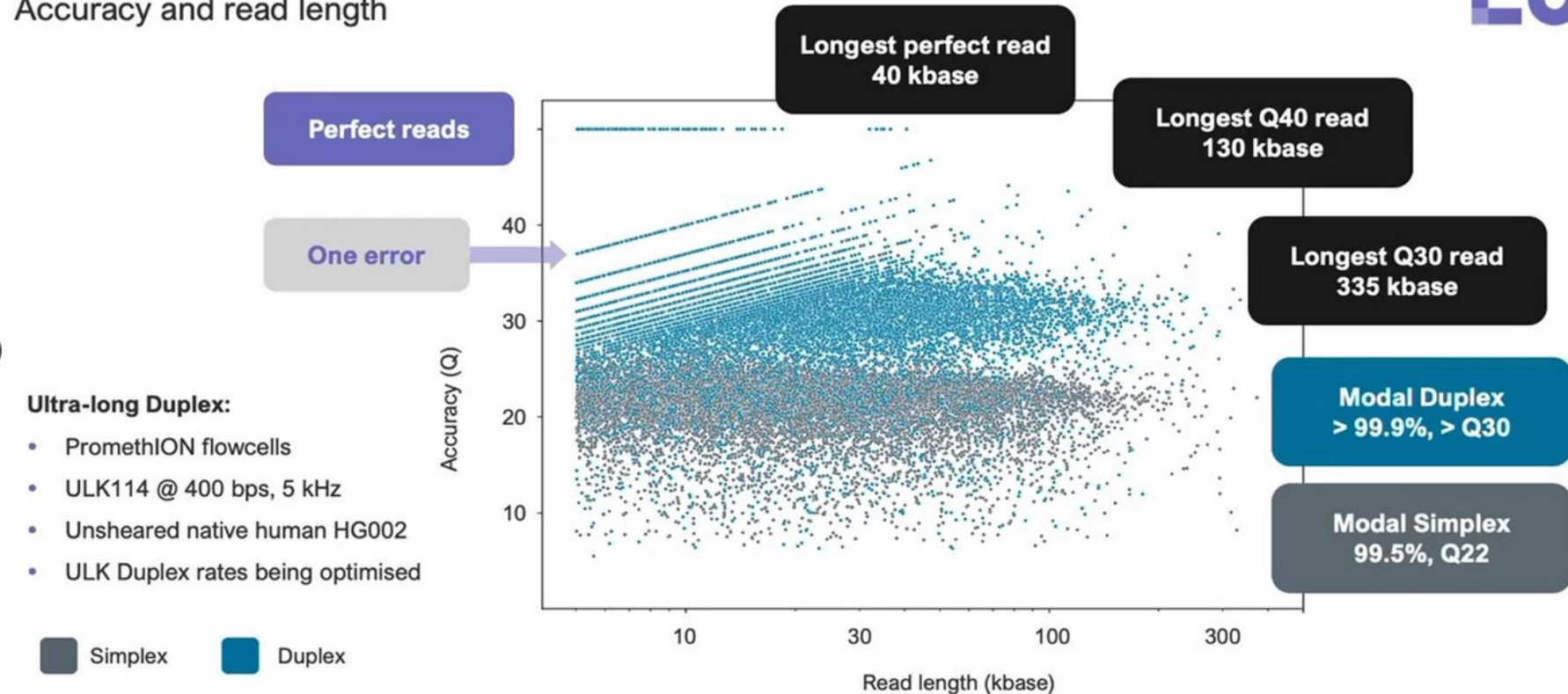


Alexander Wittenberg @AW_NGS · 30 Nov

@KeyGeneInfo received novel High duplex PromethION flow cells from @nanopore as Developer access. This pore is optimised to generate significant higher duplex data. Very high accuracy & long reads! Evaluation next week on first crops 🌱 #nanoporeconf

Duplex

Accuracy and read length

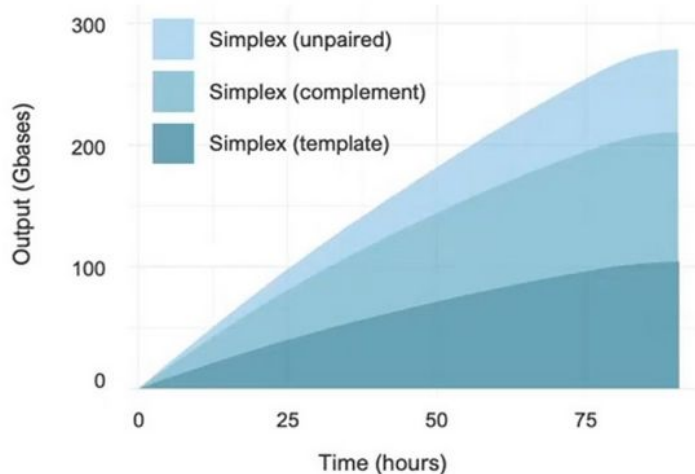


Duplex

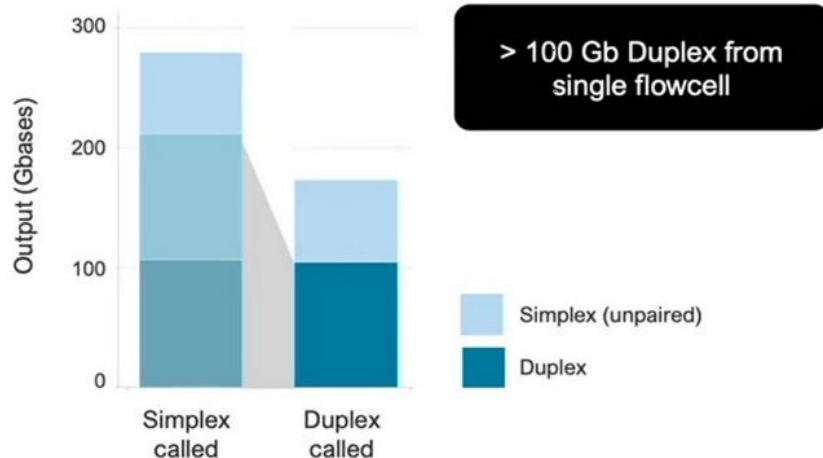
> 100 Gb of Duplex per flowcell

Higher output development Duplex runs

- Internal runs on Duplex output are very promising
- Achieving > 100 Gb of Duplex from a single PromethION flowcell
- Also provides > 50 Gb of high quality unpaired Simplex reads



- Development run conditions used for high output run
- Library preparation of PCR E.coli model sample
- Output equivalent to ~ \$6 Gb

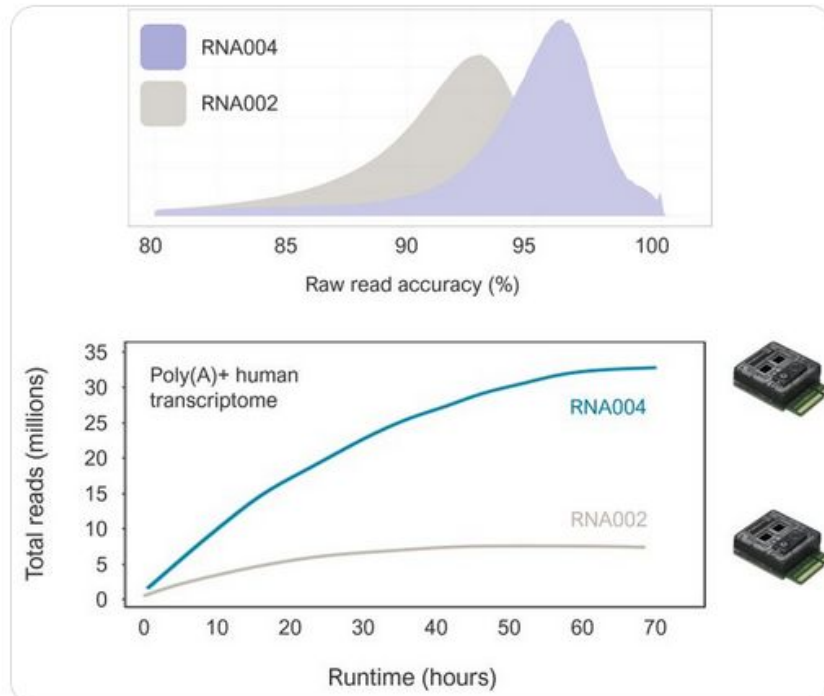




Oxford Nanopore ✨ @nanopore · May 18

...

Oxford Nanopore was the first to enable direct RNA sequencing. The new kit (RNA 004) and flow cell for direct RNA sequencing demonstrates increased accuracy and output, with the potential to unlock a new field of biological analysis. [#nanoporeconf register.nanoporetech.com/direct-rna](#)



1



48



150

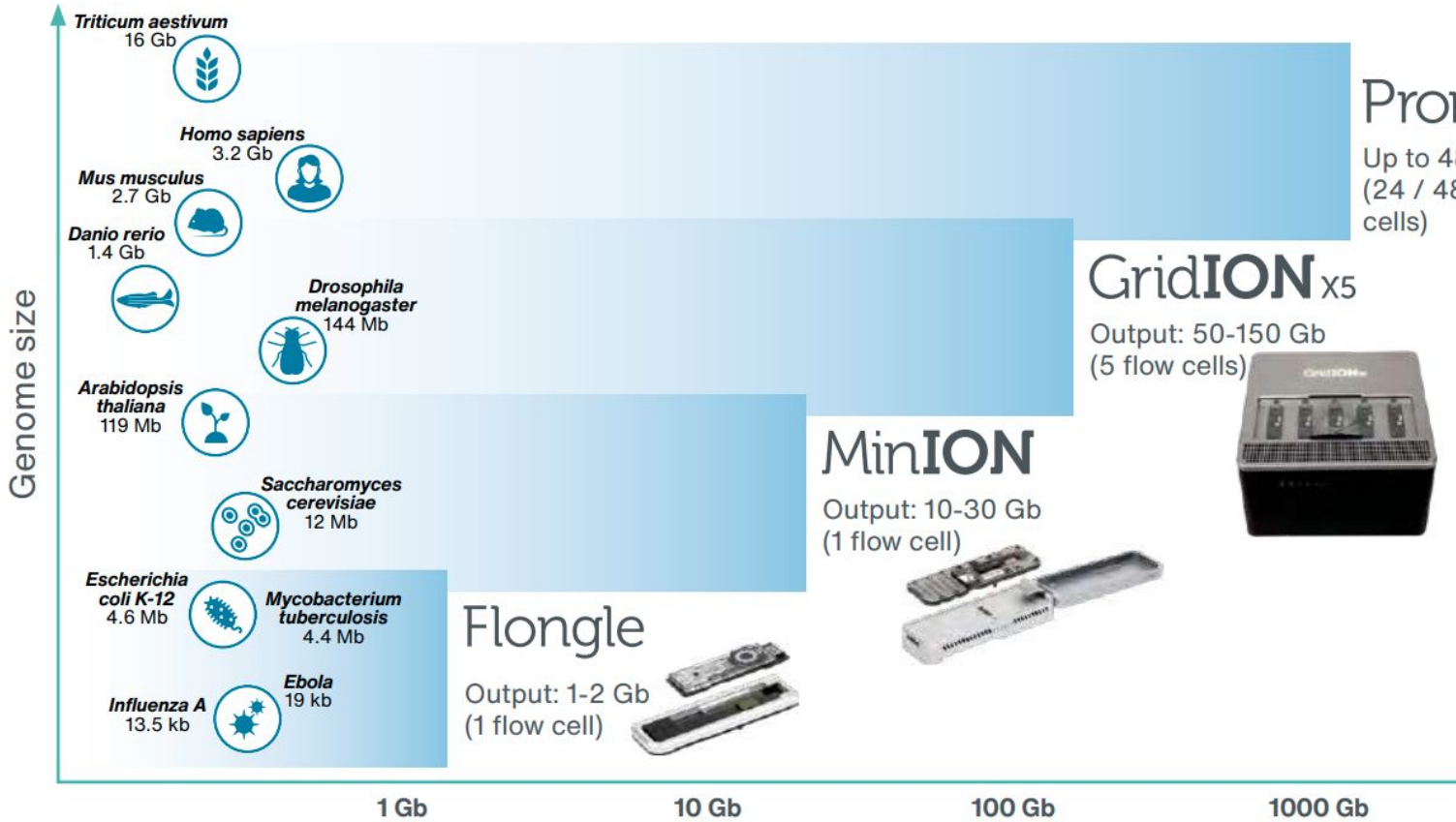


15.6K





A lot of data !



A lot of data !



MinION



MinION Mk1C



GridION



P2 Solo



P2



PromethION 24



PromethION 48



MinION and Flongle Flow Cell compatible

PromethION Flow Cell compatible

Configuration	Platform		Techniques		Tech specifications			
Number of flow cells per device	1	1	5		2	2	24	48
Maximum number of channels per flow cell	512	512	512		2,675	2,675	2,675	2,675
Run time	72 Hours	72 Hours	72 Hours		72 Hours	72 Hours	72 Hours	72 Hours
Device TMO ^t	50 Gb	50 Gb	250 Gb		580 Gb	580 Gb	~7 Tb	~14 Tb
Maximum number of flow cells per year*	104	104	520		208	208	2,596	4,992
Offer sequencing as a service	No	No	Yes		Yes	Yes	Yes	Yes

1	Platform	Run time max: (d)	Yield low: (Gb)	Yield high: (Gb)	Rate max: (Gb/d)	Reagents max: (\$)	Price per Gbp max: (\$)	Price per Gbp min: (\$)	hg-30x min instr amm (\$)	hg-30x min plus 5yr amm 10% serv 90% util (\$)	Machine: (\$ K)	Install base	Availability	Max total theoretical output per day (Gb) from current day (Gb)	Max total output per day (Gb) from installbase	
32	ONT P2 Solo 1fcell R10.4HD Q30+ data with Kit14	3.00	NA	60	20	1530-816	15.00	12.42	1242	1247	1271	10.5	430	worldwide	20	17,200
33	ONT P2 1fcell R10.4HD Q30+ data with Kit14	3.00	NA	60	20	1530-816	15.00	12.42	1242	1275	1410	60		worldwide	20	0
34	ONT P24 24fcell R10.4HD Q30+ data with Kit14	3.00	NA	1440	480	1530-655	15.00	10.00	1000	1005	1024	225	15	worldwide	535	8,025
35	ONT P48 48fcell R10.4HD Q30+ data with Kit14	3.00	NA	2880	960	1530-655	15.00	10.00	1000	1004	1018	310	60	worldwide	960	57,600

all prices of all technologies at may 2023

https://docs.google.com/spreadsheets/d/1GMMfhyLK0-q8XkIo3YxlWaZA5vVMuhU1kg41g4xLkXc/edit?hl=en_GB&hl=en_GB#gid=1569422585



Albert Vilella
@AlbertVilella

...

Oxford @nanopore has updated the pricing for the PromethION flowcells and so did I in the NGS specs spreadsheet. Numbers in the lined squares are for Q30+ Duplex data, based on the 60Gb per R10.4HD flowcell as per announced at #NanoporeConf



Matthew Miller - ↗ ... @b... · May 18

Wow, a bit of an early @nanopore surprise. Promethion flow cells now are just 900 USD, even at the four pack. As expected, this is about a 30% reduction in the cost to get a Promethion worth of Nanopore data. Huge news! #NanoporeConf



Oxford Nanopore ✅ @nanopore · May 18

Mk1D: the next generation of MinION portable sequencing.

Designed to be compatible with the iPad Pro which provides 5G connectivity & accelerated M1 processing. This combination delivers cutting edge nanopore tech with a market leading mobile platform

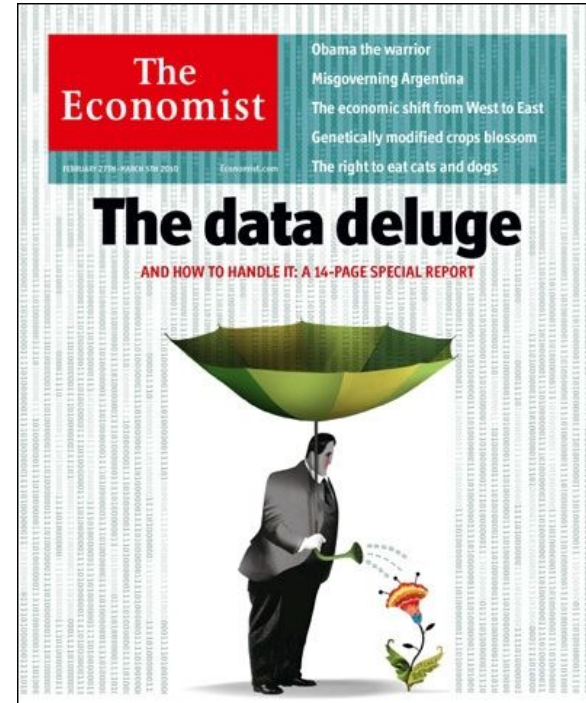
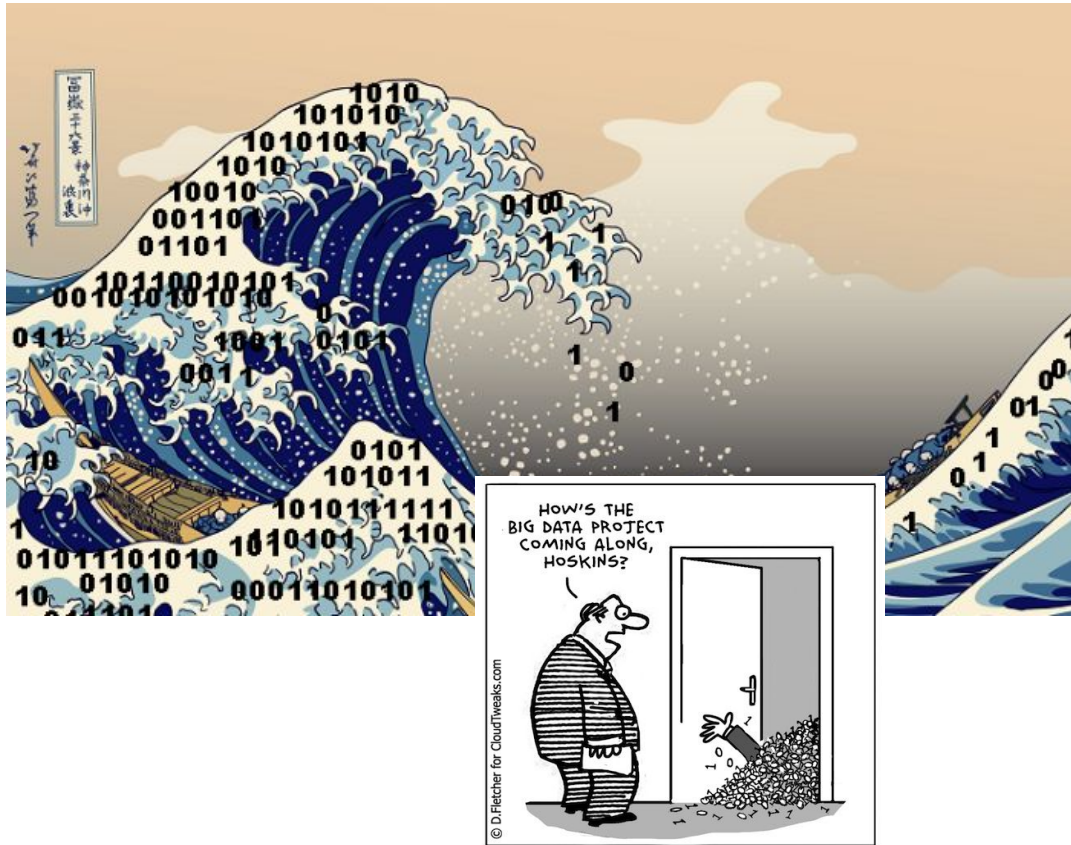
#nanoporeconf



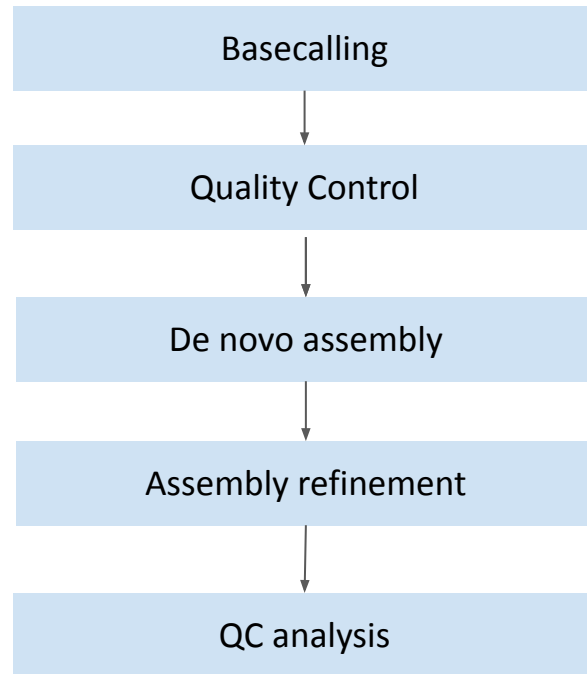
The data that these platforms produce differ qualitatively from second-generation sequencing, thus necessitating tailored analysis tools



From data rarity to data deluge

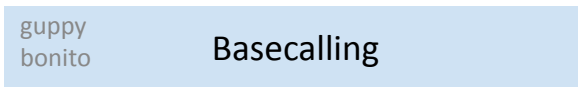


Typical long-read analysis pipelines for ONT data



Typical long-read analysis pipelines for ONT data

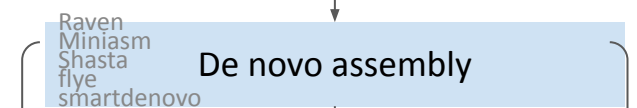
Demo



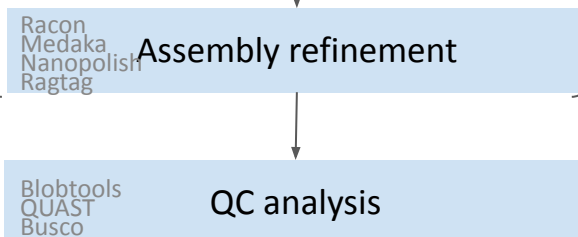
Practical 1



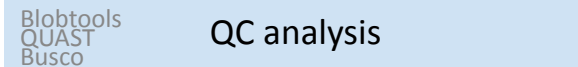
Practical 2



Practical 3



[Optional]



The Data !

The Pan-Genome of the cosmopolitan picophytoplankton *Bathycoccus prasinus*

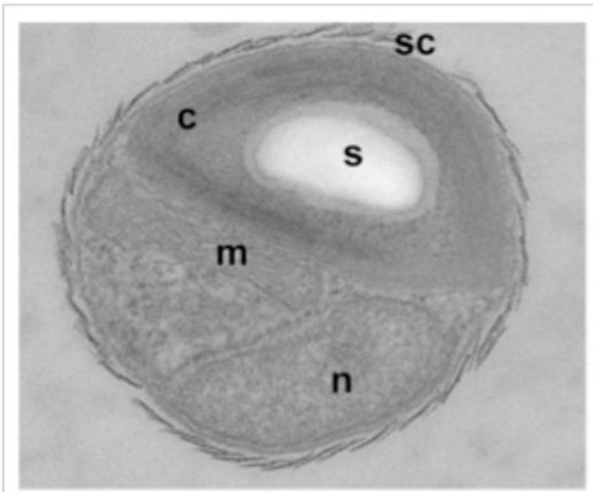
A first step towards understanding adaptation to latitude and seasons

Louis Denu, LOMIC

supervised by

François-Yves Bouget, LOMIC - Martine Devic, LOMIC - François Sabot, IRD Montpellier

Bathycoccus prasinus

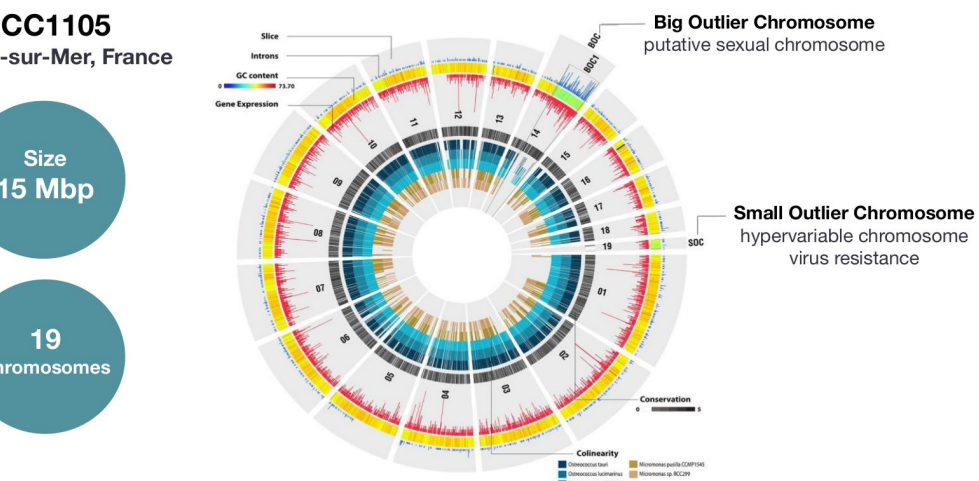


RCC1105
Banyuls-sur-Mer, France

Size
15 Mbp

19
Chromosomes

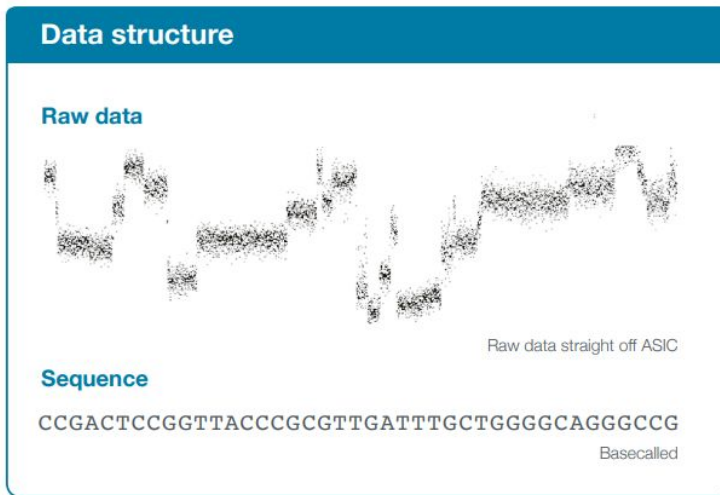
Moreau et al., *Genome Biology* 2012



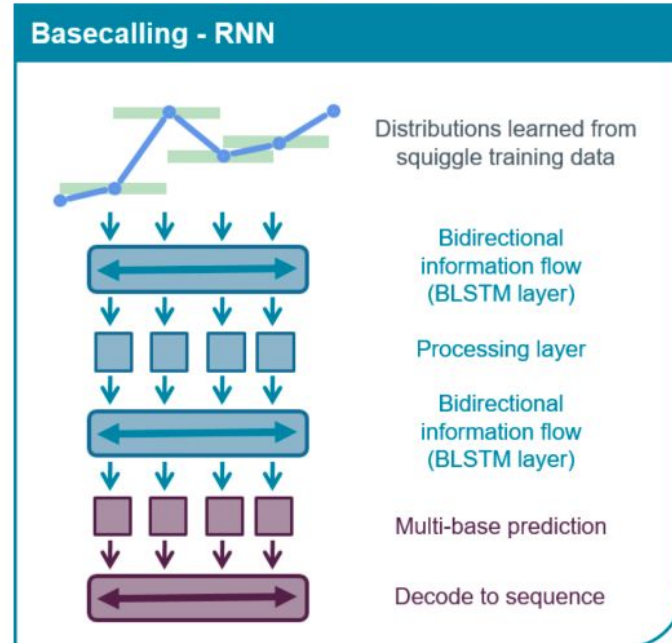
Chapitre 1

Reads Quality Control

ONT Read calling

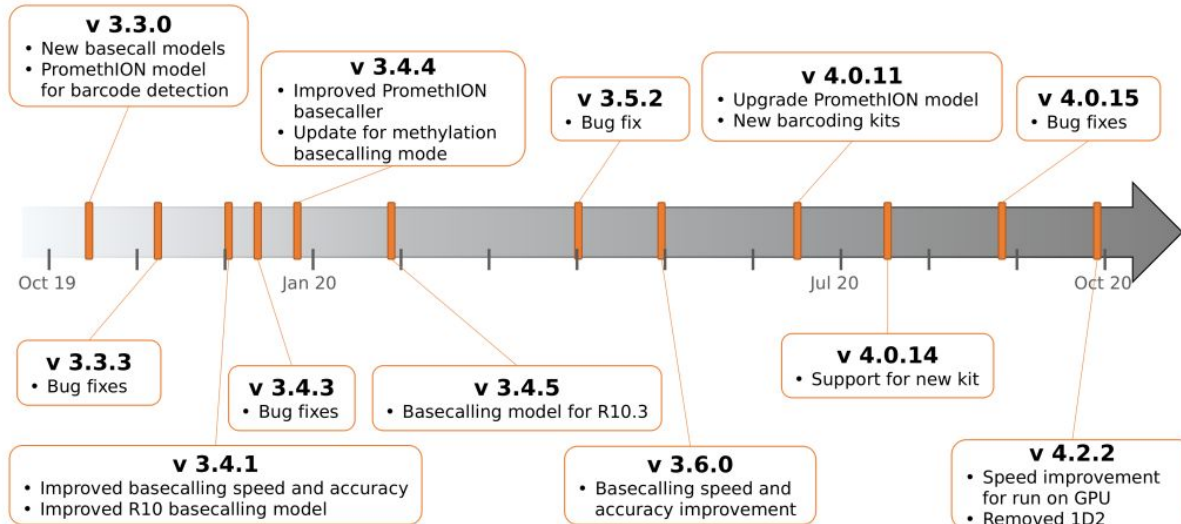


Reccurent Neural Network (RNN) – works like your brain! It can learn on the previous data and improve its performance on new data



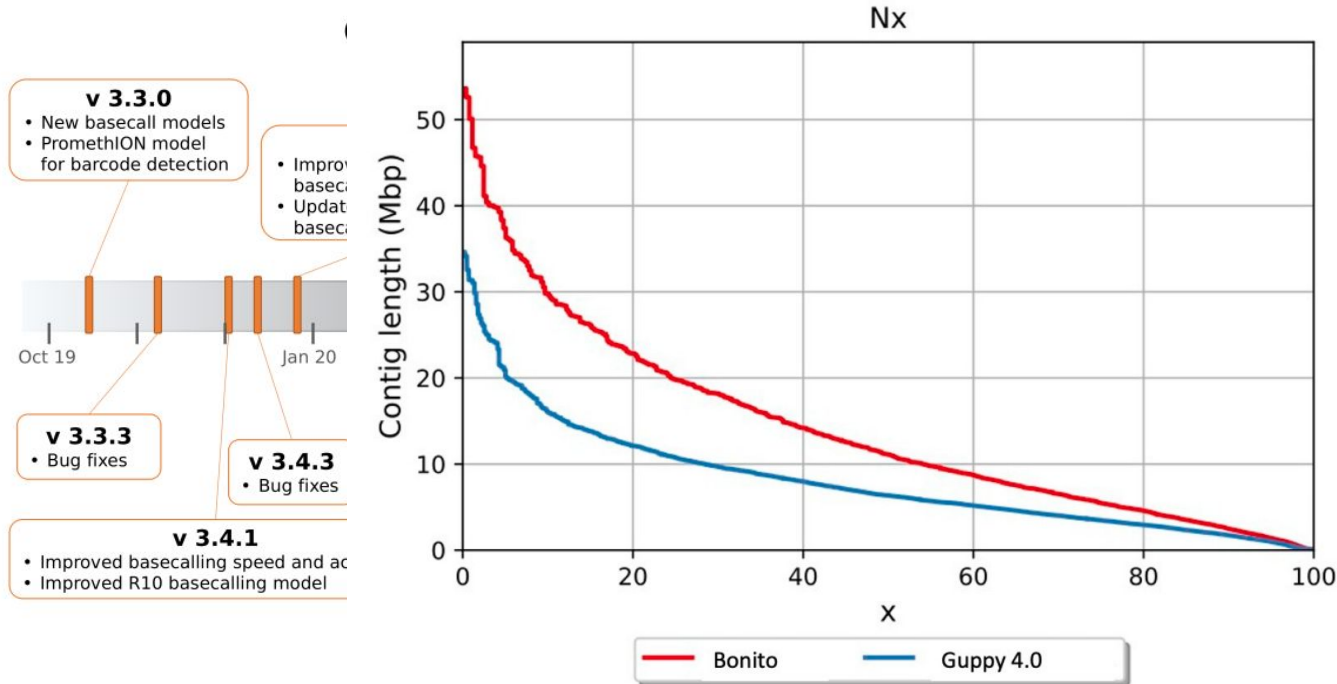
Nanopore basecallers are trained on many sequenced data, so you can run it on your data even if you are sequencing first time

Guppy basecaller releases



V 6.0.1
includes sup
models and now
can be specific
to some taxons !

(+ Many other basecallers prior to Guppy [1] and to come.)



(+ Many other basecallers prior to Guppy [1] and to come.)

Harmeet Singh @GiessenSingh ...

Contiguity comparison between Wheat @nanopore assemblies using Guppy and Bonito base calling. Looks like Bonito increases N50. #Bioinformatics #longreads Traduire le Tweet 5:25 PM · 28 avr. 2021 · Twitter Web App

16 Retweets 57 J'aime

Replies 12 Likes 12 Shares 1

Retweetez Répondre

harish @harishkt19... · 30 avr. ... En réponse à @GiessenSingh @kazumachack et @nanopore Do you have any comparisons as to how Bonito basecalled and HiFi reads behave? 1 1 1 1

Harmeet Singh @... · 30 avr. ... Not yet!! but I will have that in some time 1 2 1 1 Voir les réponses

Nick Verecke @m... · 29 avr. ... 1 1 1 1



Chris Seymour ✅ @iiSeymour · May 18

Huge dorado release 🎉

...

- 1.4x simplex perf HAC/SUP on A100
- 5khz v4.2 models
- All context 5mC, 6mA models
- Significantly faster duplex calling
- Duplex pairing handled automatically
- BAM output
- Aligned output via minimap2

#nanoporeconf

New Release v0.3.0

v0.3.0

[0.3.0] (18 May 2023)



This is a major release of Dorado which introduces: Duplex pairing and splitting for directly going from POD5 to duplex reads, major...



1

Contributor

github.com

Release v0.3.0 · nanoporetech/dorado

[0.3.0] (18 May 2023) This is a major release of Dorado which introduces: Duplex pairing and splitting for directly going from POD...

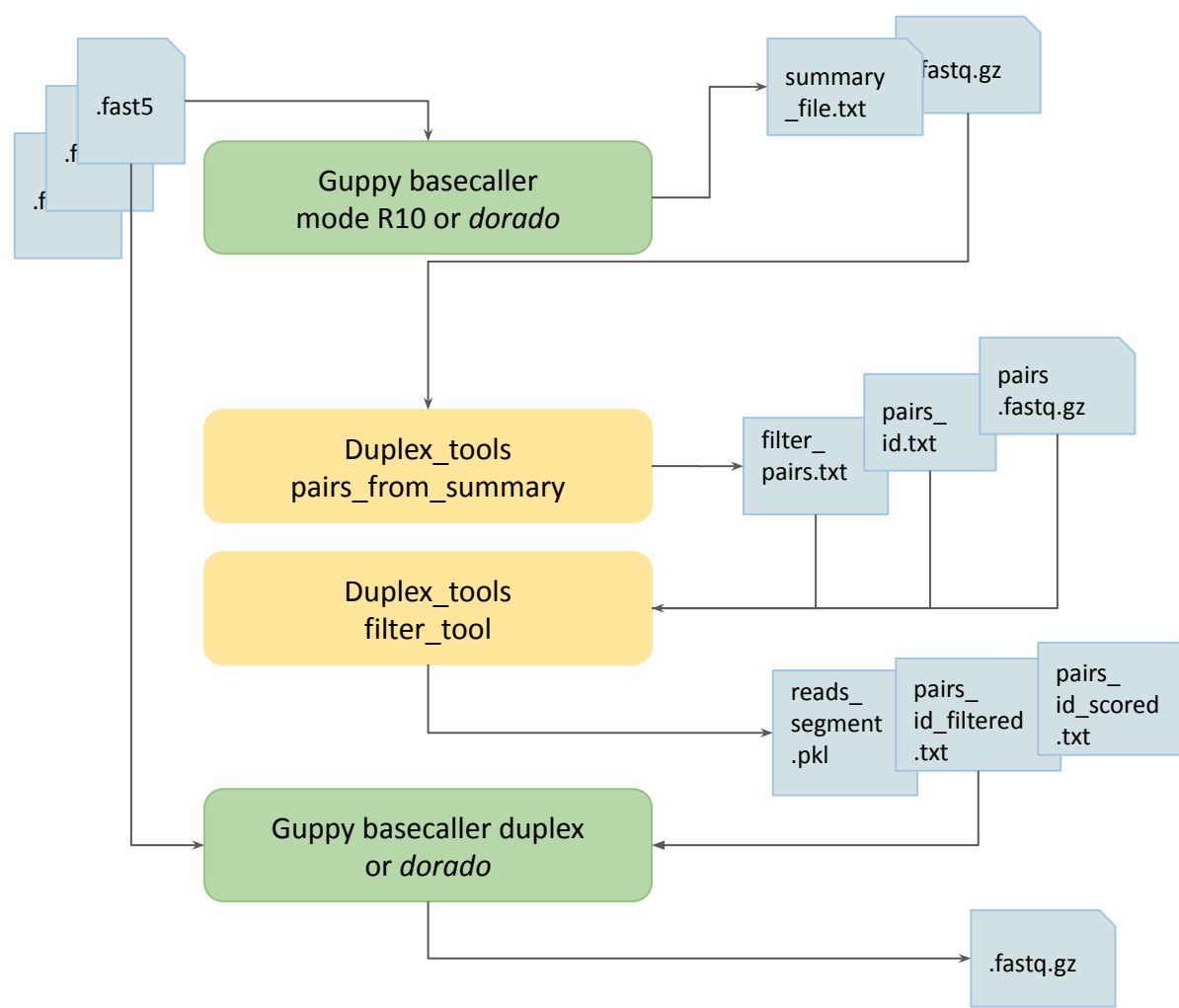
4

65

156

23.2K





<https://github.com/nanoporetech/duplex-tools>

summary_file.txt

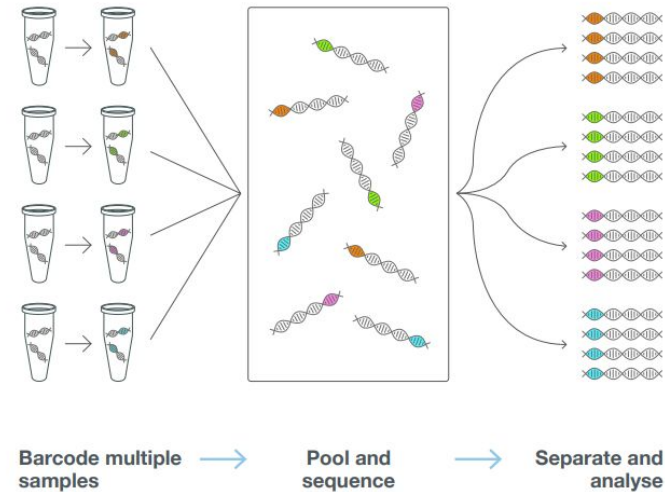
filename	FAK47038_aa36ef836fd50817477a5770772dffc63bfed2eb_30
read_id	188e2a0b-780c-440d-9223-61d8979dd002
run_id	aa36ef836fd50817477a5770772dffc63bfed2eb
batch_id	0
channel	70
mux	3
start_time	9688.985500
duration	1.610500
num_events	1288
passes_filtering	TRUE
template_start	9689.318000
num_events_template	1022
template_duration	1.278000
sequence_length_template	545
mean_qscore_template	11.462492
strand_score_template	3.165753
median_template	79.270927
mad_template	9.512511
scaling_median_template	79.270927
scaling_mad_template	9.512511

ONT demultiplexing

Deepbinner: Demultiplexing barcoded ONT reads with deep convolutional neural networks (CNN). The network is trained to classify barcodes based on the raw nanopore signal.

Guppy

In contrast to Deepbinner, guppy barcoding requires basecalling of all reads and detects barcodes in the sequence



ONT Read calling, cleaning and filtering

Sequencer ONT : raw fast5 files (or POD5)

- Transform fast5 signal in fastq standard format *Guppy Bonito/Dorado*
- Optional Demultiplexing and removing adapters *Guppy options*
- Optional Find and remove adapters from reads *Porechop*
- Optional Quality filtering using the *sequencing_summary.txt* information : *Guppy options, filtlong, nanofilt*

Guppy is a neural network based basecaller that in addition to basecalling also performs filtering of low quality reads, clipping of Oxford Nanopore adapters and estimation of methylation probabilities per base

FASTQ FORMAT

1 séquence = 4 lignes

```
@H4:C7C99ACXX:6:1101:1360:74584/2
CTGTTTCTTAGTATTTTGATGTCAATTCCGTGTTGGTTAGTTGCAAGGT
+
@@@DADFFHHFFHIIIEFEIGJGGHI4FFIEIGHI<FHGAHGGGB@3?BDB9D
@H4:C7C99ACXX:6:1101:1452:19906/2
CTGAGATCAATTGGATCCTGATGATACTGTGCTTAGCTATTACCTTGTT
+
@@@DDDD>FFFFAFBEABB4C+3?:CBB@<<A?E4A???9C@CFF*9*B3D?B
@H4:C7C99ACXX:6:1101:1476:35220/2
CATGTGCTATTACCAAAAGTCAGTAACGACCTATAAATTAAAGTAGC
+
@CFFFFFFGGHHHHJJJJIEE<HHHIJJIGBHGGEEIJJEIEIJIHHJFIIJJGHJJ
@H4:C7C99ACXX:6:1101:1491:94128/2
AGAAGTCTCGGAAAAGTTGGGTATGGCTCTAGTAGCTTTGTCTTAT
+
@C@FFFFFFGGHHDHGIIEEHIII<CGHIJJIJJ:FC9DGAFGHII?DGBFIJHBI
@H4:C7C99ACXX:6:1101:1538:34462/2
ACAAAAAAGCTAAAAGAACACAGTTGCTGAAGCAGCAAACACAAGAAC
+
B@@@DFFFFFGHHHHJIIIIJJJIIGJCHHEIII>GHIG@GHIDHGJIIFHIIJJG
@H4:C7C99ACXX:6:1101:1568:67898/2
ACAAATGGGTGTGAAGAGTTAAAAACAAATTATGAGCAACTGAGTTTC
+
@@@CFFFFFFHFFFHFGIJJIHIIJJIIHJJECGHJJCHGICDGHHJ<FGGIJJ
@H4:C7C99ACXX:6:1101:1575:18963/2
AACATGTTGTCGGGGTTGGAAATTGTCACTTCTGCTACAATGCCG
+
@<@DDDDDHFFFFDIIBDFGHGG;FGGCHHAGGGIIH@E>AEDDEECAB>
```



- @identifiant de la séquence
- Séquence
- + (id séquence).
- Qualité de la séquence = un caractère ASCII pour chaque base

Reads quality

Phred quality score: confidence score for each sequenced base
Ranging from 0 to 93 (the higher the better)

Base	T	G	A	T	A	G	T	T	A	T	G
Score	32	40	41	35	29	23	26	32	36	32	14
ASCII	A	I	J	D	>	8	;	A	E	A	/

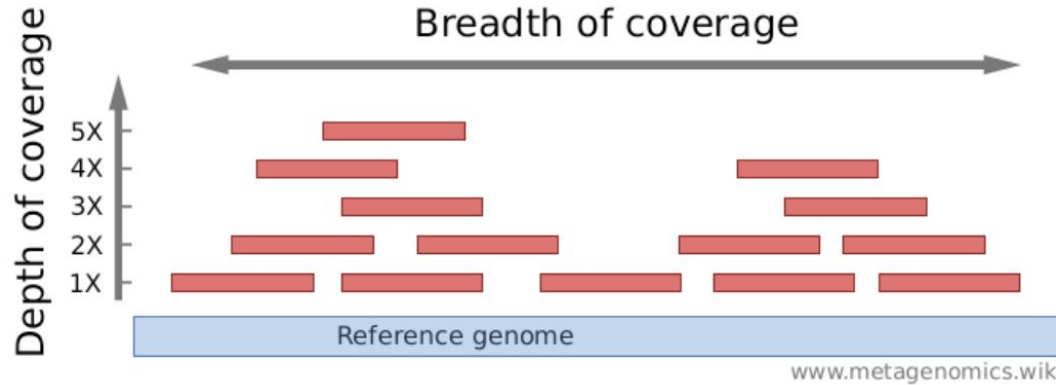
In FASTQ files scores are encoded in ASCII characters (33 to 126)

Score indicates probability P of a wrong base:

$$P = 10^{\frac{-Q}{10}}$$

Phred score of 10 \leftrightarrow 10% error rate ; score of 20 \leftrightarrow 1% error rate

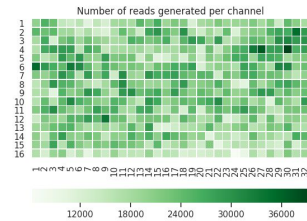
Calculate depth of coverage



depth of coverage estimation :

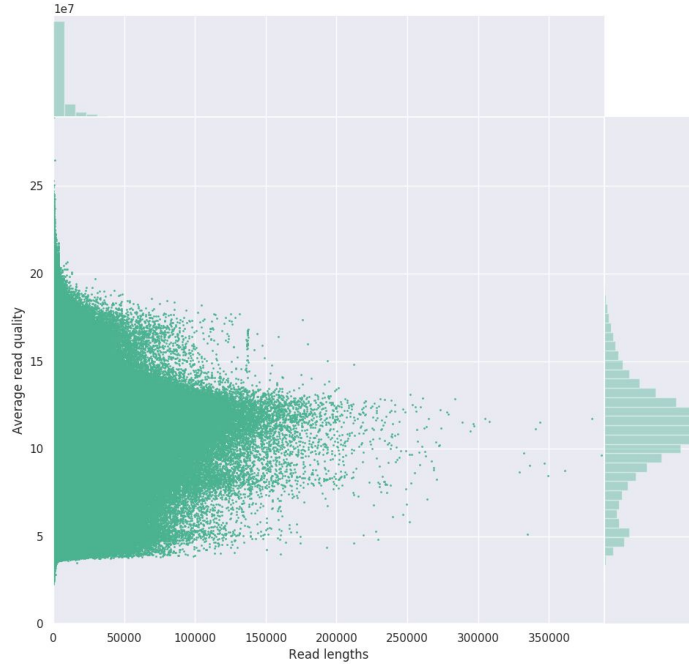
- Count how much base pairs in all sequenced reads? *total_pb*
- What is the expected genome size? *genome_size*

$\text{depth_of_coverage} = \text{total_pb}/\text{genome_size}$



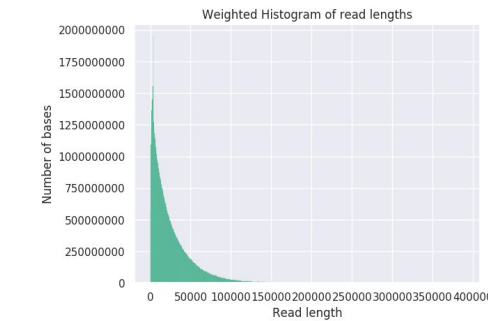
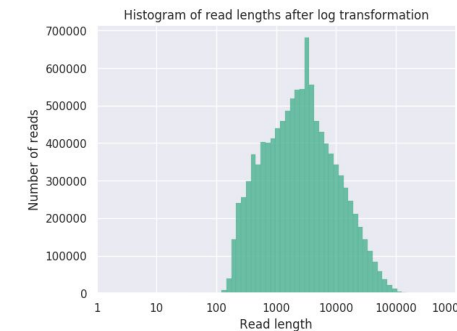
Reads Quality control : *NanoPlot*

Read lengths vs Average read quality plot

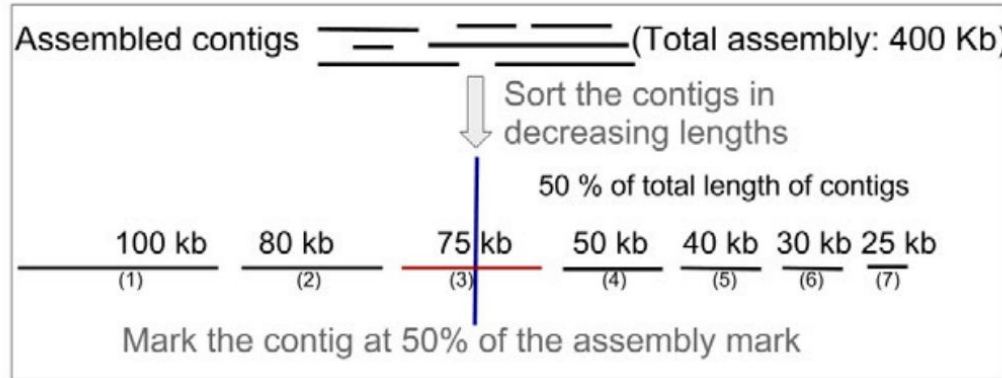


Summary statistics

General summary	
Active channels	512.0
Mean read length	6,315.6
Mean read quality	10.9
Median read length	2,517.0
Median read quality	11.1
Number of reads	10,847,854.0
Read length N50	16,816.0
Total bases	68,510,227,164.0



What is N50 and L50?



- N50, length of the contig at 50% assembly: 75 kb
→ L50, number of contigs until 50% assembly: 3

Reads Quality control

NanoPlot : <https://github.com/wdecoster/NanoPlot>

NanoComp : <https://github.com/wdecoster/nanocomp>

(mini_qc : https://github.com/roblanf/minion_qc)

Conclusion : check reads N50, reads length distribution, and calculate coverage !

TP1. Reads Quality Control

- TP1

[https://github.com/SouthGreenPlatform/training_ONT_teaching/
blob/2023_MTP/1.light_raw_quality_control.ipynb](https://github.com/SouthGreenPlatform/training_ONT_teaching/blob/2023_MTP/1.light_raw_quality_control.ipynb)

Chapitre 2

Assemblies

What do you want to do with these long reads?



Research areas

- Microbiology
- Microbiome
- Environmental
- Plant
- Animal
- Infectious disease
- Human genomics
- Clinical research
- Cancer
- Transcriptome
- Populations genomics
- COVID-19

Investigations

- Structural variation
- SNVs and phasing
- Gene expression
- Identification
- Splice variation
- Assembly
- Fusion transcripts
- Epigenetics
- Single cell
- Chromatin conformation

Techniques

- Whole genome
- Targeted
- Whole transcriptome
- Metagenomics
- Short Fragment Mode

Type	Reference	Application	
Aligners/Alignment-based classifiers			
BLAST, MEGABLAST	[58,59]	Targeted; Shotgun	
minimap2	[33]	Targeted; Shotgun	
Alignment-free classifiers			
Kraken, Kraken2	[35,64]	Targeted; Shotgun	
KrakenUniq	[65]	Shotgun	
Bracken	[66]	Targeted; Shotgun	
Metamaps	[69]	Shotgun	
Centrifuge	[34]	Targeted; Shotgun	
Mash	[72]	Targeted; Shotgun	
Long-read assemblers			
Canu	[90]	Shotgun	
miniasm	[73]	Shotgun	
wtdbg2	[91]	Shotgun	
OPERA-MS	[95]	Shotgun	
MetaFlye	[96]	Shotgun	
MetaSPAdes	[74]	Shotgun	https://doi.org/10.1016/j.csbj.2021.02.020
Sequence correction and polishing tools			
Nanopolish		https://github.com/jts/nanopolish	Targeted; Shotgun
Medaka		https://github.com/nanoporetech/medaka	Targeted; Shotgun
Metagenomic analysis pipelines			
MEGAN-LR	[60]		Shotgun
NanoCLUST	[25]		Targeted
Reticulatus		https://github.com/SamStudio8/reticulatus	Shotgun
MUFFIN	[70]		Shotgun
NanoSPC	[71]		Shotgun
BusyBee		https://ccb-microbe.cs.uni-saarland.de/busybee/	Shotgun

Type	Reference	Application
Aligners/Alignment-based classifiers		
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OPERA-MS	[95]	Shotgun
MetaFlye	[96]	Shotgun
MetaSPAdes	[74]	Shotgun



HAS BEEN !

Sequence correction and polishing tools

Nanopolish	https://github.com/jts/nanopolish	Targeted; Shotgun
Medaka	https://github.com/nanoporetech/medaka	Targeted; Shotgun

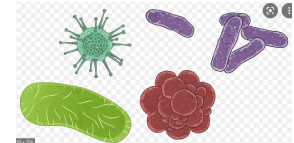
Metagenomic analysis pipelines

MEGAN-LR	[60]	Shotgun
NanoCLUST	[25]	Targeted
Reticulatus	https://github.com/SamStudio8/reticulatus	Shotgun
MUFFIN	[70]	Shotgun
NanoSPC	[71]	Shotgun
BusyBee	https://ccb-microbe.cs.uni-saarland.de/busybee/	Shotgun

Which assembler to use over my favorite organism?

Long reads simplify genome assembly, with the ability to span repeat-rich sequences (characteristic of antimicrobial resistance genes) and structural variants. Nanopore sequencing also shows a lack of bias in GC-rich regions, in contrast to other sequencing platforms. To perform microbial genome assembly, we suggest using the third-party de novo assembly tool Flye. We also recommend one round of polishing with Medaka.

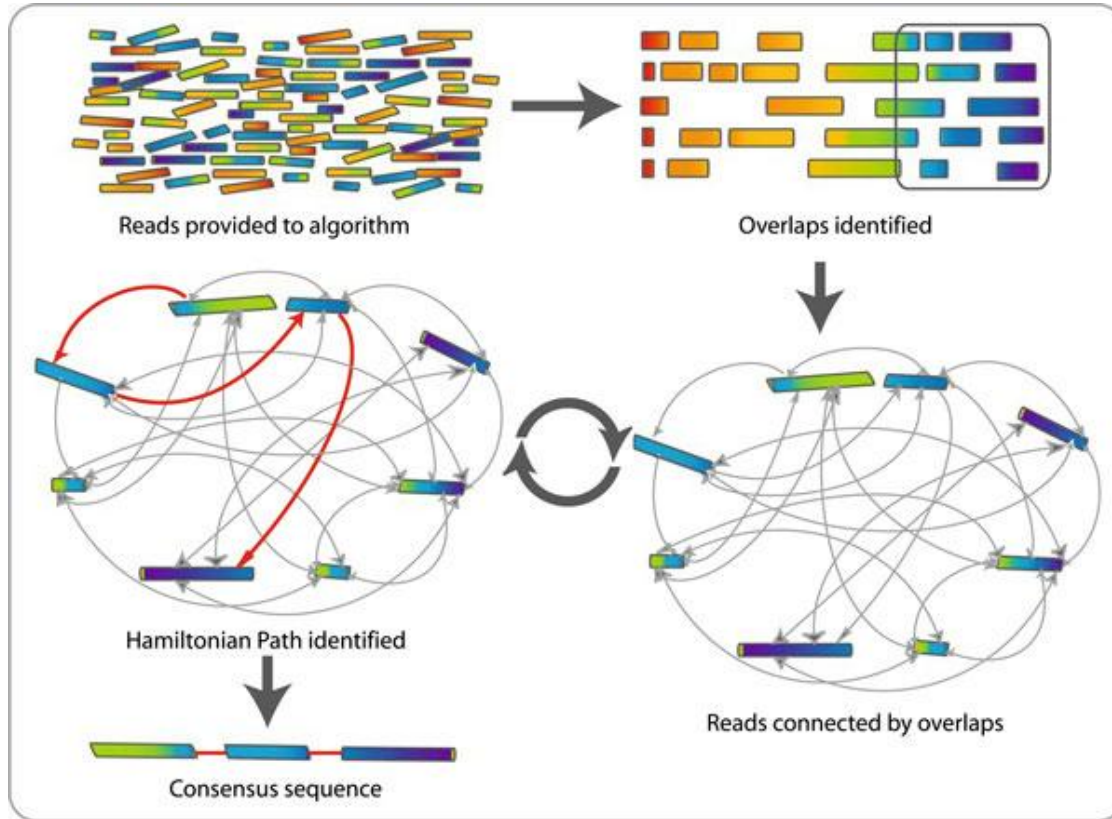
<https://nanoporetech.com/sites/default/files/s3/literature/microbial-genome-assembly-workflow.pdf>



For assembly, ONT recommend sequencing a human genome to a minimum depth of 30x of 25–35 kb reads. However, sequencing to a depth of 60x is advisable to obtain the best assembly metrics. We also recommend basecalling in high accuracy mode. Greatest contig N50 is usually obtained with Shasta and Flye. Polishing/Correction is also recommended (Racon and Medaka).

<https://nanoporetech.com/sites/default/files/s3/literature/human-genome-assembly-workflow.pdf>

Overlap–layout–consensus genome assembly algorithm (OLC)



[Canu](#), [Flye](#), [Miniasm](#), [Raven](#), [Smartdenovo](#), [Shasta](#)

<http://www.ncbi.nlm.nih.gov/pmc/articles/PMC3055744/>

Polishing / Correction

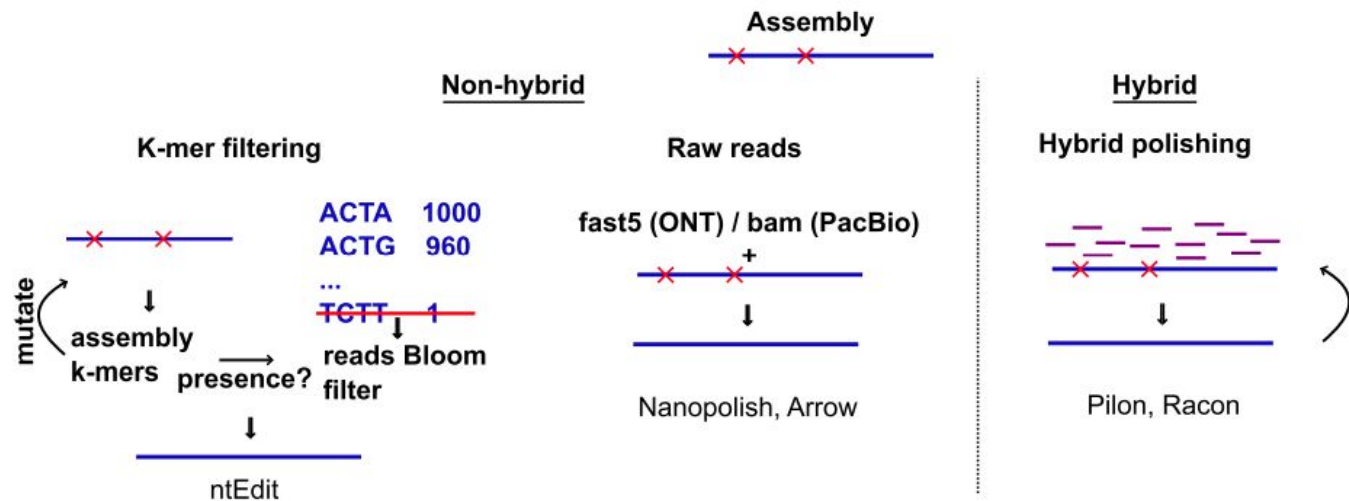
[Racon](#) correct raw contigs generated by rapid assembly methods which do not include a consensus step. It can polish with either Illumina data or data produced by third generation of sequencing. (recursive use)

[Medaka](#) and [Nanopolish](#) create a consensus sequence of nanopore sequencing data. (mapping + consensus)

- + Medaka uses neural networks where Nanopolish uses HMMs.
- + Medaka uses basecalled reads, not the raw signal.
- + Medaka propose the ability to train one's own basecalling model

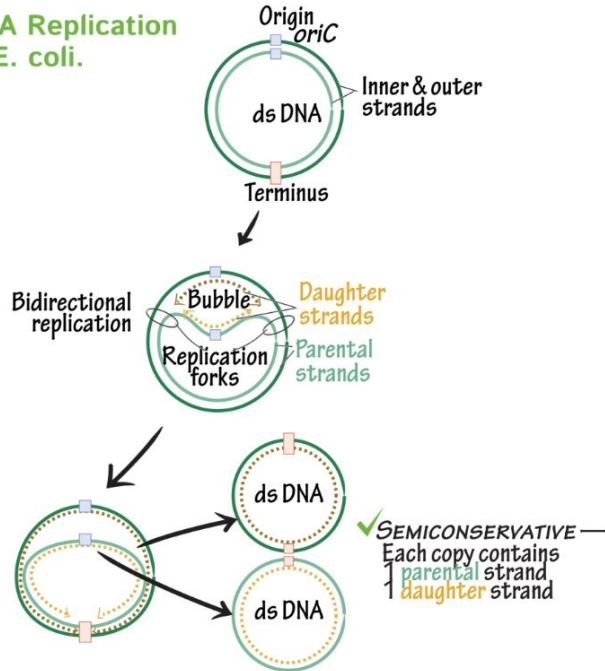
[Pilon](#) correct assemblies using illumina reads. (recursive use)

Autres : [NeuralPolish](#) , [ntEdit](#)



Circularisation ?

DNA Replication
in E. coli.



Some assemblers give you information about circularisation of assembled molecules (flye, canu).

Circularisation can be found also on GFA files generated by assemblers. (miniasm, raven, shasta)

You can try to circularise assembled molecules using tools as [circlator](#)

it could be interesting tagging and rotation of circular molecule before each polishing step.

As well as, fixing (*dnaA* gene) the start position on circular genome. This is efficient when multiple genome alignments are envisaged.

TP2. Assemblies

- TP2

[https://github.com/SouthGreenPlatform/training_ONT_teaching/
blob/2023_MTP/2.light_assemblies.ipynb](https://github.com/SouthGreenPlatform/training_ONT_teaching/blob/2023_MTP/2.light_assemblies.ipynb)

Chapitre 3

Contigs Quality Control

QUAST

Quality Assessment Tool for Genome Assemblies by [CAB](#)

26 March 2021, Friday, 07:37:40

[View in Icarus contig browser](#)

All statistics are based on contigs of size ≥ 3000 bp, unless otherwise noted (e.g., "# contigs (≥ 0 bp)" and "Total length (≥ 0 bp)" include all contigs).

Aligned to "TIGRv7_ok" | 375 096 285 bp | 16 fragments | 43.57 % G+C

Worst	Median	Best	<input type="checkbox"/> Show heatmap			
Genome statistics						
Genome fraction (%)	65.801	65.916	65.417			
Duplication ratio	1.036	1.041	1.041			
Largest alignment	2 503 013	2 501 477	1 739 590			
Total aligned length	255 403 246	257 194 821	255 339 839			
NGA50	48 559	48 062	42 714			
LGA50	1338	1333	1404			
Misassemblies						
# misassemblies	9633	9923	7666			
Misassembled contigs length	373 371 138	373 825 172	335 007 830			
Mismatches						
# mismatches per 100 kbp	2776.55	2831.25	2669.89			
# indels per 100 kbp	321.69	301.83	330.99			
# N's per 100 kbp	0	0.23	0			
Statistics without reference						
# contigs	181	250	250			
Largest contig	43 938 576	43 971 118	14 121 367			
Total length	383 158 522	384 147 370	387 291 200			
Total length (≥ 1000 bp)	383 173 133	384 197 574	387 291 200			
Total length (≥ 10000 bp)	382 901 616	383 618 037	387 291 200			
Total length (≥ 50000 bp)	381 421 486	381 880 053	387 291 200			
250	13 998 410	383 785 534	369 892 751	369 966 935	368 865 072	365 953 108
729	6 500 937	383 785 534	369 892 751	369 966 935	368 865 072	365 953 108
854	6 543 040	383 785 534	369 892 751	369 966 935	368 865 072	365 953 108
373 136 825	373 406 571	371 578 702	368 382 574			

[Extended report](#)

plus petit nb de contigs : flye+racon puis raven+racon
plus long contigs : flye+racon

<https://github.com/ablab/quast>

Genome statistics	FLYE_STEP_POLISHING_RACon	FLYE_STEP_ASSEMBLY	RAVEN_STEP_POLISHING_RACon	RAVEN_STEP_ASSEMBLY	SHASTA_STEP_POLISHING_RACon	SHASTA_STEP_ASSEMBLY
Statistics without reference						
# contigs	181	250	250	250	729	854
# contigs (>= 0 bp)	194	285	250	250	767	1149
# contigs (>= 1000 bp)	188	274	250	250	763	1000
# contigs (>= 5000 bp)	168	207	250	250	674	746
# contigs (>= 10000 bp)	139	156	250	250	564	587
# contigs (>= 25000 bp)	97	99	250	250	487	488
# contigs (>= 50000 bp)	74	75	250	250	444	445
Largest contig	43 938 576	43 971 118	14 121 367	13 998 410	6 500 937	6 543 040
Total length	383 158 522	384 147 370	387 291 200	383 785 534	369 892 751	373 136 825
Total length (>= 0 bp)	383 176 103	384 204 105	387 291 200	383 785 534	369 969 110	373 471 297
Total length (>= 1000 bp)	383 173 133	384 197 574	387 291 200	383 785 534	369 966 935	373 406 571
Total length (>= 5000 bp)	383 108 497	383 977 711	387 291 200	383 785 534	369 668 739	372 705 755
Total length (>= 10000 bp)	382 901 616	383 618 037	387 291 200	383 785 534	368 865 072	371 578 702
Total length (>= 25000 bp)	382 215 424	382 691 571	387 291 200	383 785 534	367 717 125	370 136 458
Total length (>= 50000 bp)	381 421 486	381 880 053	387 291 200	383 785 534	365 953 108	368 382 574
N50	14 538 350	14 555 248	3 455 235	3 425 125	1 355 467	1 360 886
N75	10 163 758	10 173 888	1 497 559	1 483 567	738 018	741 772
L50	10	10	28	28	79	80
L75	17	17	68	68	173	174
GC (%)	43.56	43.61	43.59	42.81	43.43	43.36
Similarity statistics						
# similar correct contigs	260	247	263	0	255	60
# similar misassembled blocks	1251	1178	1257	0	1245	499

less contigs : flye+racon puis raven+racon

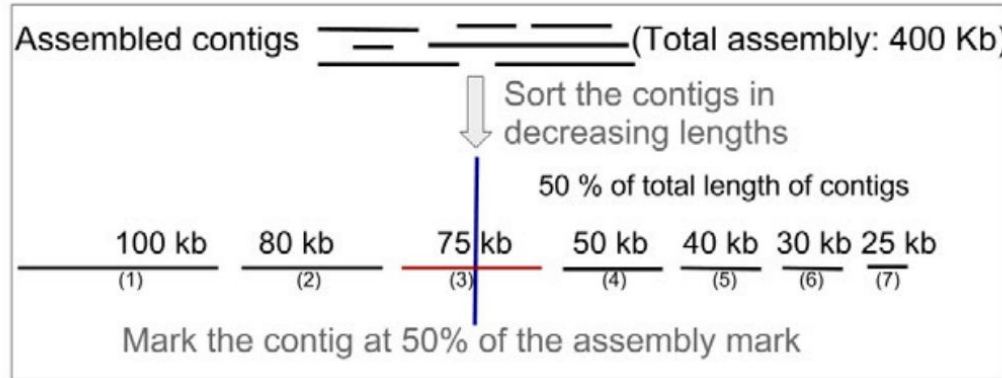
largest contig : flye+racon

largest N50 : flye

largest L50 : flye

what is N50 and L50?

What is N50 and L50?



- N50, length of the contig at 50% assembly: 75 kb
→ L50, number of contigs until 50% assembly: 3

QUAST

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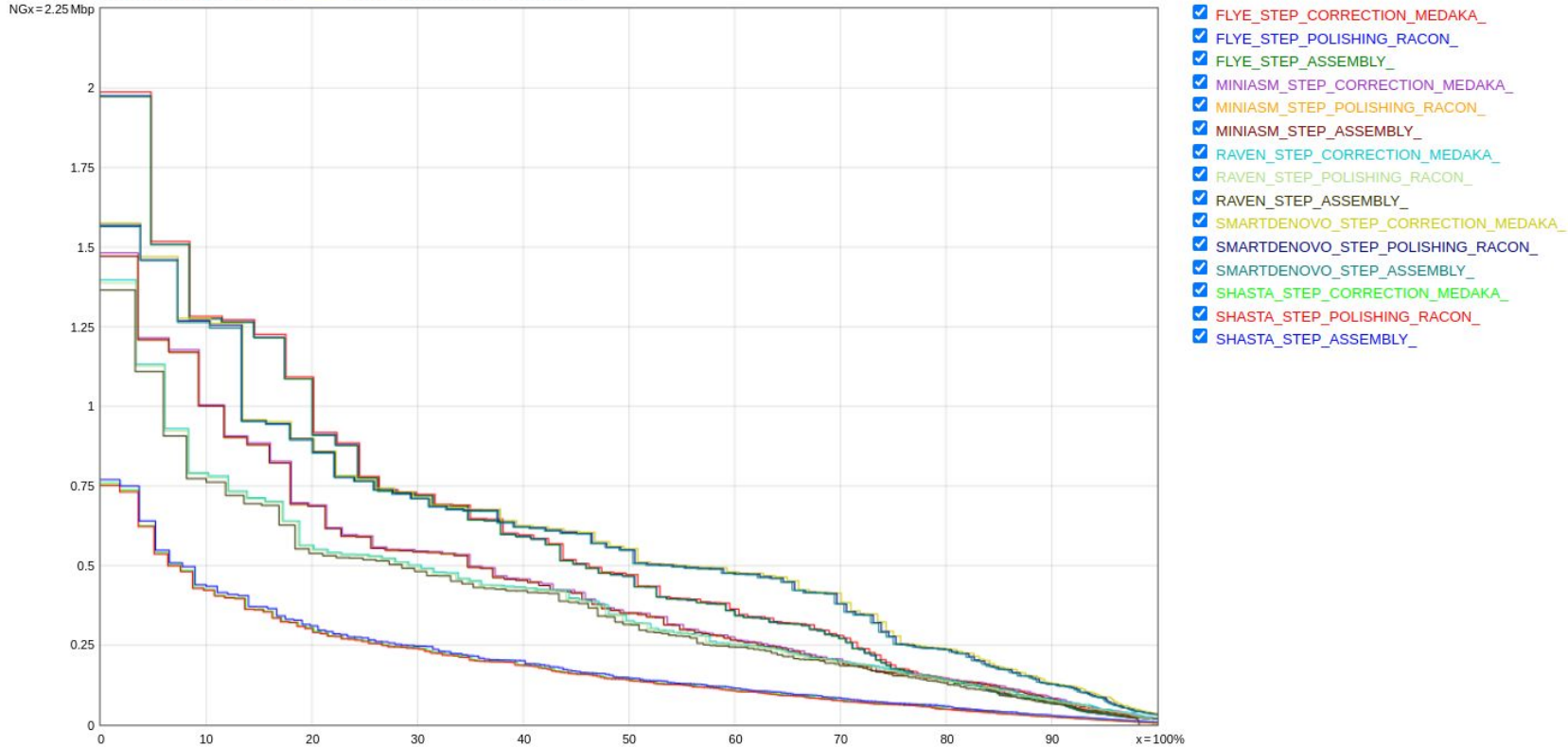
[Extended report](#)

Check misassemblies and N percentage.

BE CAREFUL! A misassembly for QUAST can be a structural variation!

Nx graph

Plots: Cumulative length Nx NAx NGx NGAx Misassemblies GC content



The greater the area under the curve AUC, the better is the assembly.
Nx represent N50 but also N10 to N100

BUSCO

from QC to gene prediction and phylogenomics

BUSCO v5.2.2 is the current stable version!

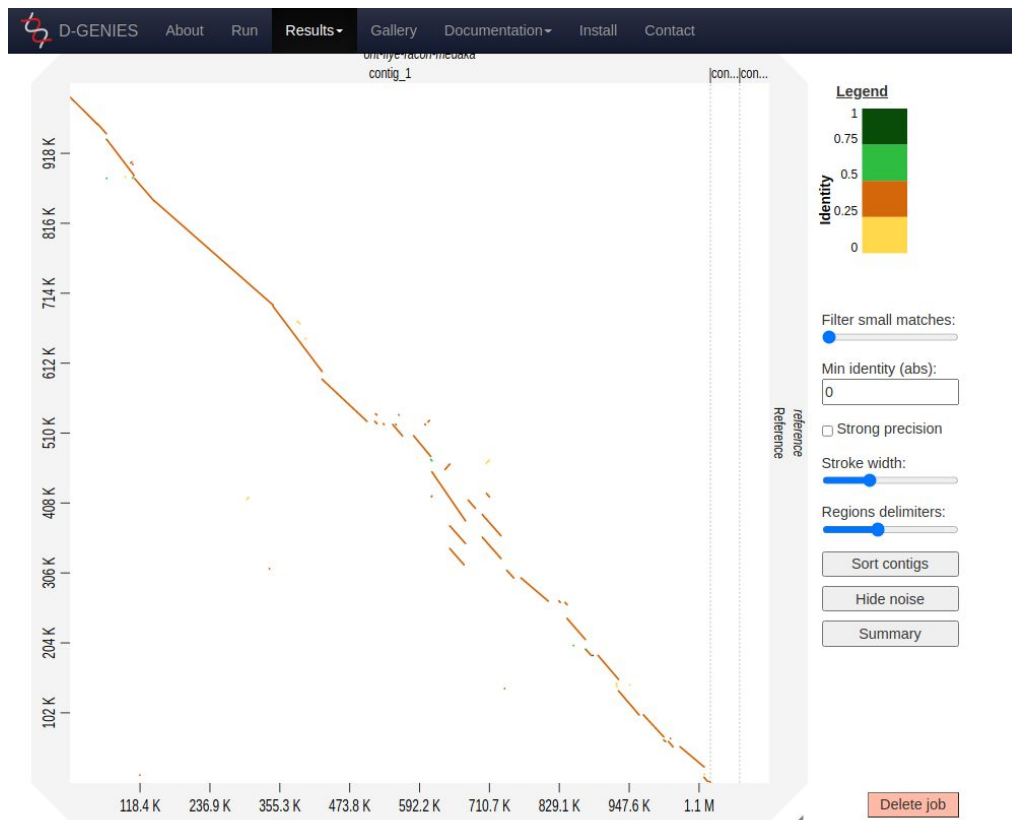
Gitlab [🔗](#), a Conda package [🔗](#) and Docker container [🔗](#) are also available.

Based on evolutionarily-informed expectations of gene content of near-universal single-copy orthologs, BUSCO metric is complementary to technical metrics like N50.

Helps to check if you have a good assembly, by searching the expected single-copy lineage-conserved orthologs in any newly-sequenced genome from an appropriate phylogenetic clade.

```
INFO Results:  
INFO C:95.6%[S:73.6%,D:22.0%],F:1.4%,M:3.0%,n:1759  
INFO 1682 Complete BUSCOs (C)  
INFO 1295 Complete and single-copy BUSCOs (S)  
INFO 387 Complete and duplicated BUSCOs (D)  
INFO 25 Fragmented BUSCOs (F)  
INFO 52 Missing BUSCOs (M)  
INFO 1759 Total BUSCO groups searched  
INFO BUSCO analysis done. Total running time: 621.2351775169373 seconds  
INFO Results written in /tmp/orjuela/BUSCO/run_trinity_busco/
```

Comparison with a reference genome



- NUCMER : Aligns a set of draft sequence contigs to a finished sequence
<http://mummer.sourceforge.net/>
- D-Genies : Online tool to compare two genomes by dot plot method
<http://dgenies.toulouse.inra.fr/>
- autre: *Gepard*

CANU

FLYE

MINIASM

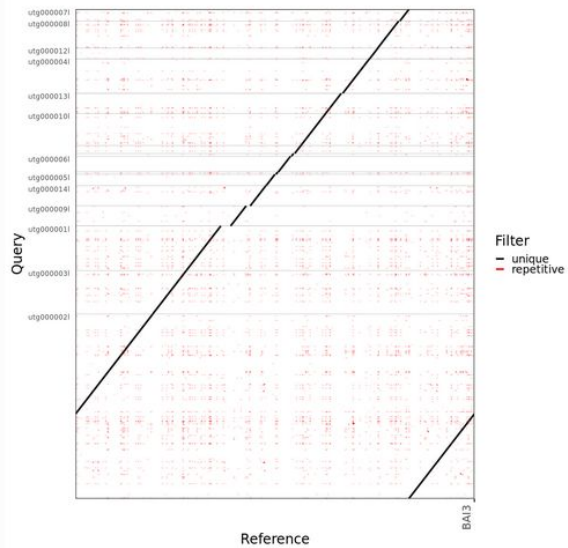
RAVEN

SMARTDENOVO

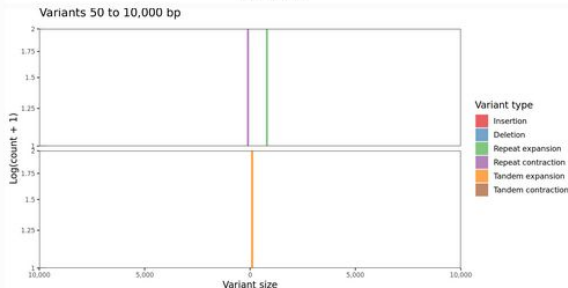
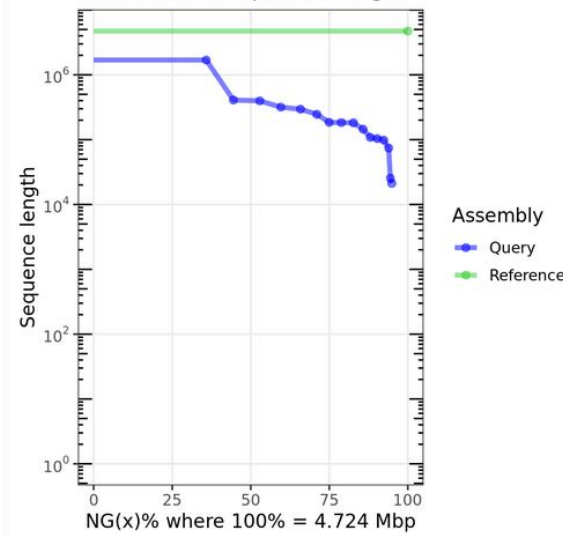
SHASTA

STEP_CORRECTION_NANOPOLISH_STARTFIXED

Dot plot of Assemblytics filtered alignments



Cumulative sequence length



TP3. Contigs Quality

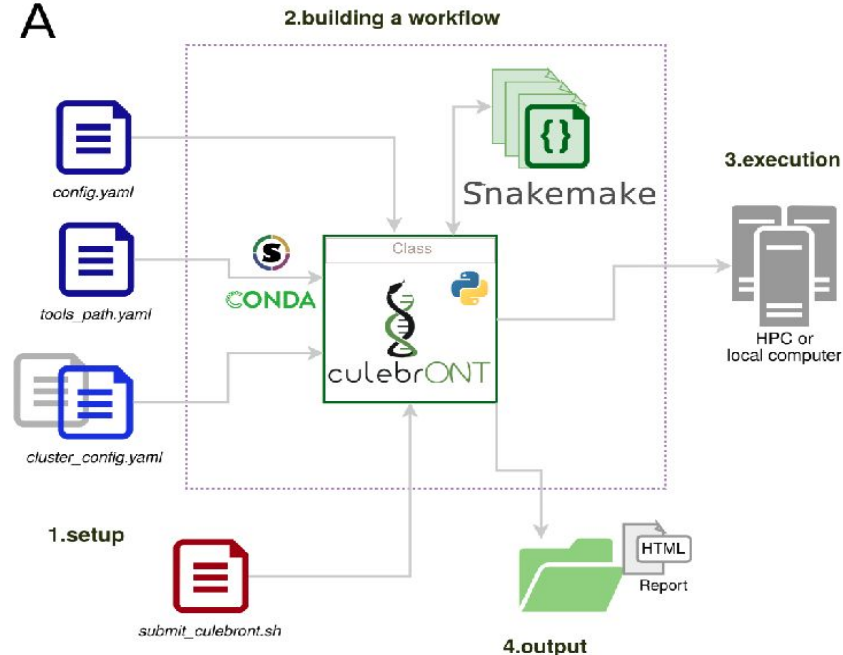
- TP3

[https://github.com/SouthGreenPlatform/training_ONT_teaching/
blob/2023_MTP/3.light_contigs_quality.ipynb](https://github.com/SouthGreenPlatform/training_ONT_teaching/blob/2023_MTP/3.light_contigs_quality.ipynb)

A flexible and reproducible pipeline for LR assembly and evaluation

pip install culebrONT

A



- A recommendation in PCI Genomics <https://genomics.peercommunityin.org/articles/rec?id=158>
- An article in PCJ <DOI:10.24072/pcjournal.153>

From contigs to chromosomes

Optical mapping : fluorescent marking of restriction sites of very long DNA molecules (up to Mb) to extract signature used to bridge contigs having these signatures.

10x chromium : shallow tagged sequencing of very long DNA fragments with Illumina machines. Read alignments enable scaffolding.

Genetic map : marker assisted contig bridging

HiC : chromosomal interaction sequencing gives the contig order on the chromosomes.

Conclusions

- DNA quality (fragment length) has a direct impact on read length
- We can assemble small to large genomes with Nanopore reads.
- Test a lot of tools to perform assemblies, ~~in any case now~~ polishing is **not** mandatory.
- There are still genomes very difficult to assemble

Formateurs

Julie Orjuela



François Sabot



Gautier Sarah (appui)



Sébastien Ravel (appui)



Special thanks to Louis Denu for the data he provided



**Merci de prendre 5 min pour remplir
l'enquête**

<https://itrop-survey.ird.fr/index.php/515725?lang=fr>

Merci pour votre attention !



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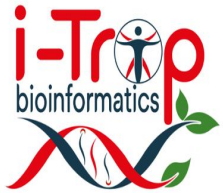
<http://creativecommons.org/licenses/by-nc-sa/4.0/>



SUIVEZ NOUS SUR TWITTER !



South Green : [@green_bioinfo](https://twitter.com/green_bioinfo)



i-Trop : [@ItropBioinfo](https://twitter.com/ItropBioinfo)



N'oubliez pas de nous citer !

Comment citer les clusters?

"The authors acknowledge the IRD i-Trop HPC at IRD Montpellier for providing HPC resources that have contributed to the research results reported within this paper. URL: <http://bioinfo.ird.fr/>"

"The authors acknowledge the CIRAD UMR-AGAP HPC (South Green Platform) at CIRAD montpellier for providing HPC resources that have contributed to the research results reported within this paper. URL:

<http://www.southgreen.fr>"



Merci pour votre attention !



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