RNA-Seq in Cancer Research

Long-reads sequencing

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Because of the broad scope of the course, and the limited time, it will be a high-level review of long-read RNA sequencing.

The participants who already have experience with long-reads RNA-seq are very welcome to contribute their comments along the lecture.

Long-reads bulk RNA-seq

Long vs Short

Technology overview

- Principles: Nanopore and PacBio
- RNAseq library prep: strand switch and full length reads
- Current hardware: Machines and Flow-cells

Accuracy overview

- PacBio: CCS; Data files Subread & CCS non-aligned BAM-s
- Nanopore: Base-callers, Pores, Consensus

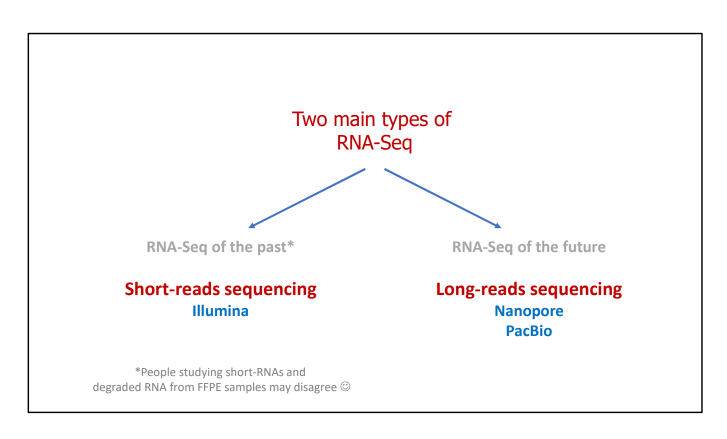
Tasks and tools overview

- Transcript isoforms identification / Genome annotation
- Quantitative analysis: DGE / DTU
- Tools, Workflows & Manufacturer supported bioinformatics

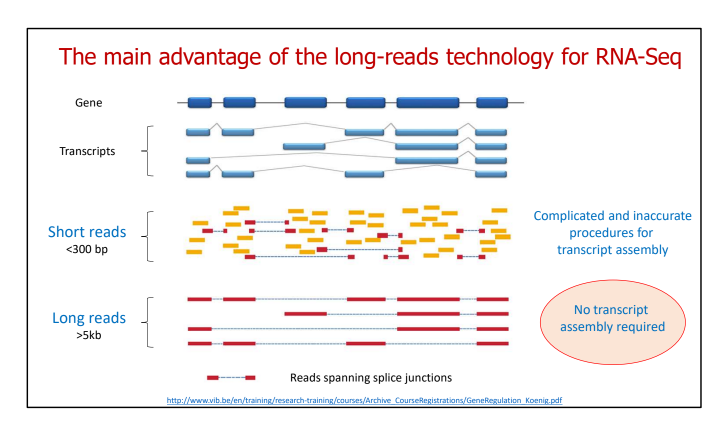
Selected examples of tools and pipelines

- PacBio Transcript Assembly: IsoSeq3, SQUANTI, TAPPAS
- ONT quantitative analysis: NanoPack, Pychopper, wf-transcriptomes pipeline

I will start with a high-level overview of the technologies, then focus on some RNA-seq specific aspects, and then will go to bioinformatics aspects discussing tools, tasks and some examples.

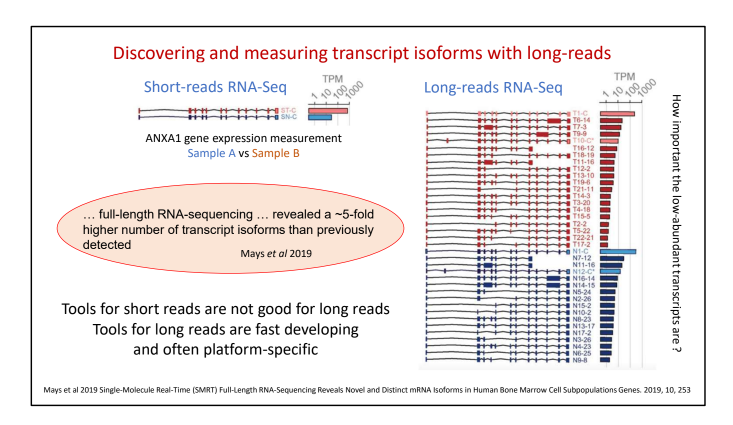


I already mentioned today that there are two main types of RNA-sequencing: the short-read and the long-read sequencing:)



The main advantage of the long-reads technology for RNA-Seq is that the long reads can span the entire transcripts, eliminating the need in the transcripts assembly (the bottleneck of the short-reads technology).

Of course, long-reads also have their limitations. Thus, later we will discus potential issues with the accuracy of some long-reads sequencing techniques. So, it is worth noting here, that the high accuracy of sequencing is not needed for RNA-seq gene/transcript expression measurement. Two or three errors per hundred nucleotides, do not complicate identifying the transcript. The length is much more important.

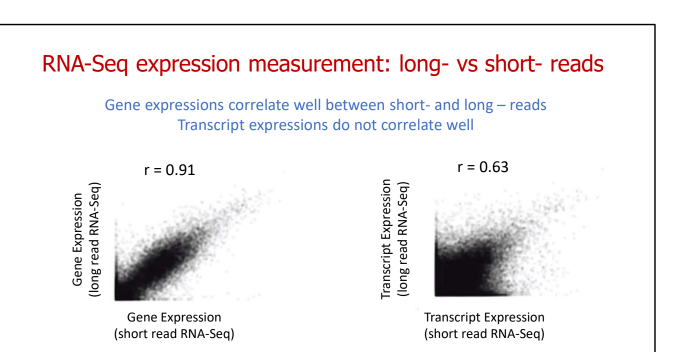


Because of the ability to read the entire transcripts, the long-reads RNA-seq reveals several times more transcripts than it was previously detectable by the short reads.

Interestingly, many of the new detected transcripts are low-abundant. So, we still don't know their biologic relevance.

Another question is:

If so many transcripts were missed by the short reads, does it mean that all the previous results obtained with the short reads were wrong?



Jonathan Göke, The SG-NEx project: nanopore long-read RNA-sequencing of human cancer cell lines Nanopore Community Webinar, 28Feb 2019

No, it doesn't.

On the left panel you can see that after aggregation per gene the results of short-reads RNA-sequencing correlate well with the long-reads. Of course, the right panel shows much worse correlation at the TRANSCRIPT level.

The point is that most of the gene expression results reported so far from the short reads were reported per gene, not per transcript. So, most of the previously published short-reads results are valid. Of course, the rare short-reads TRANSCRIPT-specific data might need to be validated.

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Now, lets review the current long reads sequencing technologies.

At the moment, there are two long-read sequencing technologies: Oxford Nanopore and Pacific Bioscience.

We will not discuss Synthetic long reads here, except for a single slide to acknowledge the existence of an alternative to the true long-reads technologies.

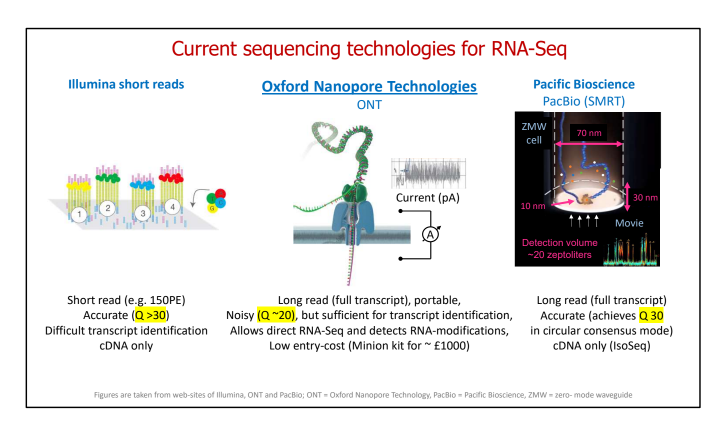
Current sequencing technologies for RNA-Seq **Oxford Nanopore Technologies Pacific Bioscience** Illumina short reads ONT PacBio (SMRT) ZMW cell Current (pA) Short read (e.g. 150PE) Long read (full transcript), portable, Long read (full transcript) Accurate (Q >30) Noisy (Q ~20), but sufficient for transcript identification, Accurate (achieves Q 30 Difficult transcript identification Allows direct RNA-Seq and detects RNA-modifications, in circular consensus mode) cDNA only Low entry-cost (Minion kit for ~ £1000) cDNA only (IsoSeq) Figures are taken from web-sites of Illumina, ONT and PacBio; ONT = Oxford Nanopore Technology, PacBio = Pacific Bioscience, ZMW = zero-mode waveguide

I included the Illumina short-reads technology on this slide (on the left) to explain why it can not produce long reads.

The Illumuna sequencing is based on amplifying each initial DNA fragment to a cluster, and then adding one nucleotide per cycle, with different nucleotides coded by different colors. There are very nice videos on the Illumina web site, which explain the technology in great details.

Unfortunately, some DNAs in clusters occasionally miss a cycle, and after 2 or 3 hundred cycles the molecules in the clusters are going out of sync. So, this technology can not read longer fragments.

The advantage of the Illumina technology is that, until the clusters go out of phase, the sequencing is very accurate: less than one error in a thousand of bases.

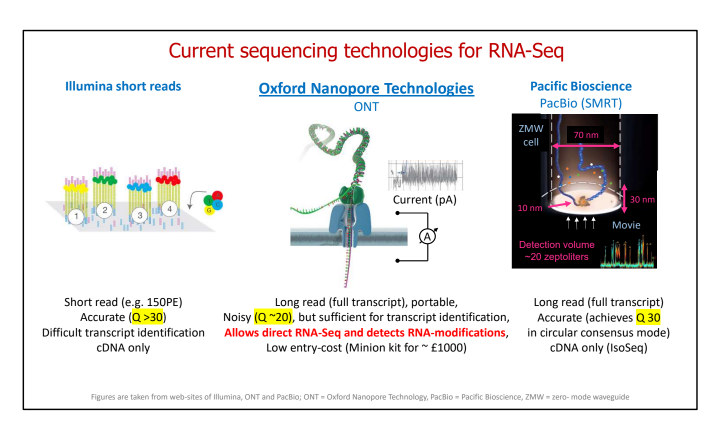


The Nanopore technology is shown here in the middle.

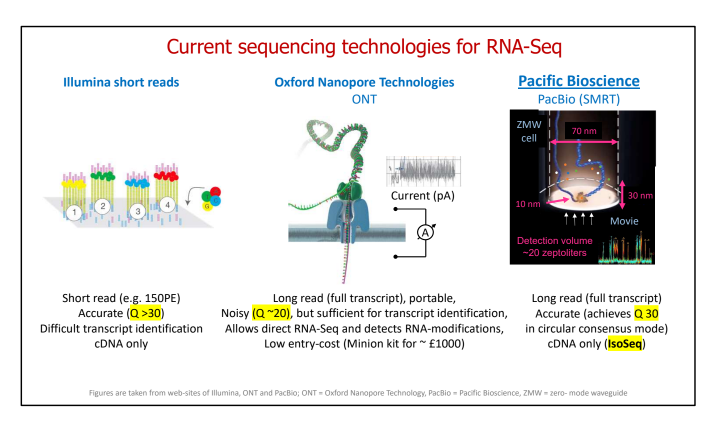
It just passes DNA fragment through a pore, while measuring the ion current passing through the same pore at the same time. Because different nucleotides have different size and charge, they block the ions' passage in a different way, and so they have distinct current signatures.

The length of Nanopore sequencing is limited just by the length of the fragment, reaching hundreds of thousand or even millions of bases.

The negative side of this technology is that it is still much less accurate than the short-reads sequencing: with some errors per each hundred of nucleotides. However, for many RNAseq applications such accuracy is absolutely sufficient.



Importantly, the nanopore can sequence RNA directly, without converting it to cDNA before sequencing.

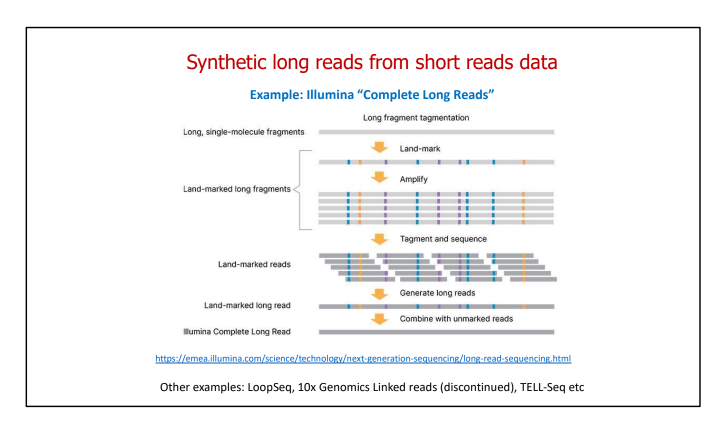


Finally, the PacBio technology is shown on the right. PacBio is also marketed as SMRT sequencing, meaning that it reads Single Molecule in Real Time.

They place a single DNA-polymerase at the bottom of a tiny well. Then the DNA synthesis is just filmed by a tiny camera in real time. Different nucleotides are labelled by different colors. When a new nucleotide is being added to the DNA, the DNA-polymerase retains it for a certain time, which is recorded as a peak in the movie.

To make such movie possible, it's important to exclude filming of the other nucleotides, which are not being retained by DNA-polymerase, but still are present in the solution. This is made by illuminating only a very small volume around the DNA-polymerase. Because of the Brownian movement, the non-retained nucleotides quickly cross the volume, so they and are not registered in the movie. Selective illumination of such a small volume is done by some physical miracle, called Zero-Mode Waveguide (ZMW). Somehow, when the diameter of the well is small enough, the light does not go through the well, but only propagates to a certain depth. A well with about 70 nm in diameter allows to illuminate a volume of just 20 zeptoliters.

Honestly, illuminating and filming a single DNA-polymerase molecule in action sounds like SciFi to me. However, somehow it works.



Finally, I would like to mention that there are alternatives to the true long-reads sequencing technologies.

For instance, Illumina provides a solution, when special "Land-marks" are placed to long DNA fragments before they are shredded and sequenced as short reads.

Then the initial long DNA fragments can be computationally re-assembled from the "land-marks" information.

There are other similar technologies that allow to assemble synthetic long reads from the short reads data.

This could be a cheaper alternative to the true long-reads technologies: the future will show.

I will not discuss Synthetic Long Reads in this lecture later. You may see a comparison between PacBio and Synthetic Long Reads following this link:

https://www.pacb.com/blog/the-hifi-difference-true-long-reads-vs-synthetic-long-reads/

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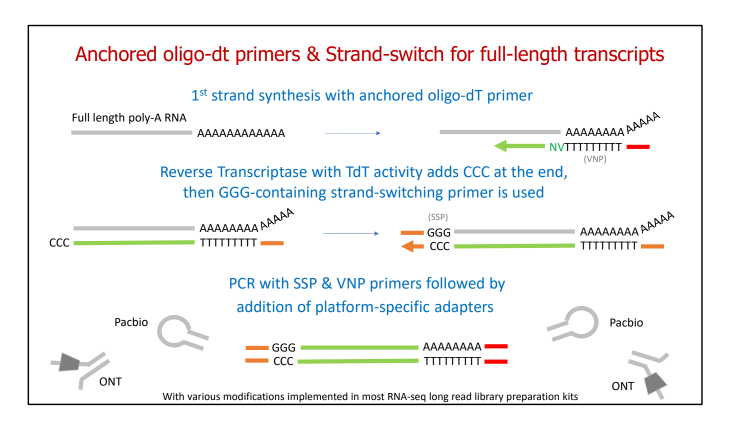
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Now let's talk about RNA-seq library preparation for the true long-reads sequencing technologies: ONT and PB.

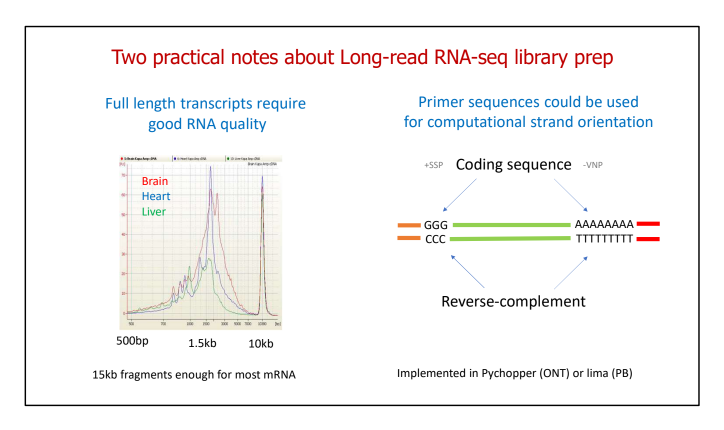


This is a *simplified* scheme explaining the widely used technique for making cDNA libraries for full-length transcripts.

SSP: Strand Switching Primer (sometime called template-switching)

VNP: V = A or G or C; N = A or G or C or T; P= Primer

TdT: Terminal deoxynucleotidyl Transferase



Of course, to make the full-length cDNA libraries, the full-length RNA molecules should be present in the extract in the first place.

If the non-degraded RNA was used, then 15kb read length should be enough to sequence most of human transcripts in any tissue.

An additional advantage of using the Strand-switching technique described in the previous slide is that it allows computational detection of the strand orientation. So, all the long-reads libraries are stranded: the coding sequence is flanked by *direct* SSP and *reverse-complemented* oligo-dT primers

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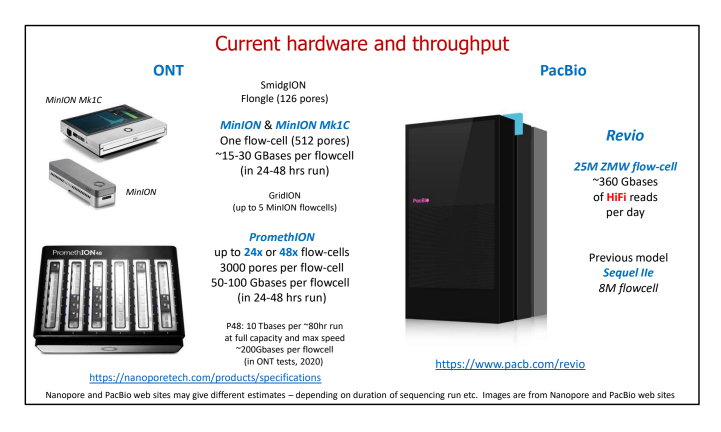
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Then the prepared libraries go to the sequencing machines.



There are many models in Nanopore.

For a long time, the entry-level ONT model was **Minion**. It's a size of 10x2x3 cm, required connection to a laptop by USB cabel.

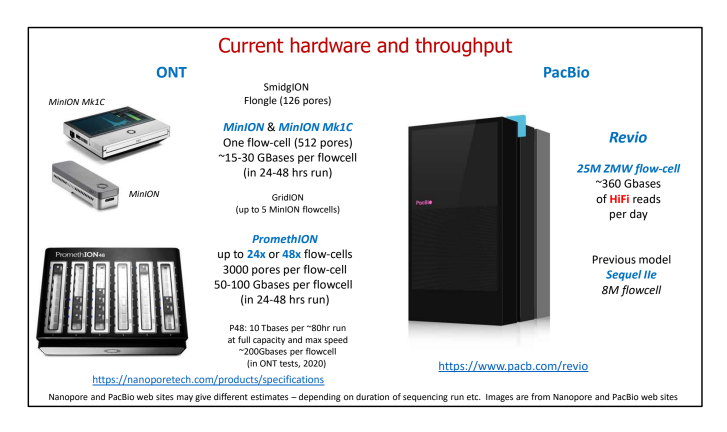
MinION Mk1C uses the same flowcell as Minion but doesn't need a laptop and includes on-board basecalling.

The highest end of Nanopore is **PromethION**, it has much larger flowcells and up to 48 of them.

1x PromethION flow-cell may allow 50-75x human genome may reach >30 consensus accuracy (caveats discussed later).

The theoretical throughput of 48 flowcell Promethion run at full speed and max capacity is mind-blowing.

There are intermediate and smaller models.



The recently launched (in 2023) "latest and greatest" model of PacBio is Revio.

It was a large step forward against the previous models.

Importantly PacBio can produce HiFi reads – their accuracy could be even higher than in Illumina (discussed later).

Although PacBio is still more expensive than ONT per base, **Revio** already allows to sequence human genome for USD1000 (with long accurate phased reads). Combined with **Kinnex** library preparation (a new technique discussed later) **Revio** may make PacBio RNA-seq even more affordable.

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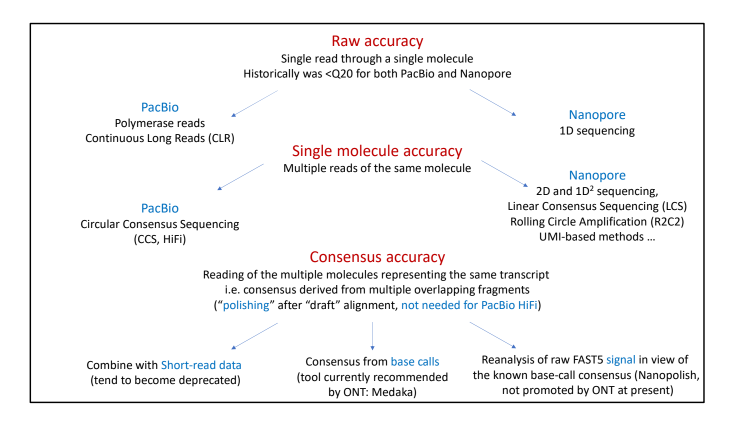
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I have already mentioned previously "consensus accuracy" or "HiFi reads".

These terms relate to the ways how long reads sequencing technologies improve the accuracy of their reads.



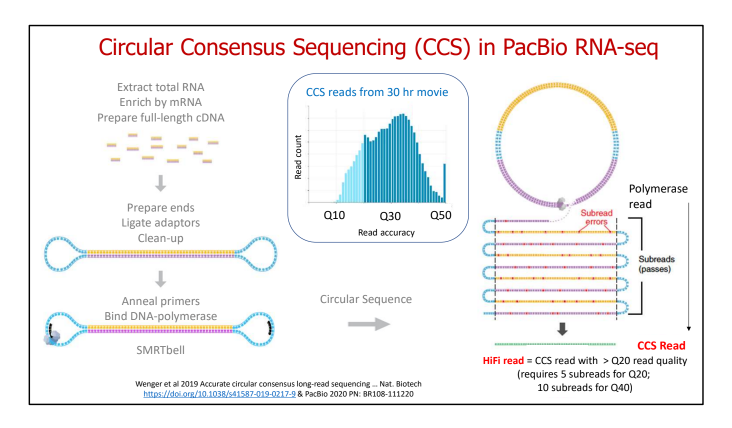
Typical *raw* Illumina base Quality is Q30-40 (which is less than 1 error per 1000 bases). The *raw* long-reads sequencing is significantly less accurate.

This slide contains many new terms. I will explain some of them now, and some later. For PacBio the *raw* data may be called Polymerase reads (or Continuous Long Reads, CLR). For Nanopore the *raw* data may be called 1D sequencing.

To improve accuracy the long-read technologies have tried to read the same sequence many times.

This solved the problem for PacBio: when a DNA fragment is circularized and red many times (as will be shown in the next slide), the Circular Consensus Reads (CCS) accuracy easily exceeds the Illumina raw base quality (if the same sequence is red for 5 and more times).

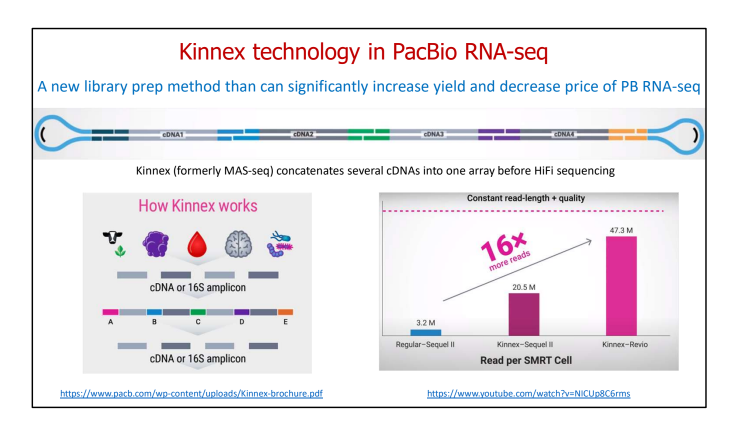
Nanopore also tried reading the same molecule twice (called 2D and 1D2). Unfortunately, because of the non-random distribution of Nanopore errors, this was less successful than in PacBio. So, Nanopore still uses consensus accuracy: reading multiple fragments from the same gene to improve the accuracy. With sufficient depth (and price) Nanopore claims that it may match Illumina accuracy (as will be shown later). However, it's still a work in progress (in 2024).



This slide explains the PacBio Circular Consensus Reads.

Because PacBio *raw* data (so called Polymerase Reads or Continuous Long Reads) have low accuracy (<20), PacBio came up with a nice trick to improve the accuracy. During the library preparation they make the fragment circular. Then, during the sequencing the same fragment is red again and again many times. Because the errors are random, the consensus sequence after multiple reads becomes as accurate as the Illumina short reads (or even better accuracy). The length of the circular consensus reads in PabBio may easily achieve 10-15 kilobases, which is enough for most of human full-length RNAs.

Importantly, unlike to Nanopore, PacBio can not sequence RNA directly: it has to be converted to cDNA before sequencing.

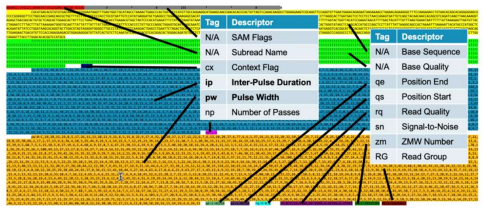


Kinnex (formerly MAS-seq) method concatenates several cDNAs into one array before taking it into CCS. This is still very new. However, combined with the high output of the Revio sequencer, this may pave the way for more affordable PacBio RNA-seq.

PacBio CCS file formats: BAM-s everywhere

- PacBio machine produces raw data as a BAM of unaligned subreads (with all base qual=0 & all read qual=0.8) *
 The CCS workflow produces a BAM with unaligned consensus reads with meaningful base & read qualities
 Alignment (mapping) programs will produce aligned BAM files that retain PacBio tags
 - * the latest versions of sequencers (Sequel IIe) may output the consensus BAM-s to reduce data size

Additional tags in PacBio BAM-s



PacBio Webinar: PacBio Data Deep Dive: A Closer Look at HiFi Sequencing, 24 March 2021 https://pacbiofileformats.readthedocs.io/en/12.0/index.html

A short side-tracking about PacBio file formats.

In addition to the base sequence and base quality PacBio needs to record the kinetic information (pulse width, inter-pulse duration) and some other information about their movies. So, they decided not to use FASTQ file format. Instead, they initially decided to use their own version of BAM files.

Also, PacBio uses a special sort of XML file format. You may see more details here: https://pacbiofileformats.readthedocs.io/en/12.0/DataSet.html

In the recent machines, PacBio implements CCS workflow on sequencer, by default outputting only CCS reads, and discarding the sub-reads information to reduce the data size.

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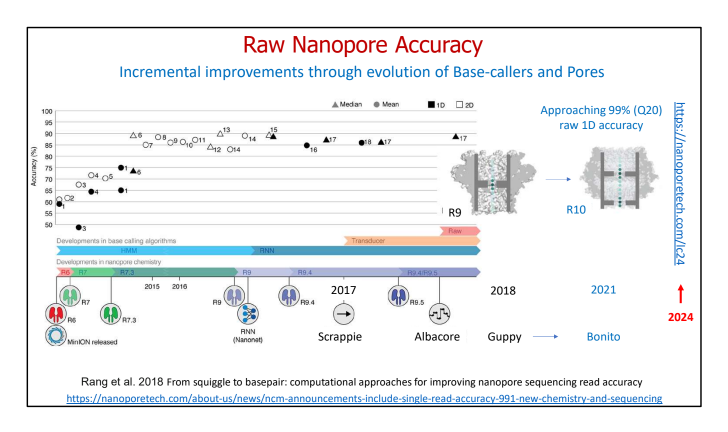
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Coming back to the accuracy of long reads sequencing.



Once upon a time (about ten years ago) the accuracy of Nanopore reads was awful: 50% errors.

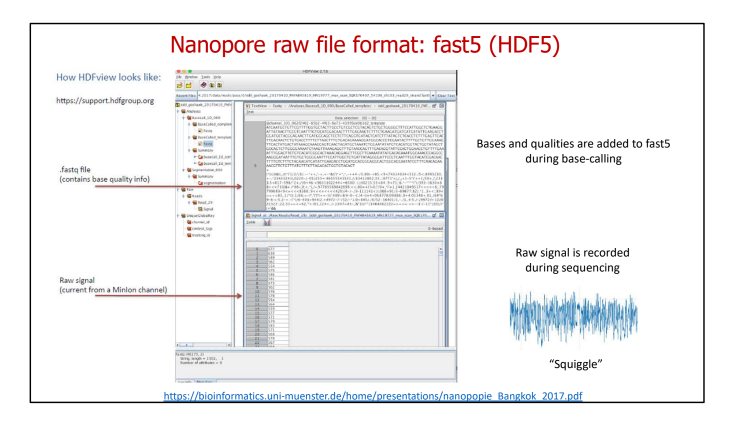
However, it was dramatically improved since that by developing the new pores, and most importantly, by the new algorithms for calling bases from squiggles (it's a term to describe the ionic current fluctuations recorded by the pore).

The breakthrough in the basecalling algorithms development happened when Nanopore decided to use machine-learning for this.

Practically, this means that the training sets and even some hyperparameters (such as depth of the network etc) may vary even between different models of the same basecaller. For instance, model trained on human DNA may not be perfect for bacteria or plants ...

Of course, the best available models were trained on the human data, which is handy for our Cancer research context \odot

A couple of years ago, Nanopore announced that the latest at the time version of basecaller **Bonito** with data from **R10 pores** approached **99% raw accuracy**, which is a remarkable progress comparing with initial 50% just 10 year ago.



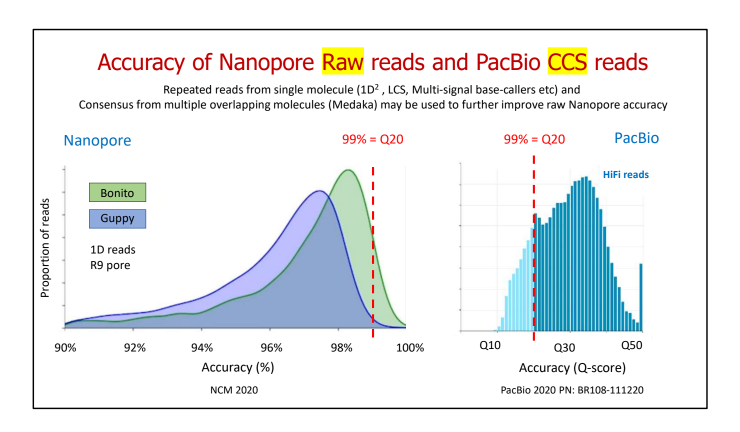
Again: a side-tracking about file formats.

To record raw data ("squiggle") along with the bases, their qualities and some other data, Nanopore initially used a sort of XML data format, called fast5.

There are viewers for such files. This slide illustrates the structure of the file and some data contained in different tags.

However, the fast5 files could be really big (many hundreds of gigabytes), and processing fast5 files may be slow. A newer version of the Nanopore data files is called POD5.

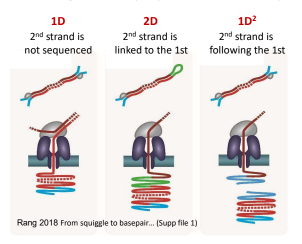
You may find more information about the Nanopore data formats here: https://community.nanoporetech.com/technical_documents/data-analysis/v/datd_5000_v1_revr_22aug2016/file-formats



However, 99% accuracy is still just Q20. So, the accuracy of *raw* ONT data is still much lower than *HiFi* PabBio.

Nanopore: techniques for repeated reading of the same molecule

ONT problem with repeated and consensus approaches: the errors are not fully random e.g. a homopolymer or secondary structure will always cause the error



Methods in development ...

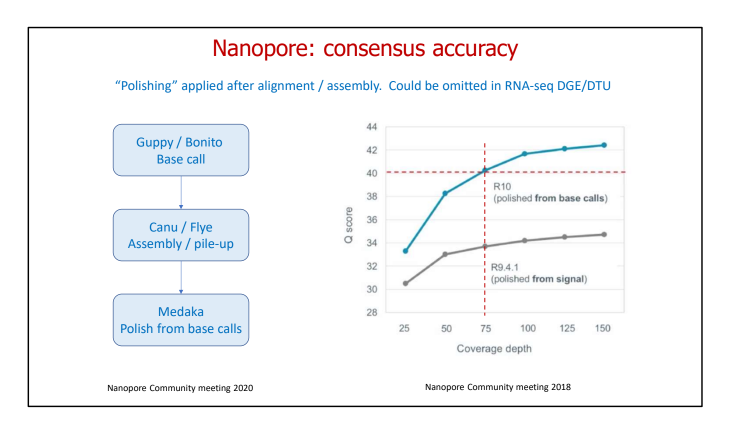
- Rolling Circle Amplification (R2C2)
- Linear Consensus Sequencing (LCS)
- UMI-based "multi-signal" base callers hits Q30 by just 3 reads with the same UMI
 - 8B4 random base substitutes (e.g. T U) to tackle homo-polimers

In 2016 use of 2D improved accuracy from 60% to 90%. Now ONT prioritizes new base callers and pore designs. Although, ONT keeps exploring new methods for obtaining multiple reads from single molecule, currently they are not applied to RNA-seq.

Nanopore also explored with the multiple reads from single molecule, but this direction looks abandoned at the moment.

- First, it red the same molecule twice by adding a hairpin (so called 2D libraries);
- Then it tried to do the same without adding a hairpin (1D2): because the 2^{nd} strand is still in vicinity of the pore, in ~75% cases it may follow the first strand even w/o hairpin.

These methods gave a modest improvement, but nowhere near the CCS reads in Pacbio. There are some other experimental approached in development.

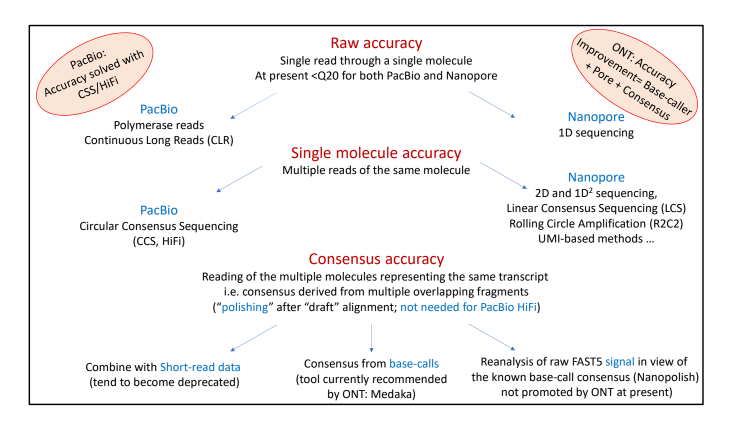


The issue complicating Nanopore basecalling is that the errors are not random. If some sequence produces a squiggle that is hard to decipher at the first pass, it is also hard at the second pass. Most errors are in homopolymers (repeats of the same nucleotide), which is not the common pattern in RNA (except for the poly-A tail).

Nanopore works on it, both on technical and marketing sides.

Their current marketing clams are that with sufficient depth of sequencing the Nanopore *consensus* accuracy may become the best of the long read sequencing method.

This is yet to happen ... However, with sufficient depth (and associated cost) they truly can rich a very good accuracy.



To summarize, The *raw* reads accuracy is low for both PacBio and Nanopore.

PacBio solved it by Circular Consensus Sequencing (**CCS**), easily reaching Q40 and above for reads of 15kb.

Nanopore achieved quality Q15-20 by new **basecallers and pores design**. This accuracy is enough for most RNAseq applications.

Because Nanopore is still cheaper, which allows to get higher depth for the same price, at the moment it is considered that Nanopore could be better for transcripts quantification and differential expression studies, while PacBio could be better for accurate transcript isoforms discovery (Pardo-Palacios et al 2023, https://www.biorxiv.org/content/10.1101/2023.07.25.550582v1)

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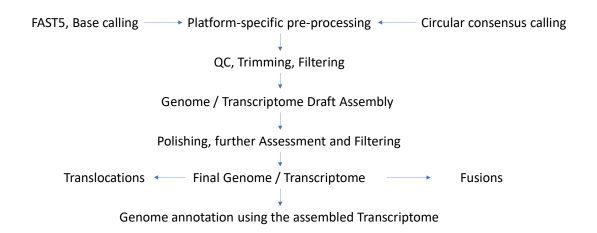
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This brings us to the next point: the tasks currently performed by long-reads RNAseq

Historically the Long Reads analysis aimed at Genome annotation / Transcriptome assembly



Till recently Differential Gene Expression was not amongst the tasks of Long-Reads RNA-seq analysis (at best: use Long Reads to get the Transcriptome, and then use Short Reads for DGE)

Till very recently the long reads RNAseq was too expensive to get sufficient depth for reliable transcripts *quantification*.

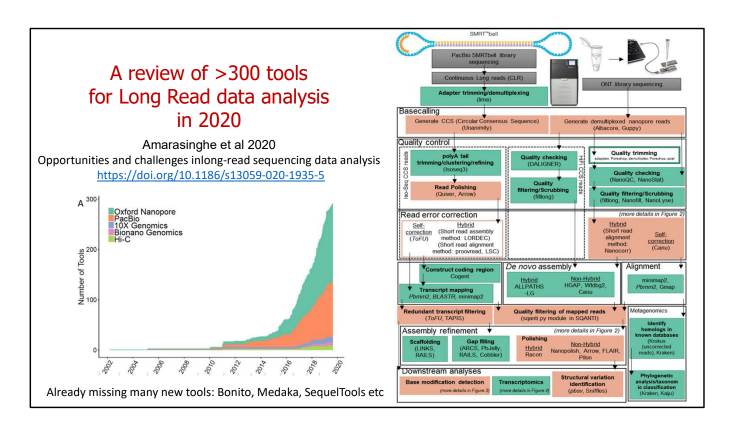
So, most of the tools, pipelines and papers published about long reads sequencing were focused on transcripts *identification*.

Identification of transcripts could be used for

- Genome annotation
- Transcriptome assembly (in absence of a reference genome) or
- Fusion detection

Such tasks as "Genome annotation" or "Transcriptome assembly" in absence of a reference genome may sound alien for human cancer researcher because human cancer research enjoys the best annotated reference genome and transcriptome available (i.e. human genome and transcriptome :)

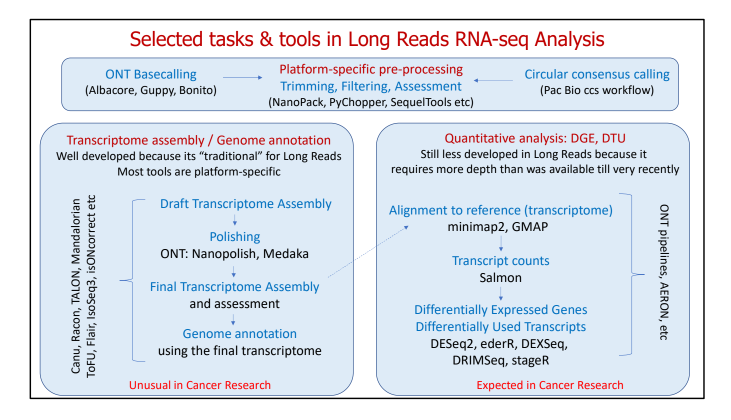
However, after adoption of Nanopore Prometheon (and maybe PacBio Revio+Kinnex in the near future) the long-reads technologies are producing sufficient and affordable depth of sequencing for the quantitative analysis too.



It looks like the tools for the long-reads data analysis are proliferating even faster than the tools for short-reads analysis:)

Only this review mentions hundreds of tools ...

It's now a separate branch of bioinformatics – not the tools development but the tools comparison :))



To simplify the chart shown in the previous slide, for the purpose of this short introductory lecture,

here I split long reads RNA-seq tools to 3 large categories:

- Pre-processing
- Transcriptome assembly/Genome annotation
- Quantitative analysis

I do not claim that mentioned tools are better than many others ...

The specific tools mentioned here are just to illustrate what was used in the literature, some of them are already deprecated.

And, of course, there are many RNAseq tasks and tools that are not even mentioned here, such as Nanopore Direct RNA-seq, RNA-modifications, poly-A tail length etc

Workflow managers

Bioinformatics tasks when writing a pipeline

- Design the workflow that puts the right tools in the right order.
- Install and configure all dependencies (i.e. tools and resources requird for analysis).
- Align outputs of the upstream tools with input requirements of the downstream tools.
- Arrange the locations of the source data, interim files and results.
- Write scripts that assemble all the pieces together, log and paralellize the computation, etc.





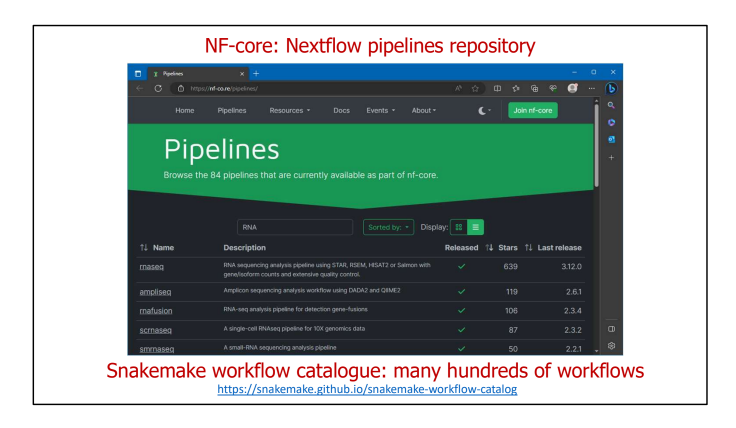


https://snakemake.github.io

https://www.nextflow.io/index.html

https://cromwell.readthedocs.io/en/stable

Even a simple bioinformatics task often requires multiple steps and tools. It takes lots of effort and expertise to write a pipeline connecting different tools. Workflow managers help to connect multiple tools into pipelines and facilitate reproducible research.



Writing and publishing pipelines is currently a strong trend in bioinformatics community.

Manufacturer supported bioinformatic solutions

| | Nanopore | PacBio |
|---|---|--|
| Software to control machines and for low-level tasks | MinKNOW | Instrument Control Software (ICS) SMRT-link |
| GUI solutions for standard tasks (could be on Server, Cloud, HPC, etc) | Epi2Me | SMRT-link |
| Command-line tools and pipelines for non-standard analyses | Epi2Me Labs Snakemake & Nextflow pipelines | SMRT-tools Cromwell & Nextflow pipelines |



Community developed tools and pipelines

Publications, GitHub. Workflow repositories (Snakemake, Nextflow, Cromwell)

Both Nanopore and PacBio provide extensive Bioinformatics support. With some pinch of salt, their manufacturer supplied bioinformatics is summarized in such table.

Along with the separate tools, both Nanopore and PacBio develop and publish the entire pipelines, using different workflow managers.

We will try one in our practical session ...

Long-read bulk RNA-seq

Long vs Short

Technology overview

- Principles: Nanopore and PacBio
- RNAseq library prep: strand switch and full length reads
- Current hardware: Machines and Flow-cells

Accuracy overview

- PacBio: CCS; Data files Subread & CCS non-aligned BAM-s
- Nanopore: Base-callers, Pores, Consensus

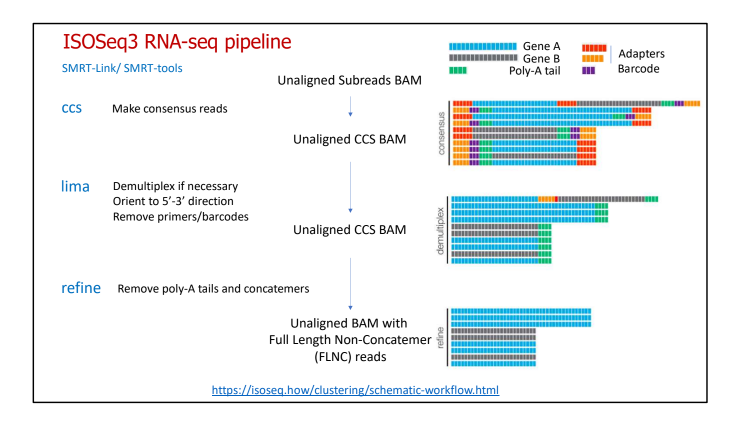
Tasks and tools overview

- Transcript isoforms identification / Genome annotation
- Quantitative analysis: DGE / DTU
- Tools, Workflows & Manufacturer supported bioinformatics

Selected examples of tools and pipelines

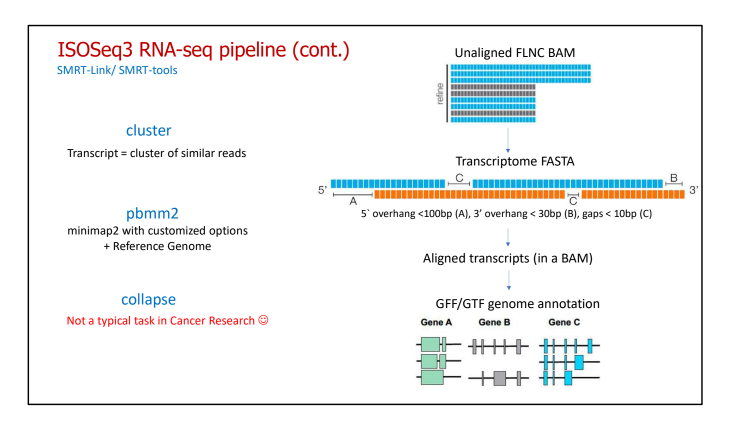
- PacBio Transcript Assembly: IsoSeq3, SQUANTI, TAPPAS
- ONT quantitative analysis: NanoPack, Pychopper, DGE DTU pipeline

Finally, I will briefly discuss some examples of the long-reads RNA-seq tools and pipelines.



I will start with IsoSeq (=Isoforms Sequencing) pipeline provided by PacBio. It was designed for transcripts isoforms identification.

It includes preprocessing that is required for virtually any PacBio RNAseq task. Tools used in this pipeline are available separately from their pages on PacBio's GitHub, as well they are included into SMRT-Link software or SMRT-tools toolset available from PacBio.

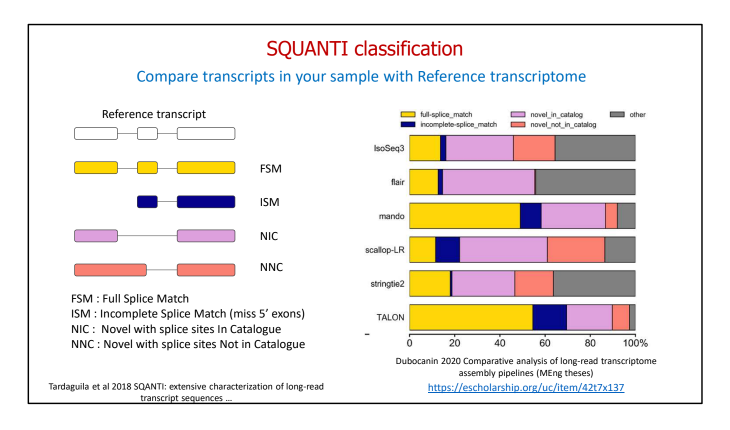


You can see that after alignment IsoSeq pipeline does not count specific transcripts (as we would expect for short-reads analysis).

Instead, it just "collapses" similar transcripts and generates genome annotation.

Not a typical task in Cancer Research.

However, the initial steps of IsoSeq pipeline still would be needed for the Iso-Forms quantification too.

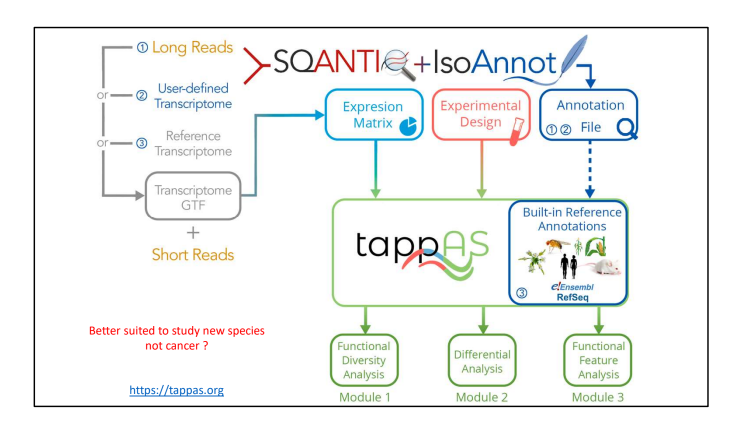


The down-stream step after the transcript isoforms discovery is to classify these isoforms.

Squanti is a very popular tool downstream of the transcriptome assembly. It compares the detected transcripts with the previously available reference transcriptome.

Surprisingly to me, despite the long reads supposedly spanning entire transcripts, there could be quite a disparity between results of different transcriptome assembly pipelines.

(shown on the right)



Squanti is a part of a wider ecosystem of tools for the transcriptome annotation, called **TAPPAS.**

Apparently, it also includes a differential expression functionality.

However, initially the *TAPPAS* quantification module relied on the additional short-reads data.

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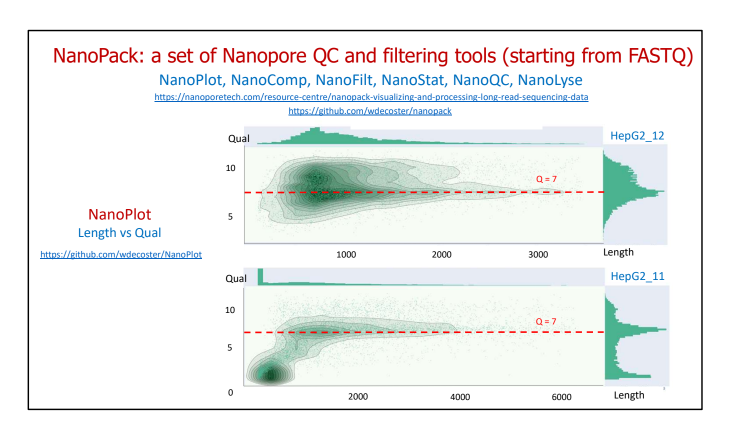
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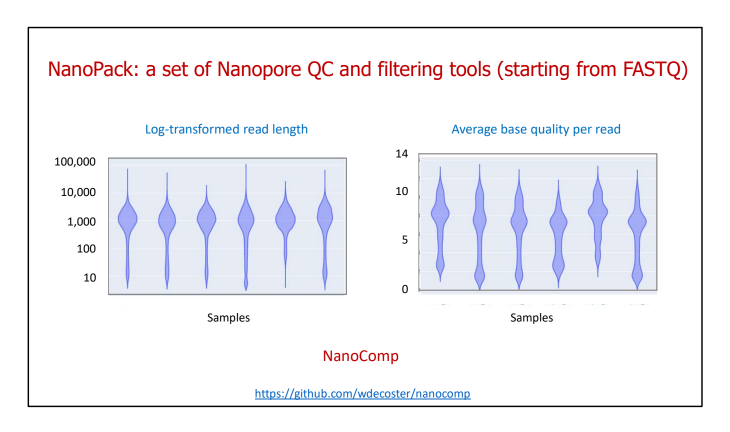
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The last part of the lecture will illustrate some Oxford Nanopore tools that we will use during the practical session.



NanoPack - Simple and straight-to-the-point toolset for QC and filtering Nanopore data (using FASTQ files).

NanoPlot shows a simple and very informative plot of reads Length vs reads Quality.



NanoComp allows to compare quality metrics between multiple samples. Actual plots could be interactive (showing additional information when mouse hoovers over the plot).

NanoPack: a set of Nanopore QC and filtering tools (starting from FASTQ)

Options:

-l, --minlength Minimum read length

-q, --quality Minimum average quality score --threads Number of threads to use

--headcrop Trim N nucleotides from the start --tailcrop Trim N nucleotides from the end

--maxgc Maximum GC content --mingc Minimum GC content --maxlength Maximum read length

Example

gunzip -c reads.fastq.gz | chopper -q 10 -l 500 | gzip > filtered_reads.fastq.gz

copper

https://github.com/wdecoster/chopper

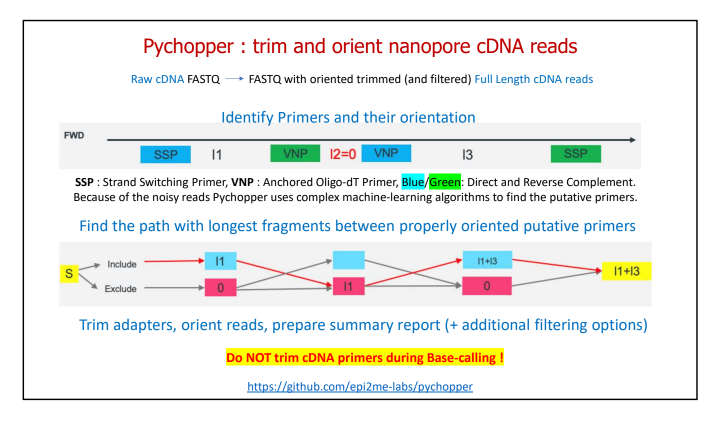
There is an older tool called NanoFilt

After evaluating the quality of the reads, we need to remove bad reads.

A common practice for Nanopore data includes removal of reads by mean qual > 7 (compare to minimal Q20 in PacBio HiFi :)

Copper can filter reads basing on many different parameters too.

Important: NanoPack's *copper* is not the same as *Pychopper* discussed on the next slide!



Pychopper trims, orients and filters RNAseq reads.

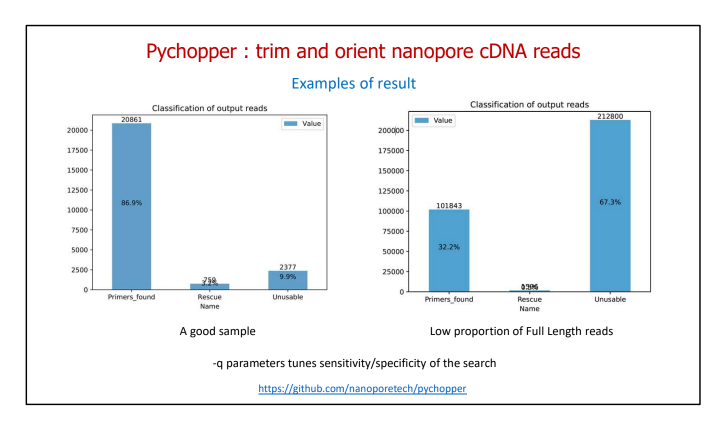
If you remember, initially raw RNAseq long reads should include strand switching primer (**SSP**) on one side and anchored oligo-dT primer (**VNP**) on the other.

The strand and orientation can be detected by the position and sequences (*direct* or *reverse-complement*) of these primers.

Because of the low quality of raw Nanopore reads (especially in old Nanopore data) **Pychopper** uses complex approximation algorithms to detect the primer sequences.

These are diagrams from the **Pychopper** web page, that supposedly should explain some of the employed algorithms.

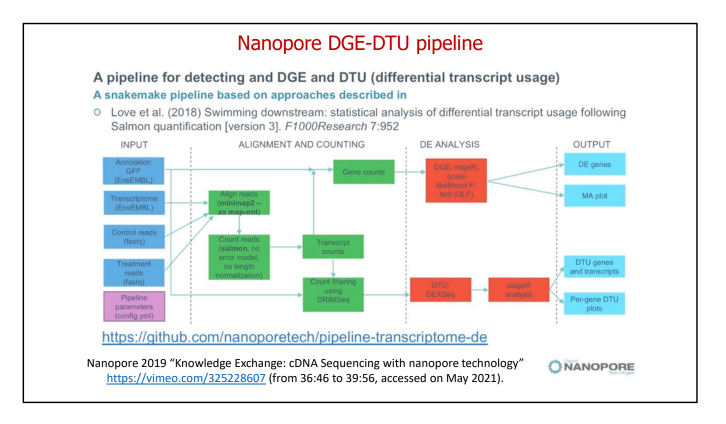
To be honest, I struggle to understand these diagrams, but you are welcome to try it yourself (the link to web page is provided :)



The good thing about **Pychopper** is that it provides informative plots about the preprocessing results.

Here you can see a good and a bad example.

The proportion of rejected reads may be tuned by –q parameter when running Pychopper (in the recent versions of Pychopper it is tuned using a sub-set of reads).



This slide illustrates the **DGE-DTU pipeline** that was recommended by Nanopore for quantification of transcript isoforms some years ago.

The pipeline started with properly oriented full-length reads prepared by Pychopper. Then the pipeline

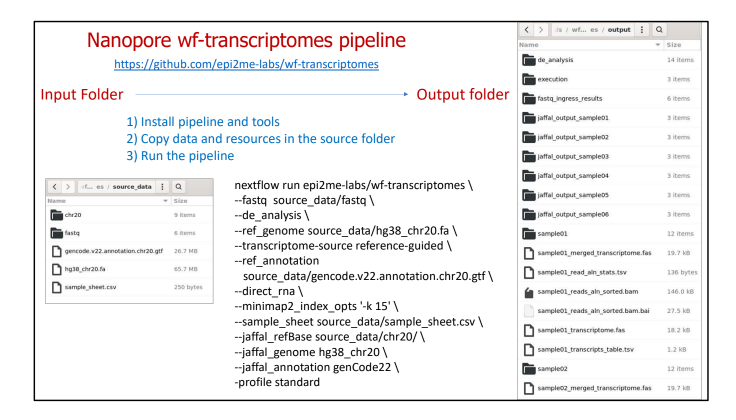
- Mapped reads to the known transcriptome with minimap2
- Passed mapped reads to Salmon for count
- Used these counts for DGE (Differential Gene Expression, edgeR) and
- Differential Transcript Usage analysis (DTU, DEXSeq & stageR)

All the required scripts were provided in the pipeline GitHub: https://github.com/nanoporetech/pipeline-transcriptome-de
The sequential steps included in the pipeline were orchestrated using the *Snakemake*

The sequential steps included in the pipeline were orchestrated using the **Snakemake** workflow manager.

The currently recommended version of this Nanopore pipeline is called **wf-transcriptomes**:

https://github.com/epi2me-labs/wf-transcriptomes
It implements the same steps, but uses the *Nextflow* workflow manager.
We will run the *wf-transcriptomes* pipeline during our practical session.



In practice, you first need to install the pipeline and tools following the authors' recommendations.

This step may require some IT knowledge, so this was already done on your VM.

Then, all what you need to is:

- Add your source data (FASTQ files, reference genome, etc)
- Start the pipeline

After the pipeline completes, the results will appear as many files and sub-folders in the output folder.

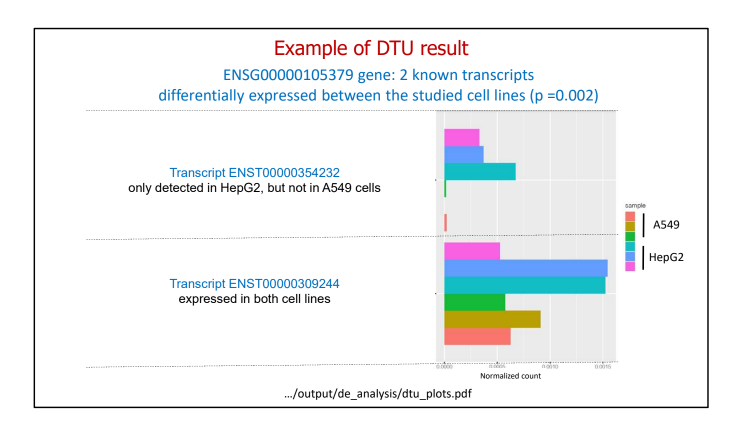
Nanopore aims to develop user-friendly GUI tool to install and run its pipelines: EPI2ME Desktop.

At the moment, Nanopore plans works on connecting EPI2ME Desktop to a cloud, because user's machines may not have enough resources to run the analysis.

A copy of EPI2ME Desktop was installed into your VM.

However, it is not yet intuitive enough to use.

So, you will run the pipeline using the provided script.



This is an example of DTU analysis produced by the pipeline. In wf-transcriptomes output folder you may find such plots .../output/de_analysis/dtu_plots.pdf file.

This example compares two cell lines: A549 and HepG2.

You can see that a specific gene (ETFB = ENSG00000105379) has two alternative transcripts.

One of them (ENST00000309244) is strongly expressed in both cell lines. While the other (ENST00000354232) is only expressed in HepG2.

The output folder contains much more results, including the outputs of every involved tool, such as FASTQs trimmed by Pychopper (if cDNA RNAseq was analyzed), BAM files produced by minimap2 etc. You will be able to explore the content of the **wf**-**transcriptomes** output folder during the practical session.

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Selected references

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Love M. et al **2018**: Swimming downstream: statistical analysis of differential transcript usage following Salmon quantification *F1000Research*, https://doi.org/10.12688/f1000research.15398.3

Practical session

| Data | Tools |
|--|--|
| Illumina, Nanopore and PacBio BAM files | IGV |
| Raw Nanopore FASTQ files | NanoPack: NanoPlot, NanoComp, chopper |
| (SG-Nex PCR-based cDNA sequencing data) | Pychopper |
| Slice of a Nanopore direct-RNA sequencing data (Chr20) Provided by Nanopore for testing wf-transcriptomes | wf-transcriptomes workflow |

Like in the Short-reads RNA-Seq practical session, it may be very intense for a person new to bioinformatics. You will be provided the detailed handouts and the fully-functional examples of scripts.

Samples are selected by size!

Don't over-interpret data quality and biology:)