

Session 2.1 – Secuenciación de genomas bacterianos: Aplicaciones

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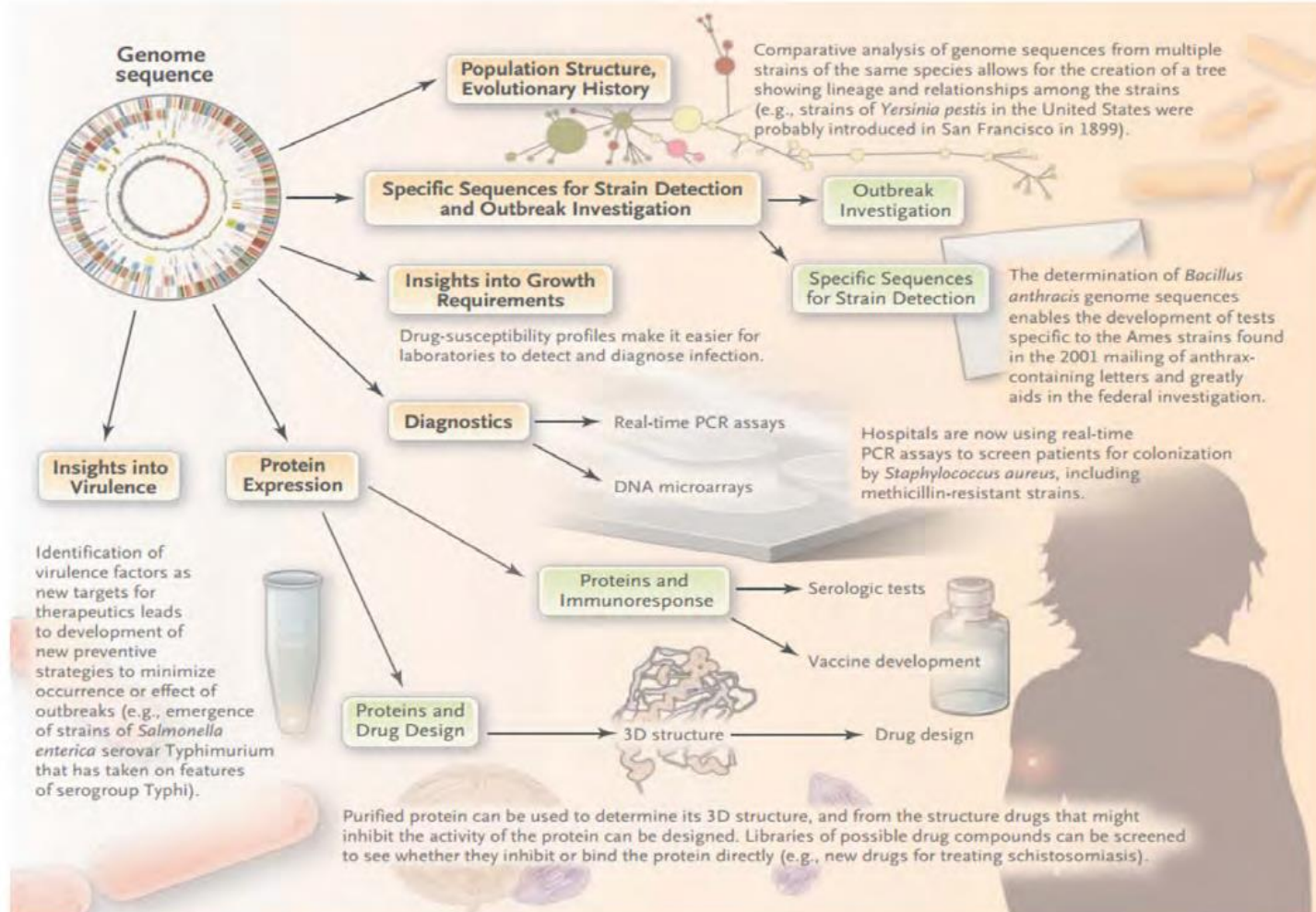
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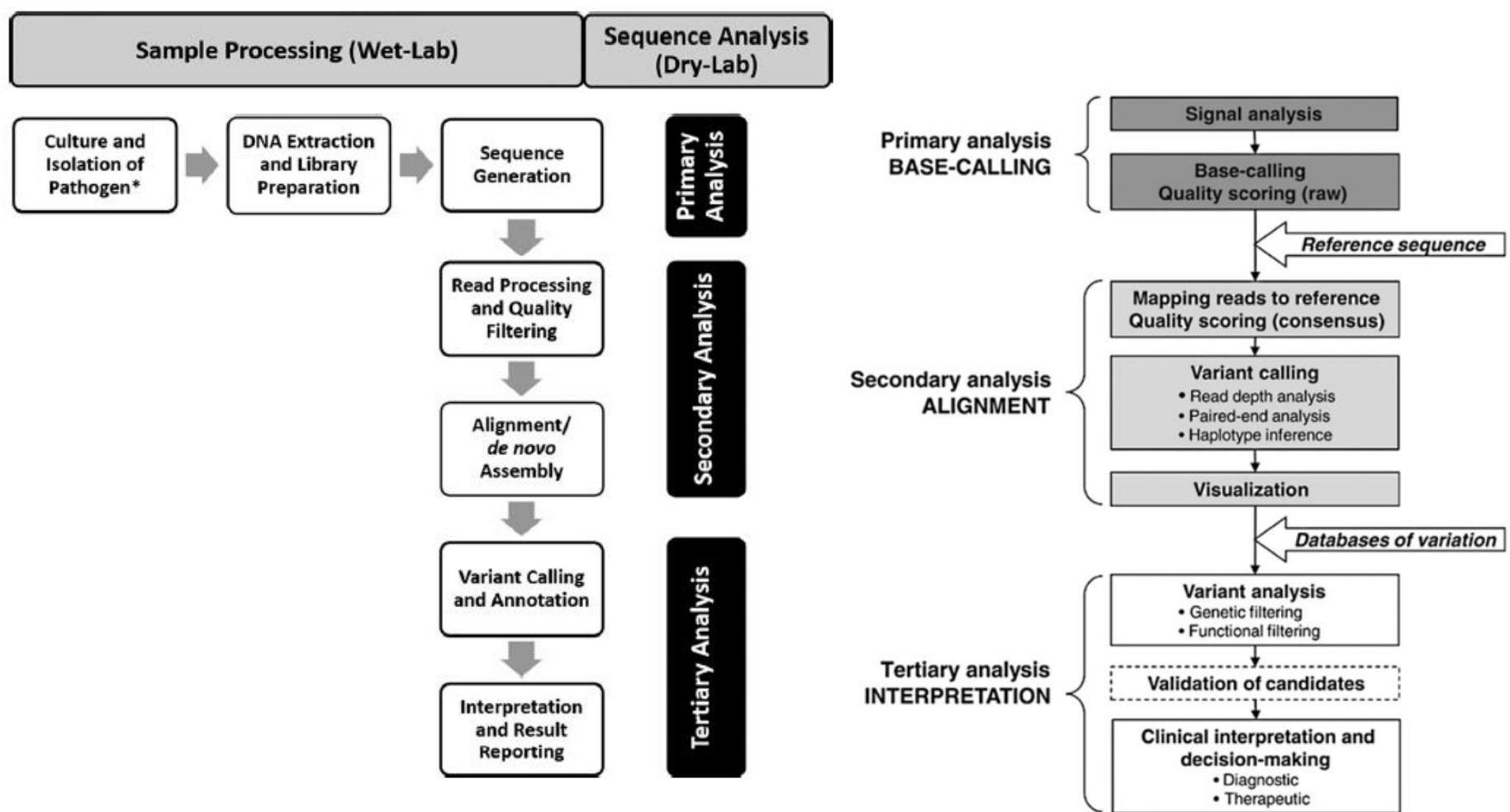
- Information obtained from WGS
- Microbial bioinformatics analysis
- Clinical use of WGS
- Considerations for implementing WGS in the laboratory routine

Use of microbial genomics for tool development

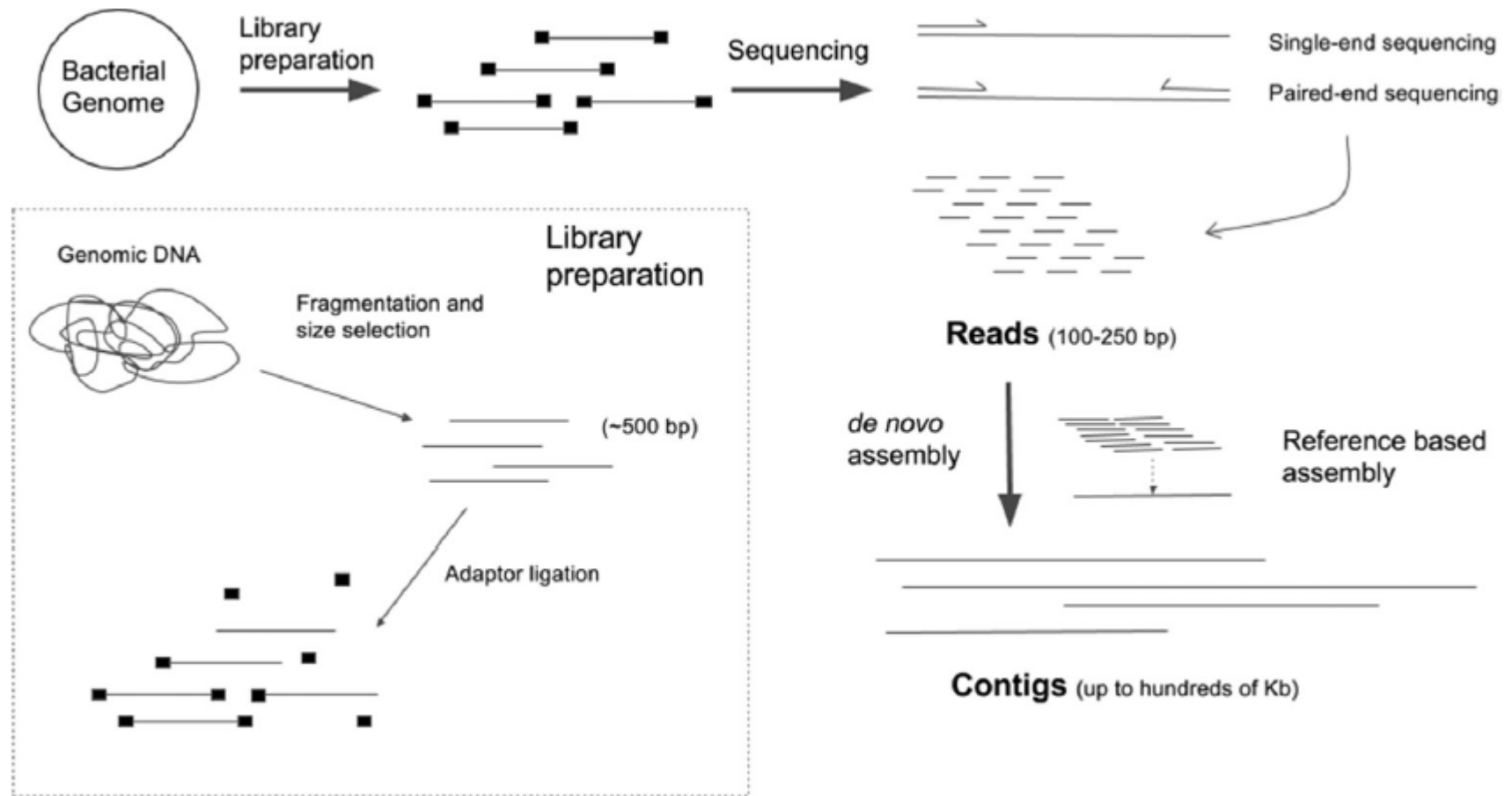
Report from *The American Academy of Microbiology*, 2015



General NGS workflow



Common microbial bioinformatics analyses



Draft genome analysis

Carrico et al., CMI 2018

- Genome annotation is the process of identifying the location and biological role of genetic features present in a DNA sequence (e.g., CDS, tRNA, rRNA, operons, CRISPR elements and genomic islands).

Software pipelines that use multiple external feature prediction algorithms

NCBI prokaryotic genome annotation pipeline (PGAP)

RAST server

PROKKA (local)

- Sequence-based microbial typing information (MLST, serotype, antimicrobial resistance or virulence genes)
- Gene-by-gene approaches: core genome = cgMLST, core + accessory genome = wgMLST
PubMLST (<https://pubmlst.org>) hosts schemas
Pasteur Institute (<http://bigsdb.pasteur.fr>) Listeria
Enterobase (<https://enterobase.warwick.ac.uk>), Salmonella, Escherichia/Shigella, Yersinia.
Pipelines: Genomic Profiler, ChewBBACA, TARANIS.
Roary: presence or absence without requiring a predefined schema
Neptune: identify differential abundance of genomic regions without requiring genome annotation information

Read mapping approaches

Carrico et al., CMI 2018

- Phylogenetic relation based on SNVs identification: if multiple strains are mapped against a single reference genome, the common variants can be used to produce a phylogenetic tree.

Software: CFSAN SNP Pipeline, Snippy, Lyve-Set, SNVPhyl, WGSOutbraker

Challenges

- SNVs in recombinant regions or in multiple-copy regions of the genome may mask the true phylogenetic signal.
- Regions with high SNV density have a high likelihood of having a recombinant origin.

Read mapping approaches

- Reads to type: using a set of target genes or genomic regions instead of using a complete or draft genome as reference for mapping.

- MLST, antimicrobial resistance genes, virulence genes

Software: SRST2, ReMatCh

Advantages over gen-by-gene approach

The ability to analyse noncoding regions, which has the potential to reveal changes in gene regulatory regions, that can be translated in phenotypic differences → limited by the reference genome used

Challenges

- Phenotype inference from genotypic data needs to be validated on a case-by-case basis

Visualization software

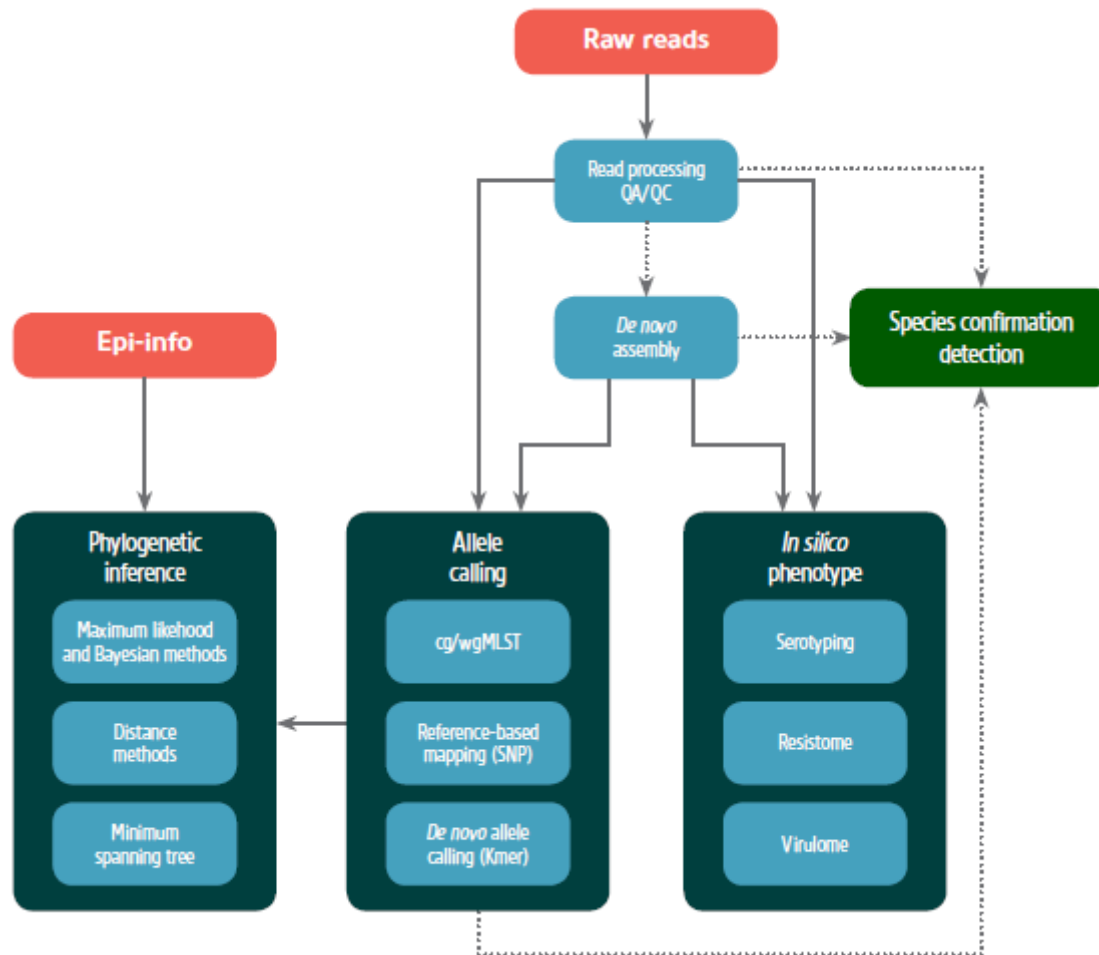
Epidemiologic and genomic data integration

- Microreact: web service where phylogenetic trees and associated geographic, genetic and epidemiologic data can be uploaded, visualized and dynamically explored, which promotes the sharing of large data sets in the platform.
- GenGIS 2: geospatial data analysis that can use the trees and epidemiologic data provided by the user.

Allelic profiles derived from gene-by-gene methods or SNV analysis

- PHYLOVIZ 2.0: representation of minimum spanning trees or hierarchical clustering together with associated metadata.
- PHYLOVIZ online: web browser, allows data sharing and visualization of large trees and metadata

Schematic representation of WGS pipeline

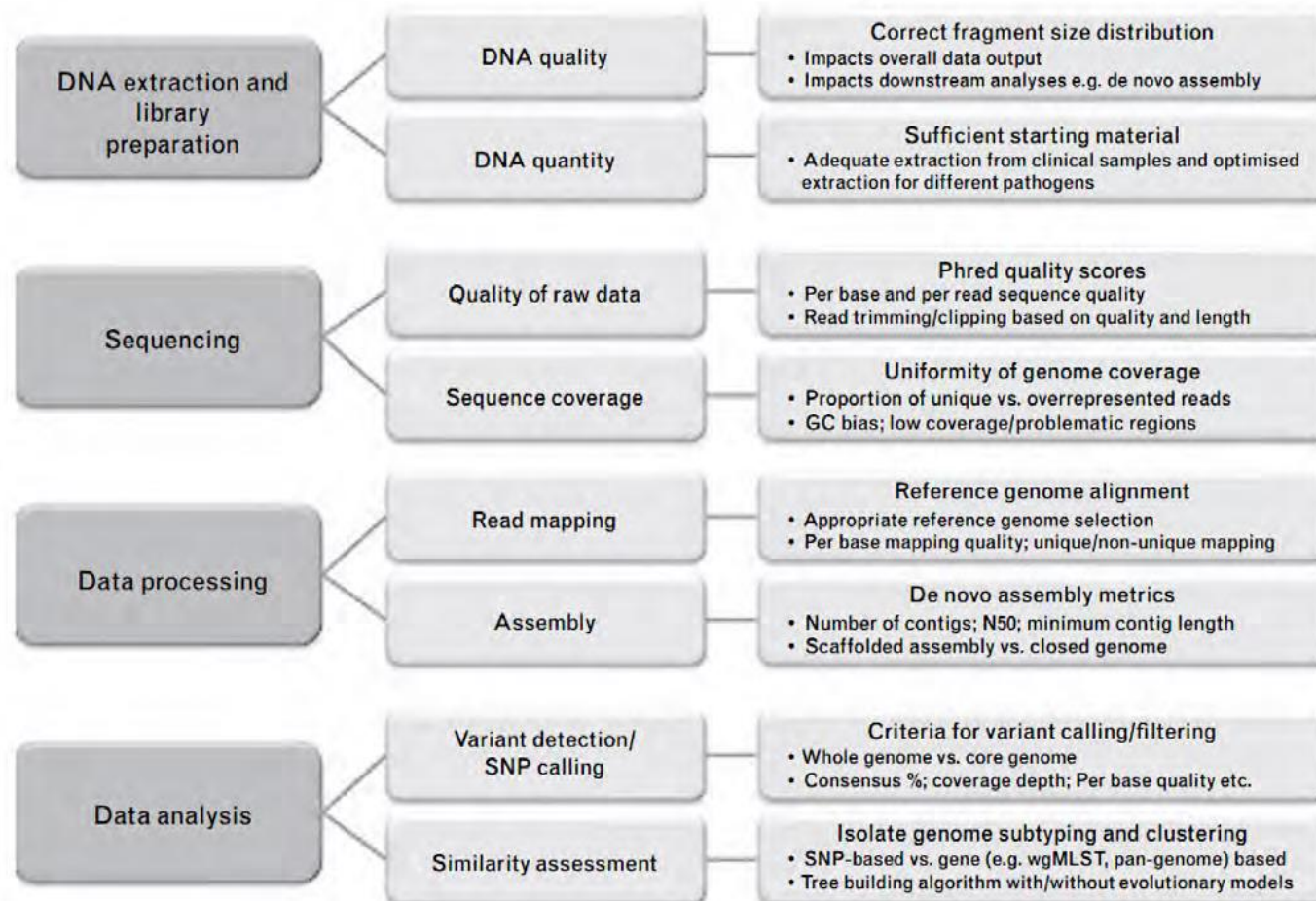


Software available for NGS, pros & cons

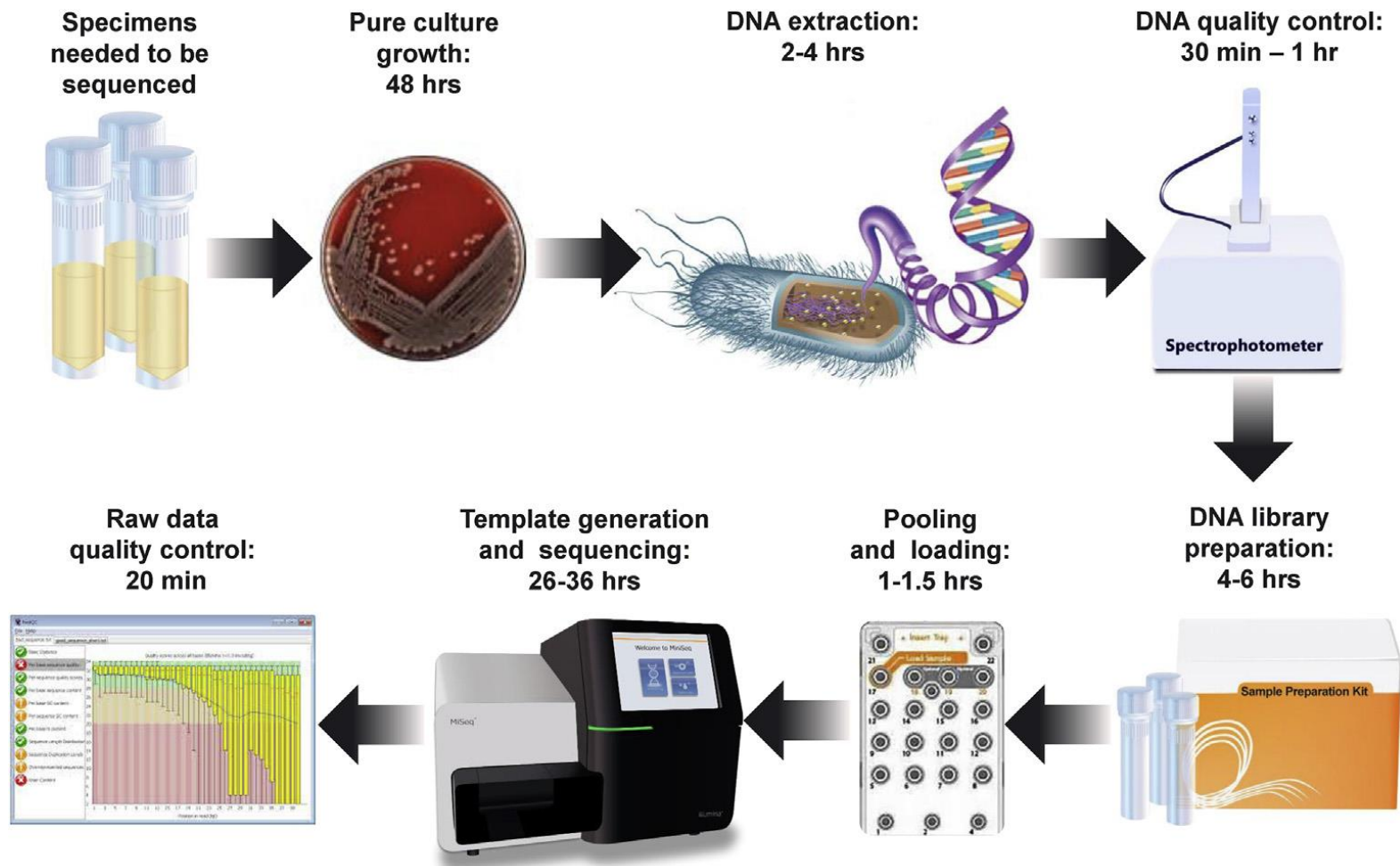
	User access	OS	Large-scale	Data-adaptable	licencie	e.g.
Web Service	Web	Browser	Limited computation	Limited or not-configurable	free	Galaxy, Different webs
Commercial	Graphic interface	Windows	Not HPC	Limited or not-configurable	\$ / €	Geneious, Bionumerics, Seqsphere, Lasergene
Open-source	Comand line	Linux	HPC	Configurable	Open code \$/ € ??	Most of them

Quality assessment considerations for WGS analyses

Applications of Clinical Microbial NGS, AASM, 2016



Typical whole genome sequencing workflow in a clinical or public health laboratory



Besser et al., Clin Microb Infect, 2018

Secuenciación de genomas bacterianos:
herramientas y aplicaciones

Applications (Clinical) of microbial genomics

1. Those requiring bacterial isolate

1. Species identification

(MALDI-TOF MS, rapid, but fails to identify unusual species)

2. Bacterial typing (WGS vs PFGE, MLVA, MLST)

2.1. Transmission pathways

2.2. Outbreak monitoring

i.e. E coli O104:H4, WGS is key to understanding the determinants and modelling the evolutionary events that may lead to a hypervirulent strain.

3. Determination of virulence factors, resistance genes

2. Those applied directly on the sample

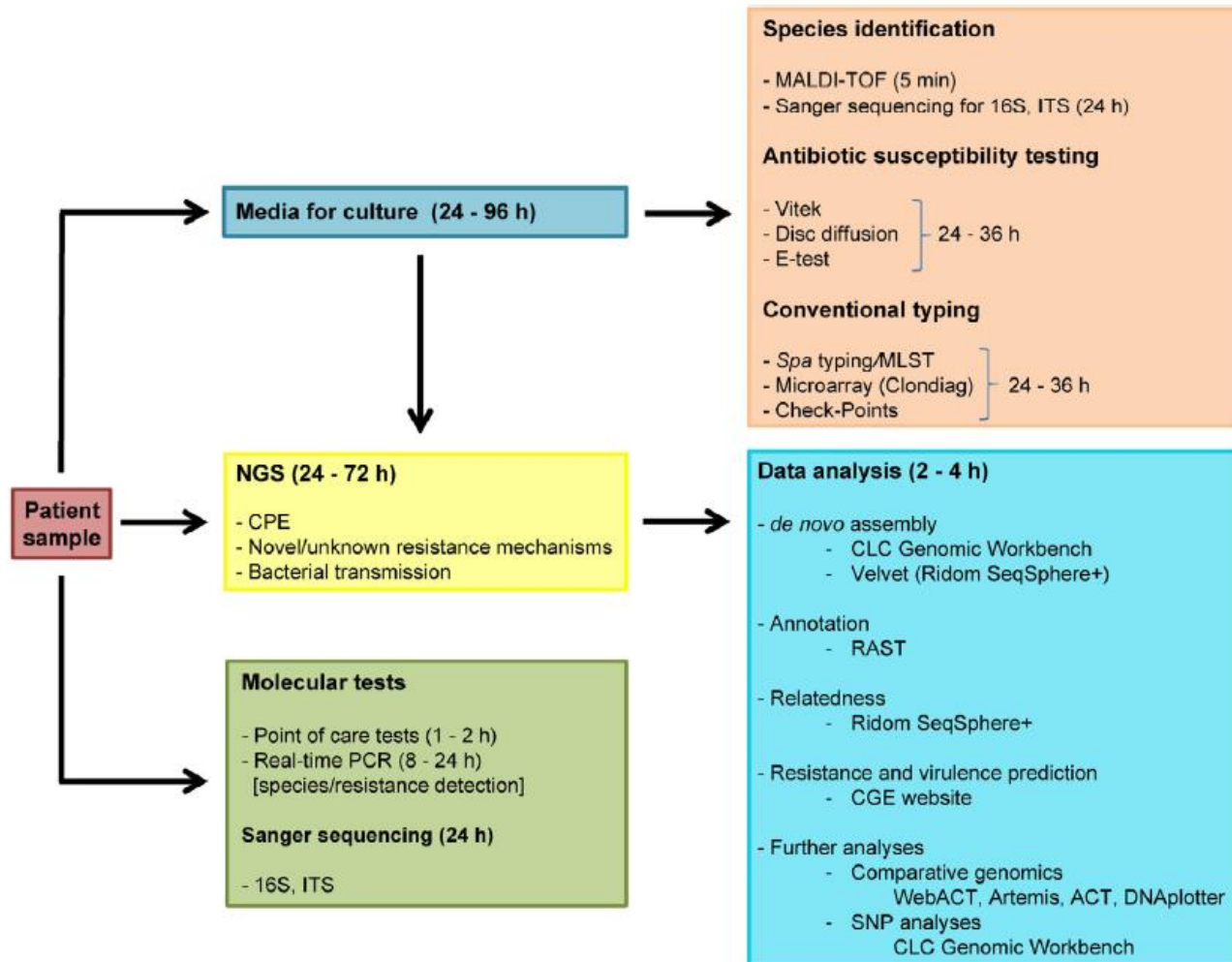
1. Metagenomics

2. Community profiling

For direct detection of known or unknown disease-associated pathogens in clinical specimens; for investigation of microbial population diversity

A schematic overview of the general workflow of diagnostic procedures including NGS in a hospital from Netherlands

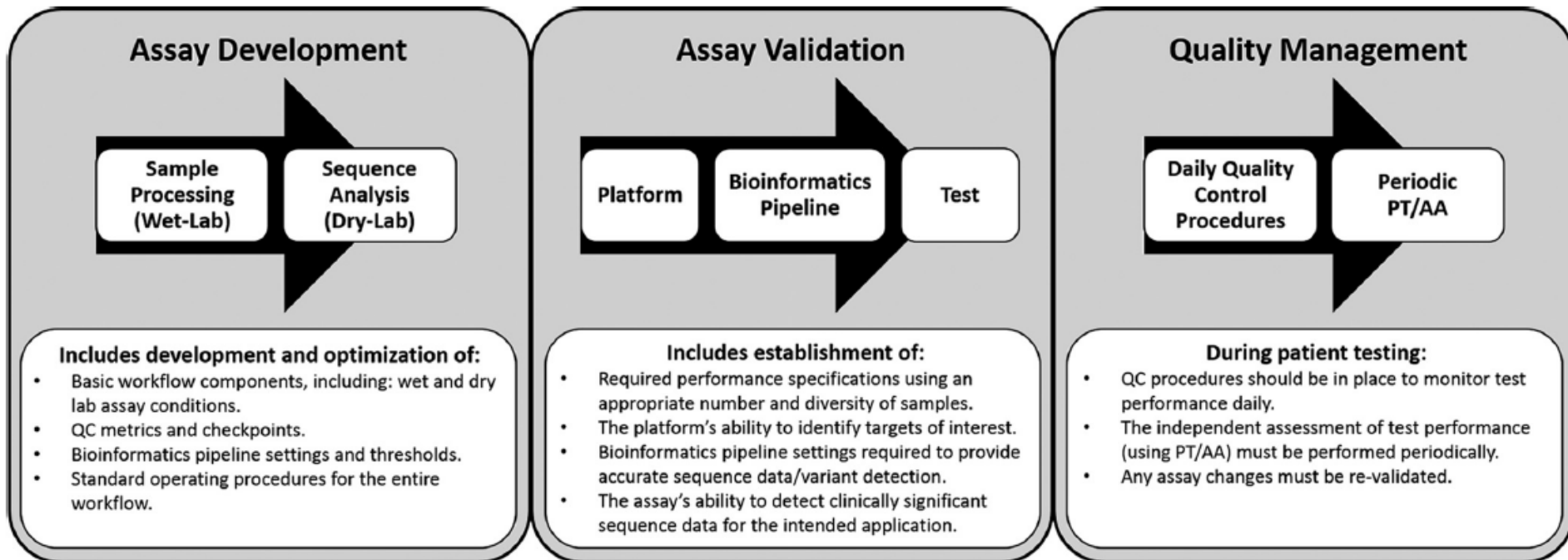
Deurenberg et al., Jbiotech, 2017



Proposed frameworks for clinical NGS implementation

Gargis et al., JCM 2016

Transition of NGS from research to the clinical and public health laboratory setting



No clinical microbiology NGS tests have been approved by the FDA, and the limited number of clinical infectious disease NGS-based assays currently offered are being performed as LDTs

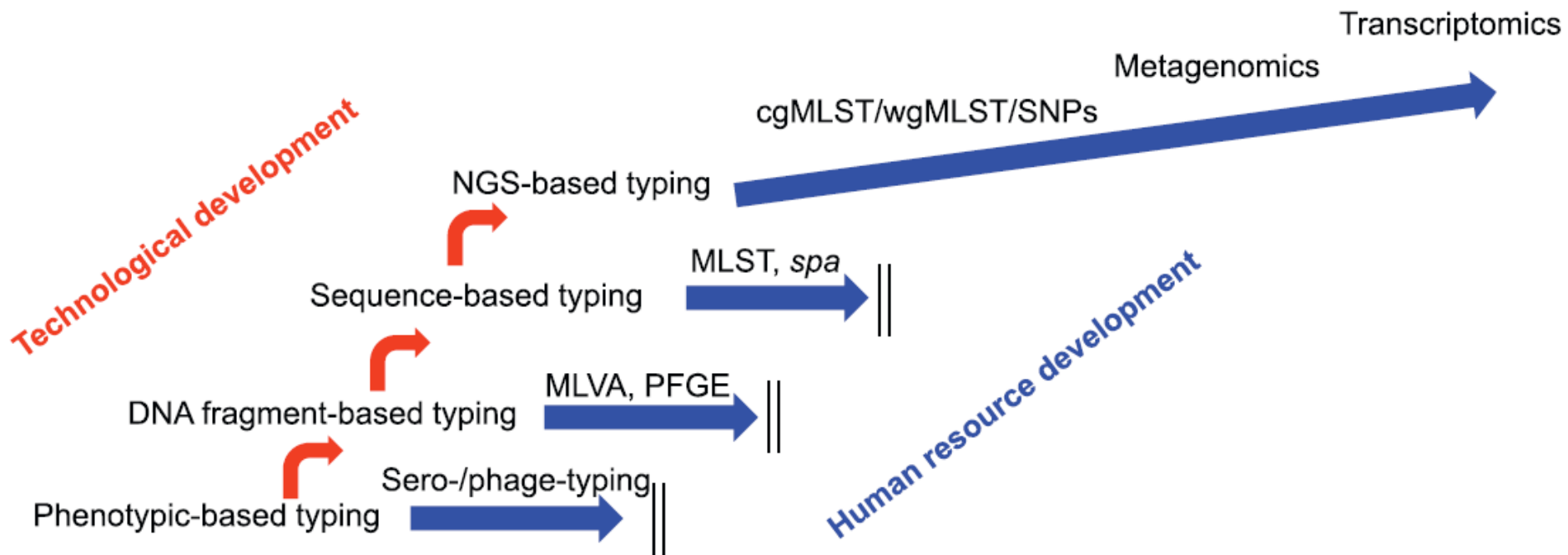
Practical issues in implementing wgs in routine diagnostic microbiology

- When and how to integrate NGS in the routine workflow?
- The place of NGS in the diagnostic hierarchy of microbiology
- Quality control issues for using NGS in microbiology
- Proficiency testing for WGS in microbiology
- Maintaining backwards compatibility with WGS

Rossen et al., CMI 2018

When and how to integrate NGS in the routine workflow?

Integrating interventional genomics in clinical microbiology settings requires development of NGS technology going hand in hand with human resource development



Rossen et al., CMI 2018

When and how to integrate NGS in the routine workflow?

Research or Surveillance:

WGS fits best in a batch-wise approach for analysing samples

Routine diagnostics:

Case-by-case approach, a balance should be kept between costs, quality, speed and complexity of the wet and dry processes.

- more samples tested in parallel: costs decrease

- longer reads may facilitate the downstream analysis: longer running times and higher costs

Rossen et al., CMI 2018

Practical issues in implementing wgs in routine diagnostic microbiology

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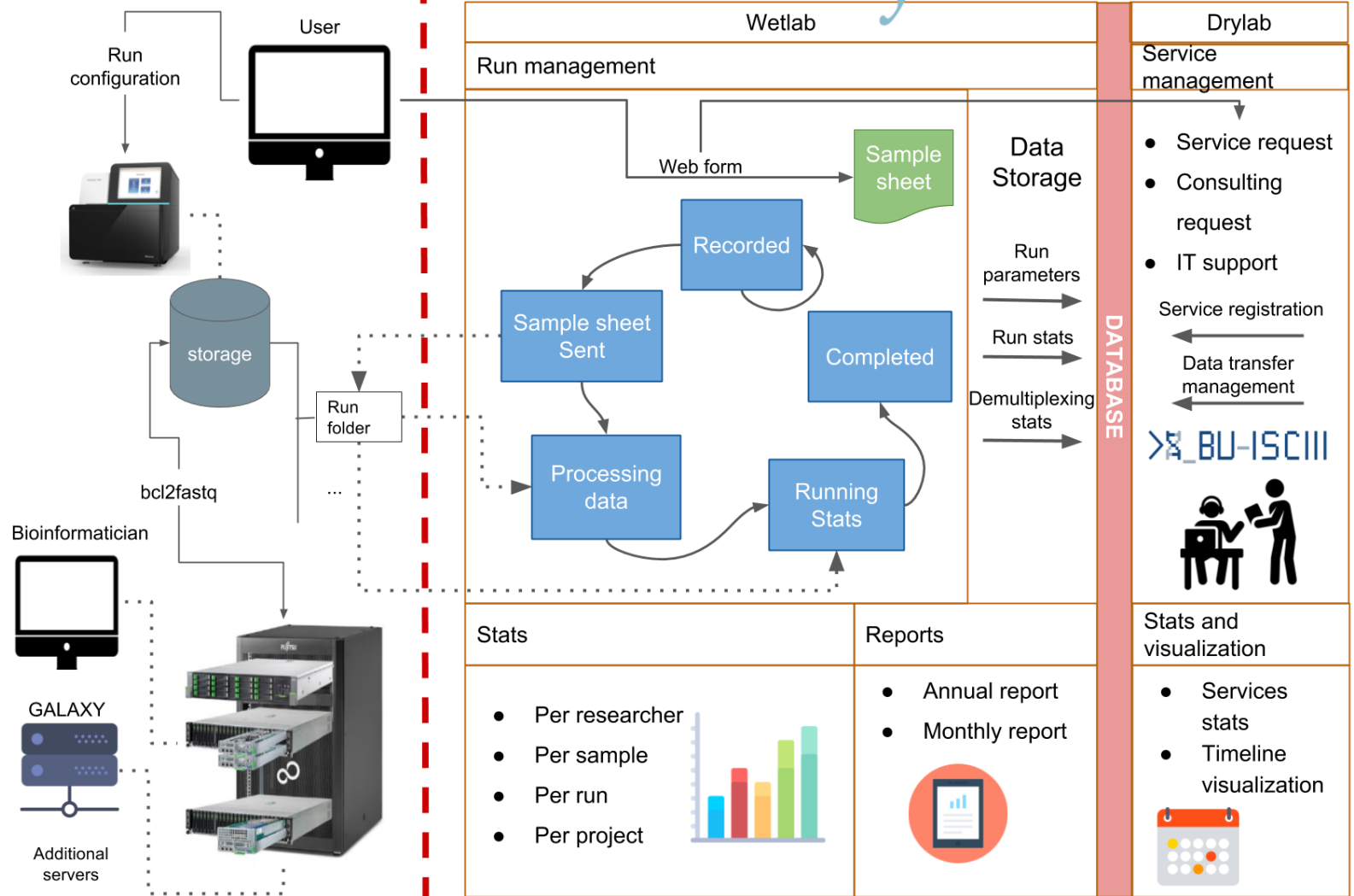
Rossen et al., CMI 2018



LIMS for NGS Labs

For NGS labs (Genomics Unit), there are common sub-processes that comprise the overall workflow, these include:

- Request for Services
- Sample Receiving
- DNA/ RNA Extraction
- Library Construction
- Sequencing
- Post Sequencing Processes such as Primary and Secondary Analysis

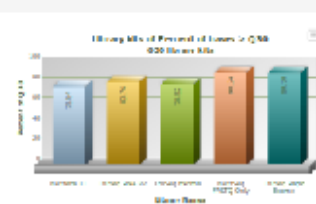


Collecting Statistics per RUN

Graphics of all Library Kit

The following graphics show the information all libraries that marches past, and time periods.

Percent of >Q30 Statistics per Library Kit



Mean Quality Score Statistics per Library Kit

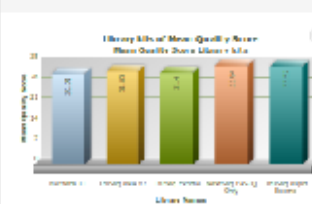


Fig. 3 Library kits, Samples with $Q > 30$.

Statistics comparison for Q > 30 for Investigator bmartinezd



Statistics comparison for Mean Quality Investigator benartinez

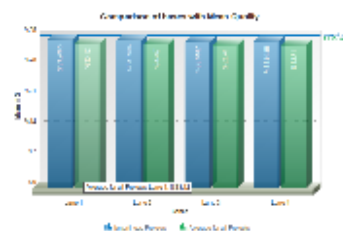


Fig. 6 Quality Statistics per Investigator.

Overall Quality of the Sample

Quality for the Sample SS-529-16

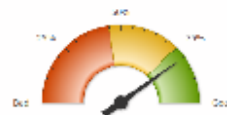


Fig. 7 Overall Quality Sample.

Chart for the Top **Unknow** Barcodes In the Run.

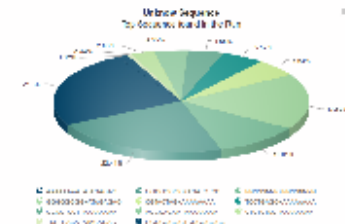


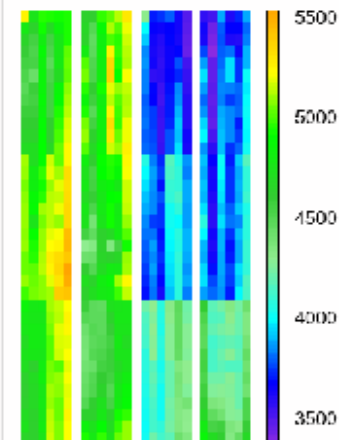
Fig. 4 Pie graphics for Unknow Barcodes.

Statistics for Mean Quality for Investigator [blindness](#)

Fig. 5 Mean Quality of Investigator.

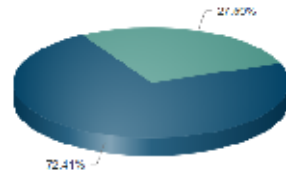
Flow Cell Chart

HI WIMPECKY Intensity



A) Relation of Completed and Unfinished Runs.

Number of Runs performed on the year

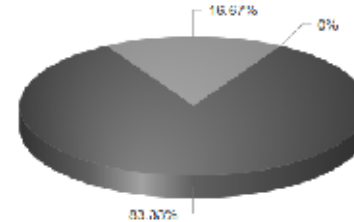


Completed Runs Not Finished Runs

Error Rate for the year 2017.

B)

Projects done per investigator on year 2017
Percentage

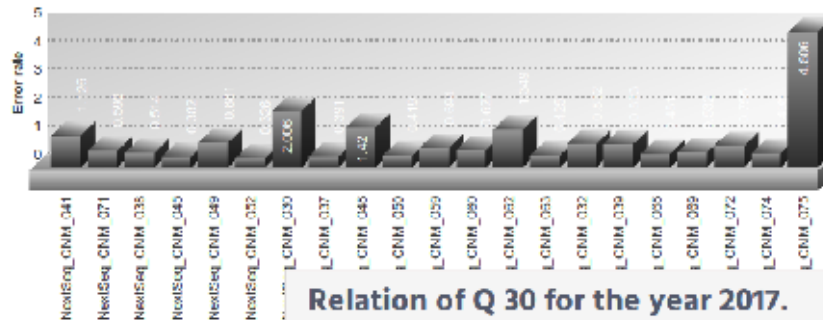


1 - 5 6 - 10 more than 10

C)

Error Rate for the runs done on year 2017

Error rate



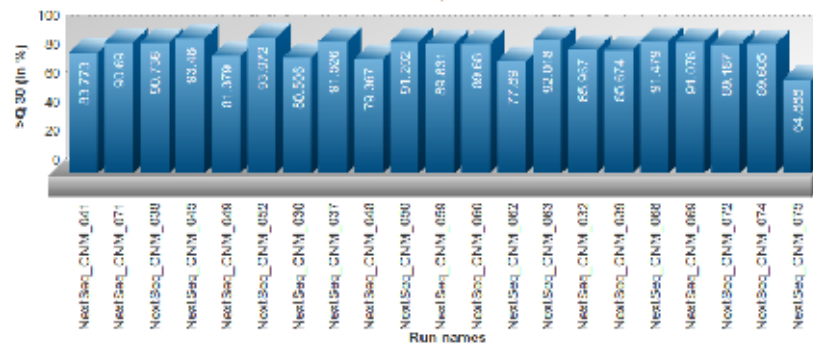
Relation of Q 30 for the year 2017.

D)

Fig. 8 Annual Reports A) Pie chart with relation of completed and unfinished runs. B) Pie charts with projects done per investigator during a year. C) Error rate percentage in the runs completed in a year. D) Relation of quality of runs done in a year.

>Q30 for the runs done on year 2017

Q30



Practical issues in implementing wgs in routine diagnostic microbiology

Table 1
Turnaround time and cost implications for routine WGS

Step	Turnaround time		Cost implications	
	Estimated time (hours)	Determinants	Estimated cost per sample ^a (euros)	Determinants
DNA extraction	1–2h	Choice of kit, additional steps (e.g. enrichment), automation	10	Kits vs. reagents, technician hands-on time vs. automation
Library preparation	4–6h	Method (enzymatic vs. shearing), choice of kit, automation	30	Choice of kit, automation
Sequencing ^b	50h	Platform, chemistry, read length, run protocol	75	Platform, chemistry, read length, number of samples per run/coverage
Initial analysis ^c	1–2h	Depending on number of samples, computing power, available software and pipelines	NA	Commercial vs. free software, availability of bioinformaticians, computer infrastructure
Specific analysis ^d	4h			

^a For performing all steps in house; direct costs and consumables not including personnel.

^b Assuming Illumina Miseq 250×2, 16–24 samples in one run.

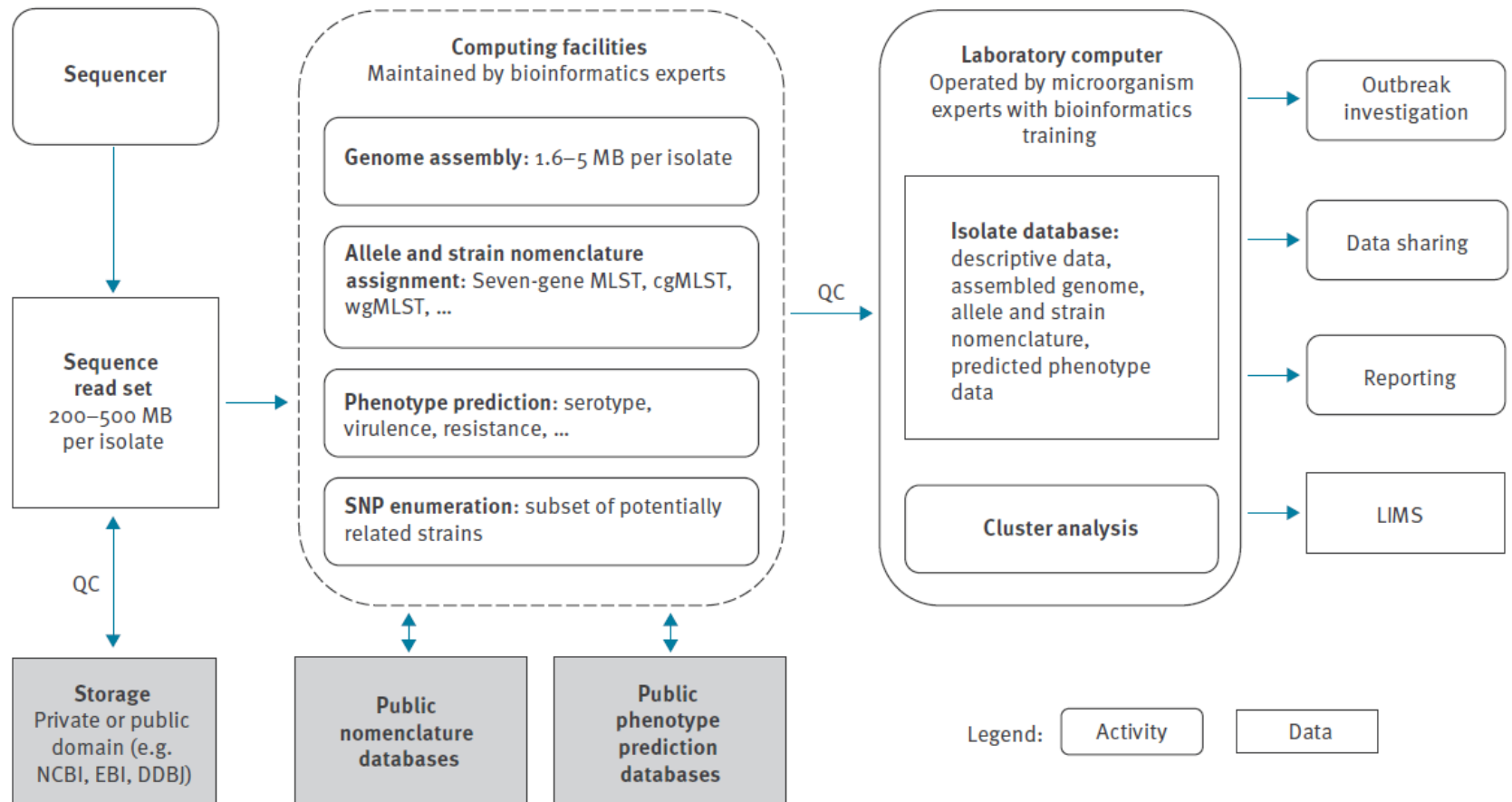
^c Quality control, read trimming, assembly, annotation.

^d Specific analysis dependent on the clinical/epidemiological circumstances.

Rossen et al., CMI 2018

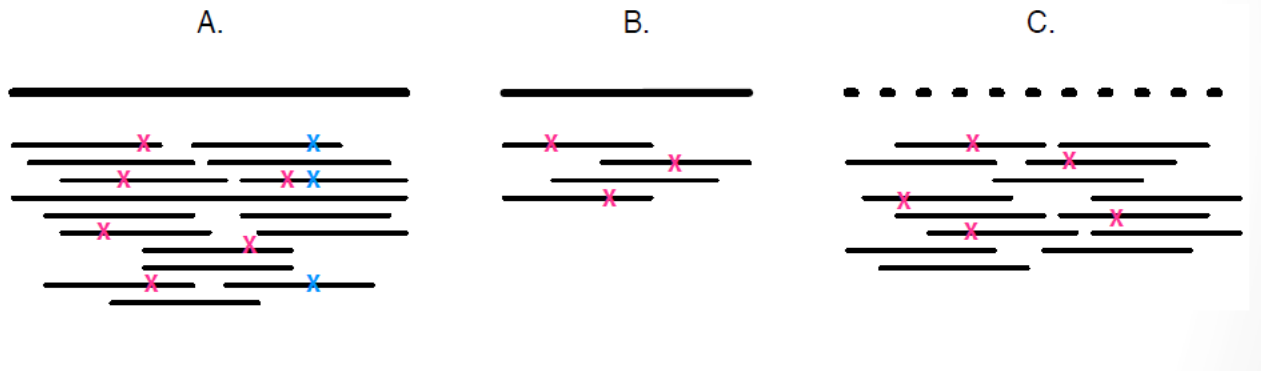
Potential solutions for computing and storage in PulseNet International laboratories, May 2017

Nadon et al., Eurosurv. 2017



Sequencing terms, three aims

(A) Resequencing (B) Read counting (C) Assembly



Sequencing terms

Breadth of coverage

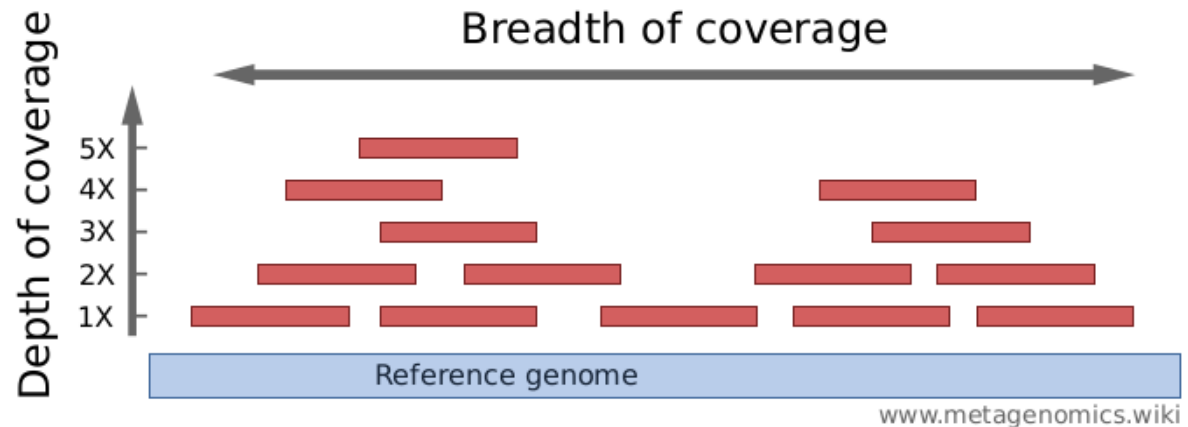
How much of a genome is "covered" by short reads? Are there regions that are not covered, even not by a single read?

Breadth of coverage is the percentage of bases of a reference genome that are covered with a certain depth. For example: 90% of a genome is covered at 1X depth; and still 70% is covered at 5X depth.

Depth of coverage

How strong is a genome "covered" by sequenced fragments (short reads)?

Per-base coverage is the average number of times a base of a genome is sequenced. The coverage depth of a genome is calculated as the number of bases of all short reads that match a genome divided by the length of this genome. It is often expressed as 1X, 2X, 3X,... (1, 2, or 3 times coverage).



Glossary of some commonly used sequencing terms

Besser et al., Clin Microbiol Infect, 2018

Adapter	Any short piece of DNA of known sequence that one adds to the ends of their unknown DNA of interest, usually for the purpose of eventually allowing a sequencing primer to hybridize at this position
Amplicon sequencing	Ultra-deep sequencing of PCR products for analysing genetic variations
ANI	Average nucleotide identity—an analysis method that assesses the nucleotide identity between genetic regions shared by two isolates
Assembly	Genome assembly is the process by which many short DNA sequence fragments, such as those generated by next-generation sequencers, are reassembled into a representation of the original genomic sequence
Bridge amplification	A PCR technique that embeds DNA on a solid surface for sequencing. It is used by Illumina's platforms
Contig	A contiguous consensus sequence derived from the assembly of many short, overlapping DNA fragments
cgMLST	Core genome multi-locus sequence typing—an analysis method that detects variation in genes that are present in the majority (>97%) of strains of a given species
Coverage (read depth)	The average number of reads that include a given nucleotide in the reconstructed sequence
Draft genome	Sequence of genomic DNA having lower accuracy than finished sequence; some segments are missing or in the wrong order or orientation
Emulsion PCR	A PCR technique that is conducted on a bead surface within tiny water bubbles floating on an oil solution. It is used by IonTorrent platforms
Error rate	The per-read error rate is defined as the proportion of reads containing sequencing errors
Flow cell	A glass slide containing small fluidic channels, through which polymerases, nucleotides and buffers can be pumped during sequencing
High-quality SNP	A single nucleotide polymorphism that has been verified using specific criteria such as: sequence coverage, sequence quality, and population and allelic frequency
Homopolymer	A DNA sequence (two or more base pairs) consisting of the same nucleotide
Index (barcode)	Unique individual DNA sequences added to each sample so they can be distinguished and sorted during data analysis. Enables sequencing of multiple samples per instrument run
Massively parallel sequencing	High-throughput DNA sequencing approaches that use the concept of miniaturized massive parallel processing to sequence 1 million to 43 billion short reads per instrument run
Metagenomics	The study of genetic material recovered directly from the primary samples
Paired-end reading	Sequencer starts reading DNA fragment at one end, finishes this direction at the specified read length, and then starts another round of reading from the opposite end of the fragment
Per-base sequence quality (accuracy)	The sequence quality score for each individual base position in a sequence. Typically, phred scores are used, where $Q = -10 \log(\text{Error Probability})$. A Q30, for example, means a 1 in 1000 likelihood of an incorrect base call at that position
Pyrosequencing	Sequencing is performed by detecting the nucleotide incorporated using enzymatic reactions after which the substrate emits light
Read	A unit of continuous DNA sequence derived from target DNA
Reversibly blocked terminator	A molecule added to a nucleotide to prevent addition of multiple nucleotides per sequencing cycle. Used by Illumina platforms
Sanger sequencing	A low throughput sequencing method based on the selective incorporation of chain-terminating dideoxynucleotides by DNA polymerase during <i>in vitro</i> DNA replication
Semiconductor sequencing	Sequencing is performed by detection of hydrogen ions that are released during incorporation of the nucleotide. Used by IonTorrent platforms
Sequencing by synthesis	Sequencing is performed by detecting the nucleotide incorporated by a DNA polymerase
Single-end reading	The sequencer reads a DNA fragment from only one end to the other, generating the sequence of base pairs
wgMLST	Whole genome multi-locus sequence typing—an analysis method that detects variation in all genes (core and accessory genes) of a given genome

Thanks for your attention!

Questions???