SARS-CoV-2 NGS Training

NGS Academy for the Africa Pathogen Genomics Initiative

Module 4 - Illumina workflow





SARS-CoV-2 NGS Training

Module 4: Illumina workflow

- Session 1: Concepts, library prep, starting a sequencing run
- Session 2: Library prep QC and sequencing run QC
- Session 3: IDSeq/Data Processing





Cristina Tato, Director Rapid Response (RR) group



Manu Vanaerschot, Scientist RR group



Vida Ahyong, Scientist RR group





Katrina Kalantar, Computational Biologist

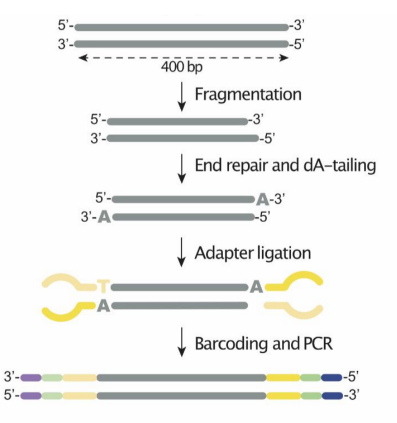


Liz Fahsbender, IDseq Application Scientist

Library prep

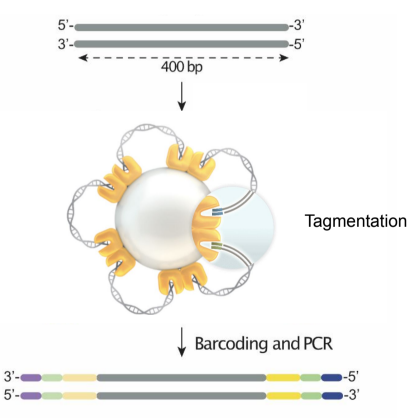
Ligation protocol

Illumina TruSeq DNA / NEBNext Ultra II DNA



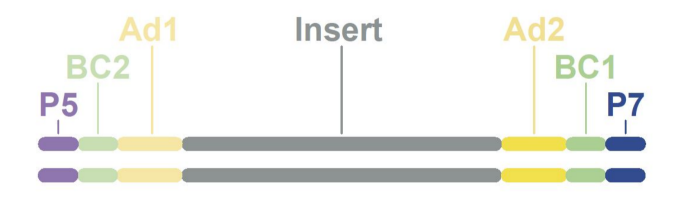
Tagmentation protocol

Illumina DNA prep (= Nextera Flex)



OR

Library prep - final library



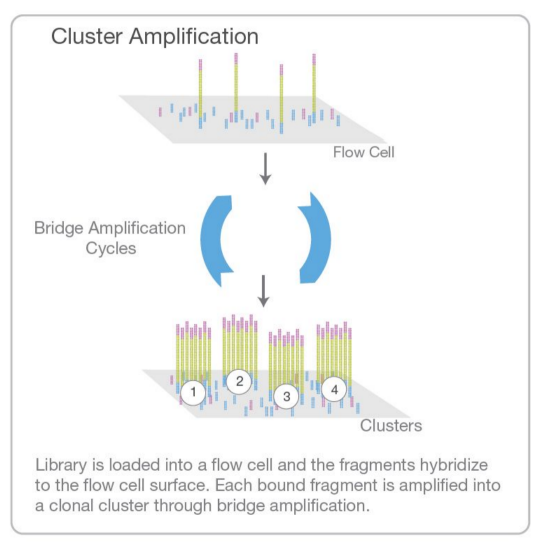
Insert = DNA fragment to be sequenced

Ad1/Ad2 = Adapters that allow initiation of sequencing

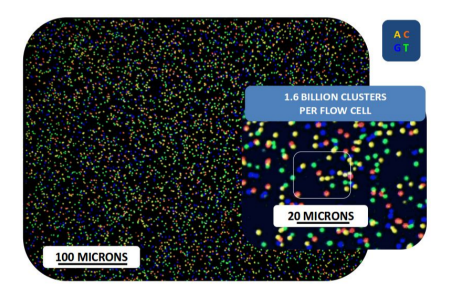
BC1/BC2 = Barcodes specific for each sample (one for each strand)

P5/P7 = Flow cell binding sites

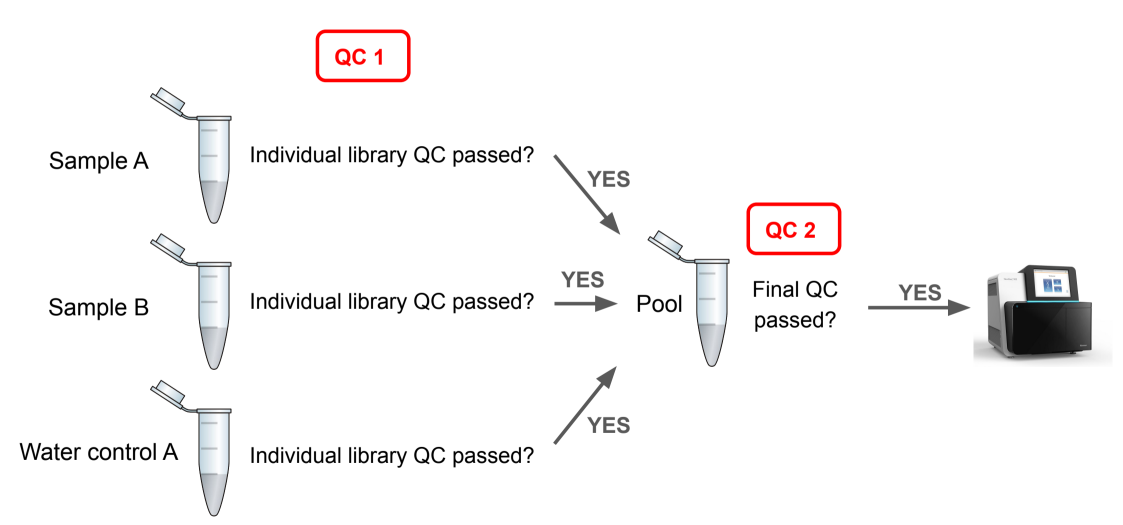
Cluster amplification is essential for Illumina sequencing



- ~ loading concentration
- ~ fragment length



Library prep QC



Library prep QC

Concentration

vs

Length

Remove adapter dimers

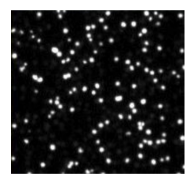
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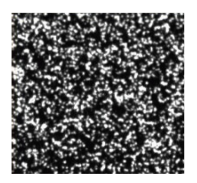
VS

or

Why is an accurate concentration important?

- Individual samples: to pool equal amounts of different samples together
 - Aim for an even amount of sequencing data per sample
- Final library: sequencer requires loading with a very precise concentration
 - o If concentration is overestimated (actual conc. is lower than what you think)
 - Underloading and underclustering of sequencing run
 - Sequencing run will yield less sequencing data than usual
 - If concentration is underestimated (actual conc. is higher than what you think)
 - Overloading and overclustering of sequencing run
 - Sequencing run will suffer from higher error rates and reduced data output





DO NOT USE to quantify libraries

Spectrophotometry (e.g. Nanodrop)



- DMA absorbs light most strongly at 260 nm
- Level of absorbance ~ concentration of DNA

BUT other molecules (RNA, proteins) & contaminants also absorb light at 260 nm, causing up to 10-fold **overestimation** of the DNA concentration!

Fluorometry

(e.g. Qubit, qPCR)





- Detects fluorescent dyes that only bind DNA
- Most reliable method for NGS

Electrophoresis

(e.g. Tapestation, Bioanalyzer)



- Compare intensity of DNA bands to standards
- Especially useful to determine concentration in certain fragment length range

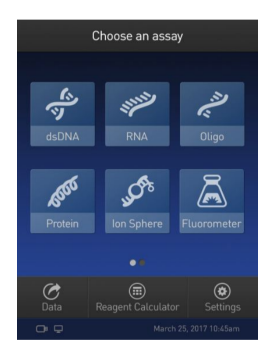
Qubit dsDNA <u>High Sensitivity</u> (HS) Assay kit



- dsDNA HS Reagent (200x concentrate)
- dsDNA HS Buffer
- dsDNA HS Standard #1
- dsDNA HS Standard #2

- 1. Prepare Working Solution:
 - Diluting HS reagent 1:200 in HS buffer
- 2. Dilute in Working Solution:
 - Standards: 10 μl standard + 190 μl Working Solution
 - Samples: 1 μl sample + 199 μl Working Solution
- 3. Vortex 2-3 seconds before running standards/samples

Qubit dsDNA High Sensitivity (HS) Assay kit



RFU Values
Standard 1 Standard 2
246.23 25454.58

Standard 2
0 100 200 300 400 500
Qubit* 3.0 tube concentration [ng/mL]





Select dsDNA HS workflow

Run the 2 standards

Enter sample volume added to tube

Run sample

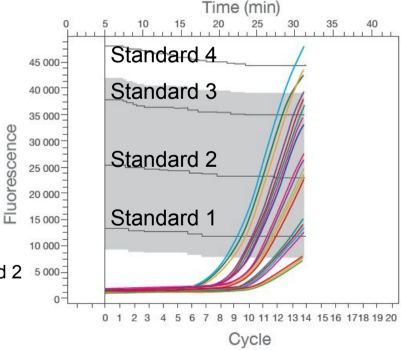
Top value = original conc.

Bottom value = diluted conc.

- What if the concentration is too low?
 - In future runs, consider adding more cycles to the indexing PCR step
 - Amplify library using P5/P7 primers and qPCR-based library amplification kit
 - e.g. Roche KK2702
- 1. Set up qPCR reaction: library + mastermix + P5/P7 primers
- 2. Treat standards as separate samples (do not add mastermix)
- 3. Start run

Temperature	Time	Cycles	
98°C	45 sec	1	
98°C	15 sec	25	
63°C	30 sec		
72°C	1m 45 sec		
Plate r	*		
72°C	20 sec		

- 4. Pause cycler <u>during the 20s at 72°C step</u> when a sample crossed the Standard 2 5000line, remove the sample and continue amplification for other samples
- 5. After amplification, perform bead cleanup on amplified libraries and QC again



Library prep QC

Concentration

VS

Length

VS

or

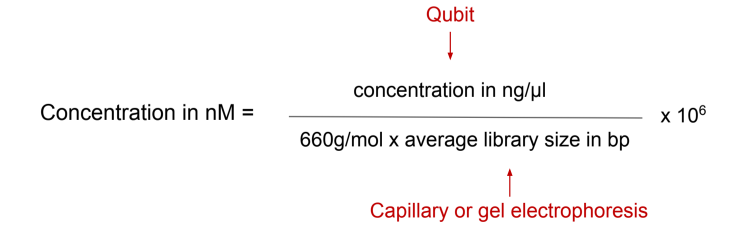
Remove adapter dimers

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Library prep QC - Length

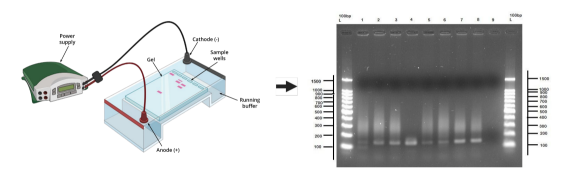
Why is it important to assess fragment length in a library?

- Long fragments (>1.2 kb) do not cluster on Illumina flow cells and are therefore not sequenced
- Short fragments are preferentially amplified (better clustering) and will thus be preferentially sequenced
- Fragment length distribution impacts calculations to go from ng/µl (Qubit) to nM for loading concentrations

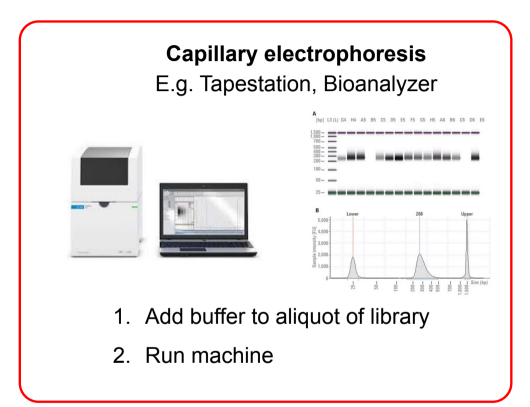


Library prep QC - Length

Gel electrophoresis



- PCR amplify an aliquot of the library using universal P5/P7 Illumina primers
- 2. Gel electrophoresis of ladder and samples
 - Takes ~ 4 hrs

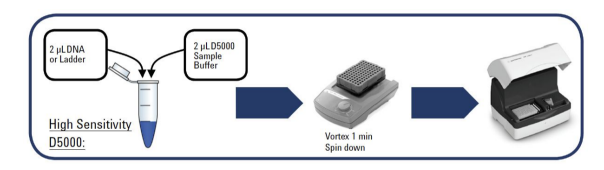


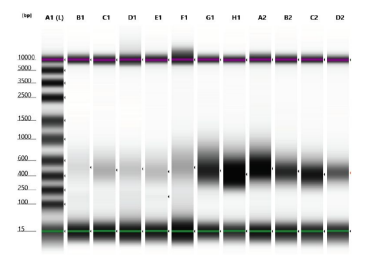
- Tapestation ~ 10 minutes (depending on # samples)
- Bioanalyzer ~ 45 minutes
- No PCR bias
- Can calculate concentration in nM

Library prep QC - Length

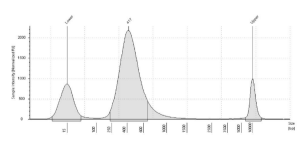
Tapestation

High Sensitivity D1000 or D5000 kits (1kb or 5kb detection limit)





Peak: H1



Location
Concentration
Description
Observations

H1
1470

Size [bp]	Calibrated Conc. [pg/µl]	Assigned Conc. [pg/µl]	PeakMolarity [pmol/l]	% Integrated Area	PeakComment	Observations
15	346	-	35500	-		Lower Marker
417	1470	1	5430	100.00		
10000	180	180	27.7	-		Upper Marker

Library prep QC

Concentration Length

vs vs

or

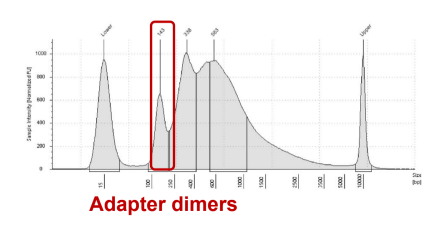
Remove adapter dimers

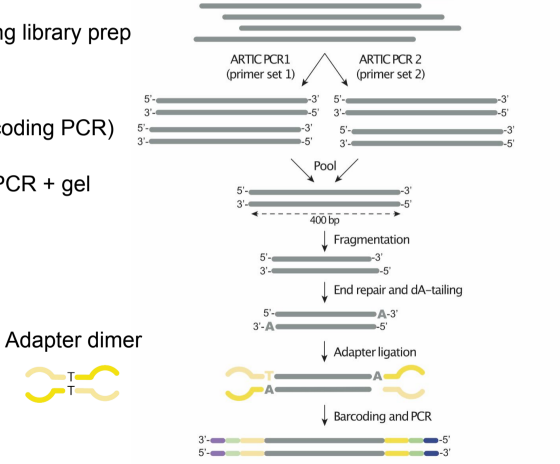
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RNA DNA

Reverse transcription with random primers

- Adapter dimers are:
 - A byproduct of the adapter ligation step during library prep (TruSeq/NEBNext Ultra II DNA kits)
 - ± 140 bp long at the final QC step (after barcoding PCR)
 - Use capillary electrophoresis (preferred) or PCR + gel electrophoresis to detect adapter dimers



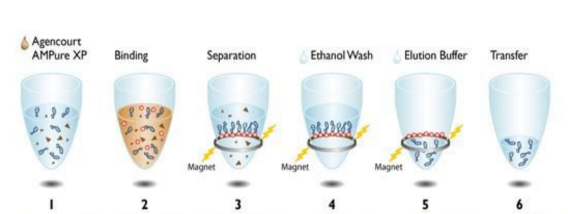


Library prep QC - Clean up adapter dimers

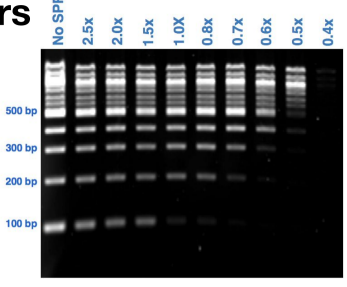
AMPure XP SPRI select beads (paramagnetic)

or homemade SPRI beads (>10x cheaper)

- The longer a fragment, the better it will bind to the beads
- The fewer beads you add, the more short fragments you'll lose
- Perform size selection using these beads to remove adapter dimers



1. PCR reaction 2. Binding of PCR amplicons to magnetic beads 3. Separation of PCR amplicons bound to magnetic beads from contaminants 4. Washing of PCR amplicons with Ethanol 5. Elution of PCR amplicons from the magnetic particles 6. Transfer away from the beads into a new plate



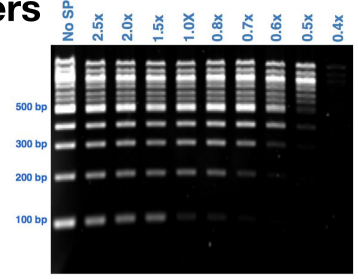
Example: for a 1.8x SPRI clean-up:

Add 18 μl of SPRI beads to
 10 μl of sample

Library prep QC - Clean up adapter dimers

AMPure XP SPRI select beads (paramagnetic)

- Size selection using beads to remove adapter dimers
- Rule of thumb: if you see a clear trace of adapter dimers, performs another round of cleanup

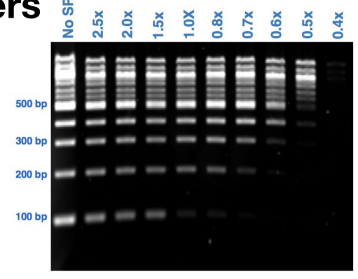


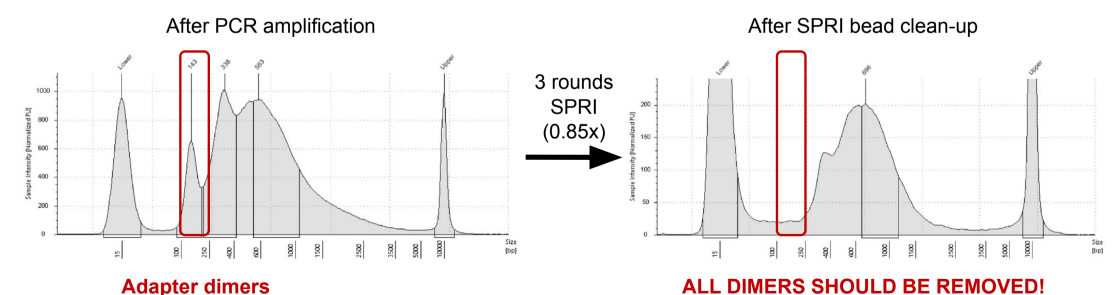
Step	To remove	SPRI ratio
After adapter ligation	Adapter dimers (64 bp) & reagents	0.9x
After P7/P5 PCR	P5-adapter dimer-P7 (143 bp) & reagents	0.8x
QC of library	P5-Adapter dimer-P7 (143 bp)	0.8x

Library prep QC - Clean up adapter dimers

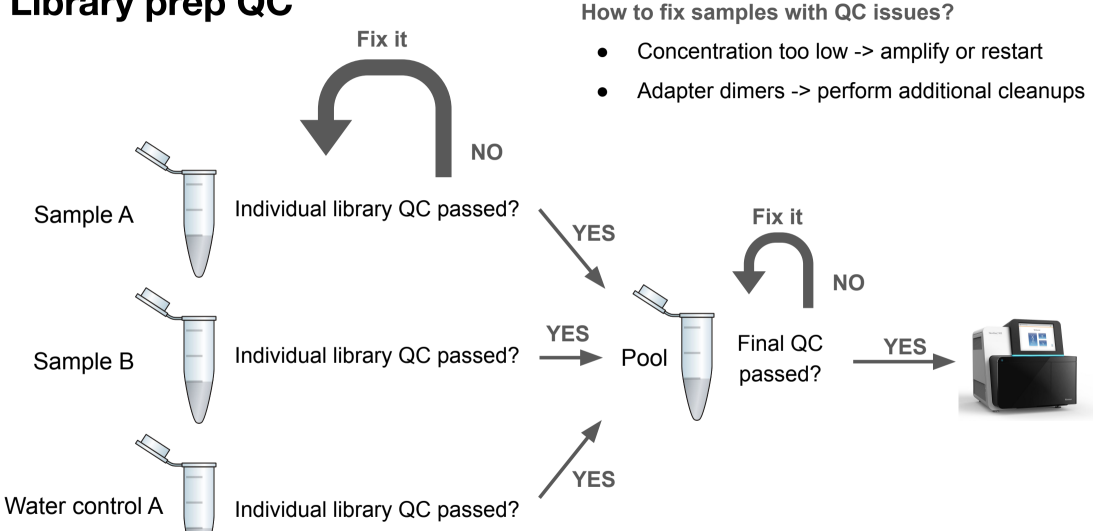
AMPure XP SPRI select beads (paramagnetic)

- Size selection using SPRI beads to remove adapter dimers
- Rule of thumb: if you see a clear trace of adapter dimers, performs another round of cleanup





Library prep QC

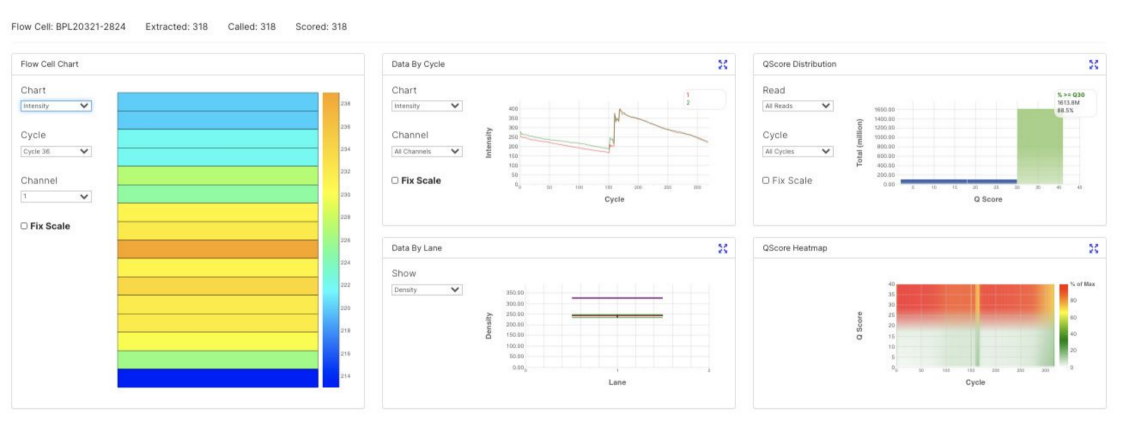


PhiX control

- PhiX is a control library that can be spiked into your library right before loading the sequencer
 - PhiX is a bacteriophage, single stranded DNA genome of 5386 -> known sequence
 - PhiX spike-in comes already fragmented and adapter-ligated -> ready to sequence
 - Spike-in concentration usually 5%

- Advantages of using PhiX:
 - Enables calculation of error rate
 - If a sequencing run fails, Illumina will evaluate PhiX data to assess whether or not the kit was the issue (and send a replacement if so)

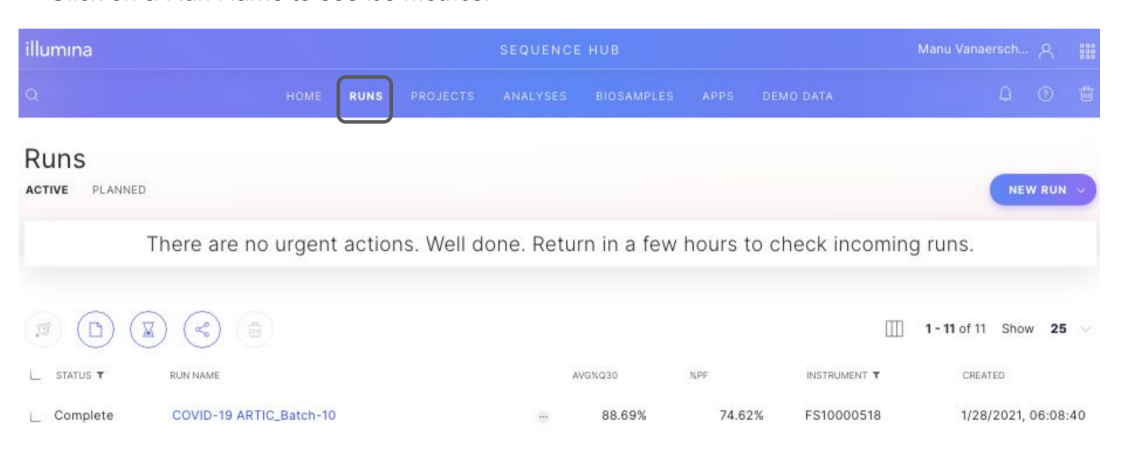
Sequencing Run QC



Did you get the expected amount of sequencing data? If not, why?

Sequencing Run QC using BaseSpace

After logging into BaseSpace, Click on 'Runs' to view a list of all your runs. Click on a Run Name to see it's metrics.



Sequencing Run QC using BaseSpace

COVID-19 ARTIC_Batch-10

SUMMARY BIOSAMPLES FILES INDEXING QC SAMPLE SHEET Instrument Instrument Type Created 74.62 88.70 FS10000518 2021-01-28 06:40 iSeq100 File Count/Size File Status 11,353 files (1 GB) Active Run Status Lane QC Status Flow Cell Status **QcPassed QcPassed** Complete Owner User Flow Cell ID Run ID Yield Latest Analysis Cycles BPL20321-2824 20210128 FS10000... 151 | 8 | 8 | 151 1.80 Gbp

AVG%Q30 and %PF



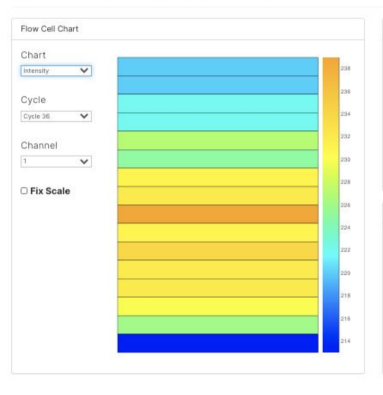
- AVG%Q30 = percentage of reads that have bases with an average quality score >Q30
 - Ideally AVG%Q30 >90%, but >70% is good too
 - If <70%: evaluate why the quality is lower than expected. Possible reasons:
 - Overclustering: too much material was loaded on the flow cell
 - Adapter dimers: these will yield reads with only ~70bp of good signal, and the rest low quality 'nothingness'.
- %PF = percentage of passed filter reads = reads based on a cluster of a single molecule
 - Ideally >70%, but >50% is good too
 - If <50%, check for indications of overclustering

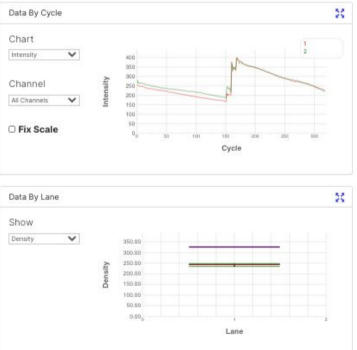
QC charts

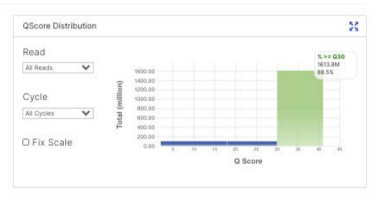
Run: COVID-19 ARTIC_Batch-10: Charts

SUMMARY BIOSAMPLES CHARTS METRICS INDEXING QC SAMPLE SHEET FILES

Flow Cell: BPL20321-2824 Extracted: 318 Called: 318 Scored: 318

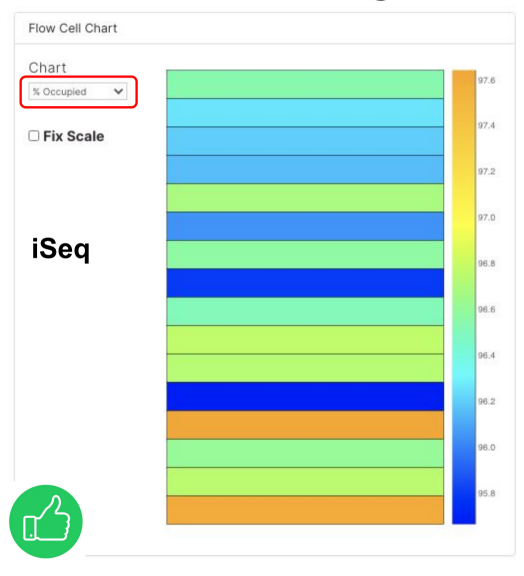








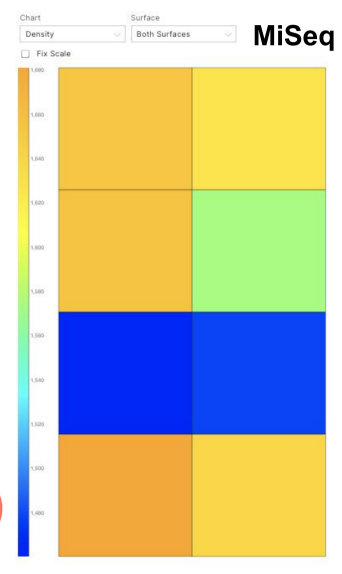
Evaluate clustering for iSeq



Flow Cell Chart: select % occupied under 'Chart'

- -> Signs of overclusterig or underclustering?
 - Best range for values is 90-98%
 - >98%: possible overclustering, which will reduce %PF and AVG%Q30 scores. Review library quantification data to check for errors or consider reducing the loading quantity for the next sequencing runs.
 - <90%: Review library quantification data to check for errors or consider increasing the loading concentration for the next sequencing runs to get more reads from a run.

Evaluate clustering for MiSeq



Flow Cell Chart: select Density under 'Chart'

Optimal Raw cluster density MiSeq:

V2 kits: 1000-1200 K/mm²
 V3 kits: 1200-1400 K/mm²

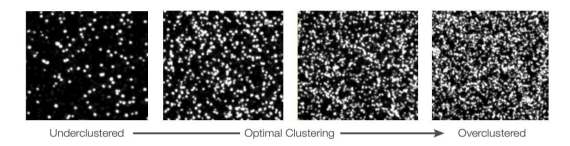
This V3 example shows a raw cluster density ~1600 K/mm²

- -> The run was overclustered!
 - -> lower %PF reads
 - -> lower yield of PF reads total

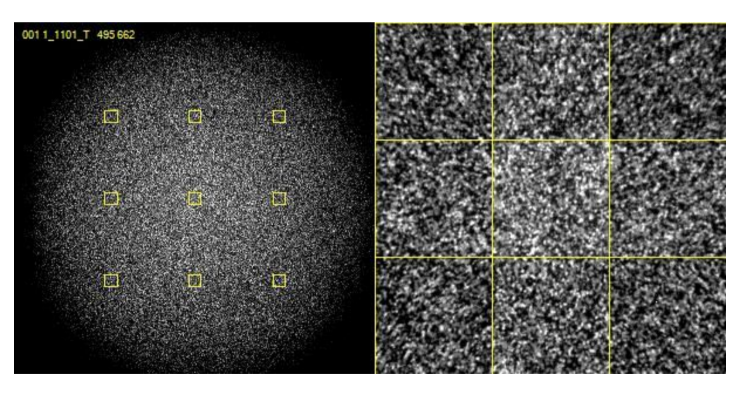
Check QC data to see why concentration in nM of final library was underestimated (length overestimated?)



Evaluate clustering by looking at Files/Thumbnail_Images

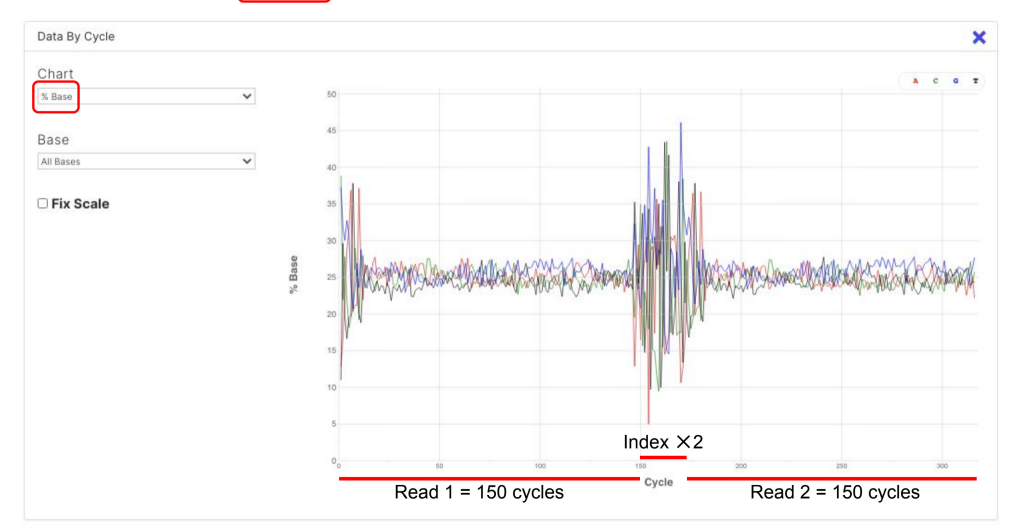


Illumina examples from here



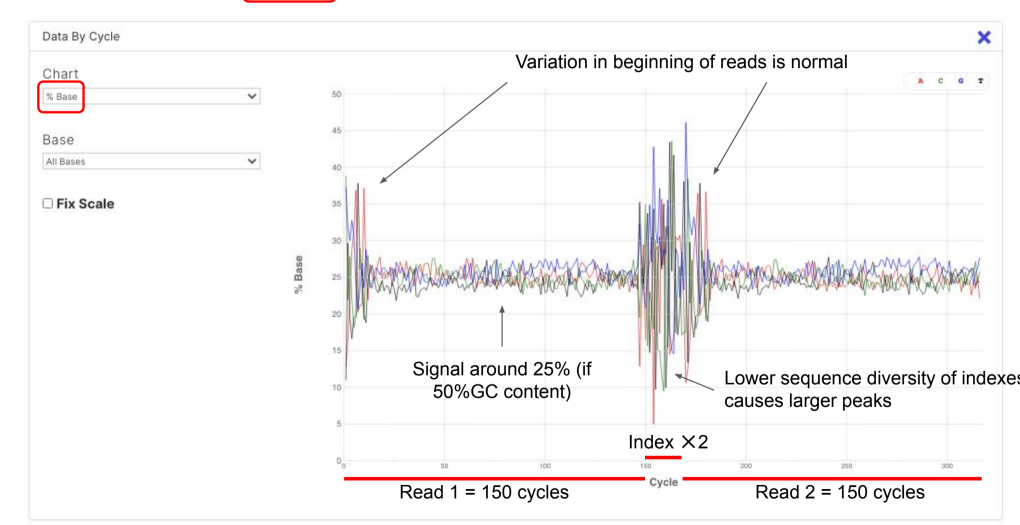
Tiles also show overclustering of this MiSeq run

Data by cycle chart: select % Base under 'Chart' -> Adapter dimers? Short fragments?

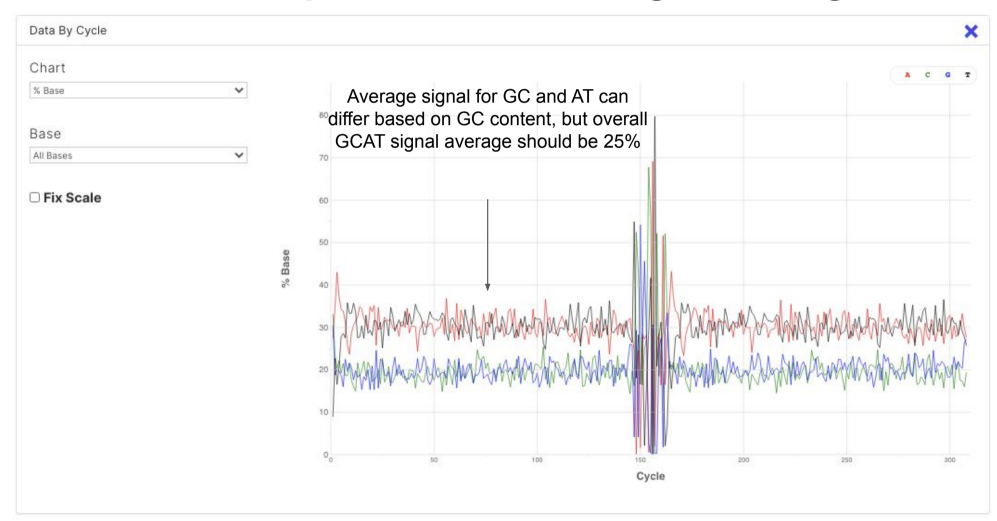




Data by cycle chart: select % Base under 'Chart' -> Adapter dimers? Short fragments?



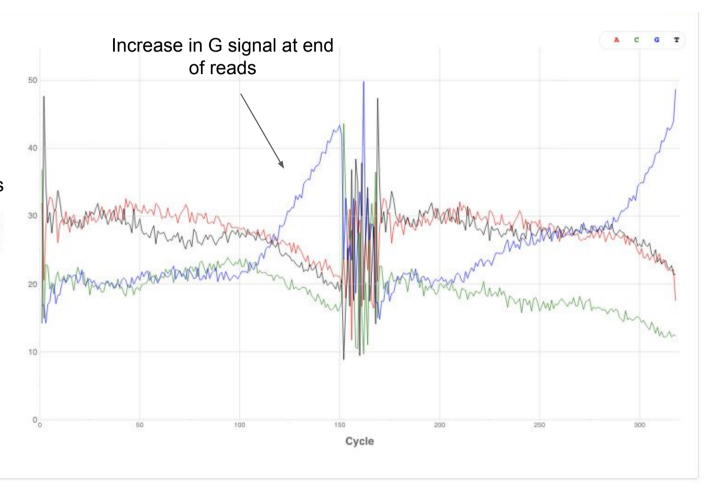






Issue 1 - short reads: Increase in G signal at end of reads

- G is the "dark base" is iSeq, MiniSeq,
 NextSeq and NovaSeq chemistry.
- Continuous increase in G indicates no bases detected for subsequent cycles
 - Inserts shorter than 150 bp?
 - Issue with elongation of strands during clustering?
- If insert size was indeed shorter than usual (check lib prep QC), reduce fragmentation time for future!

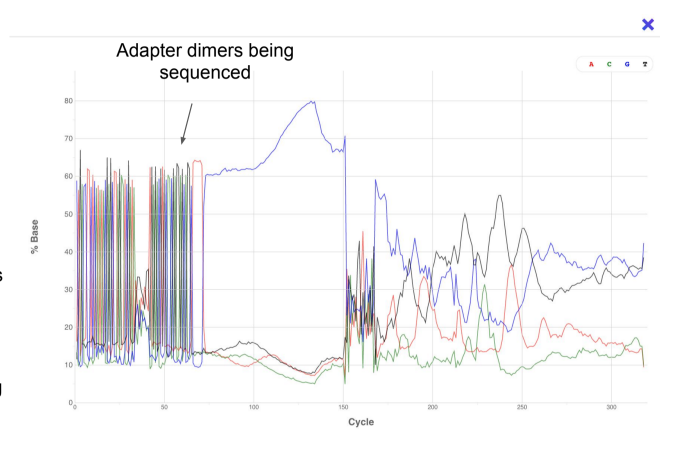


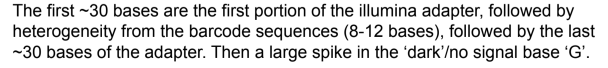


Note: MiSeq and HiSeq would call an A in case nothing is detected).

Issue 2 - adapter dimers: Large peaks for first 70 cycles, and then an increase in G signal (iSeq, MiniSeq, NextSeq, NovaSeq)

- Adapter dimers will yield reads with low sequence variation (large peaks) that are about 70 bp long
- Adapter dimers are short fragments and thus get preferentially sequenced, taking up valuable sequencing space
- Crucial to remove ALL adapter dimers during library prep QC !!

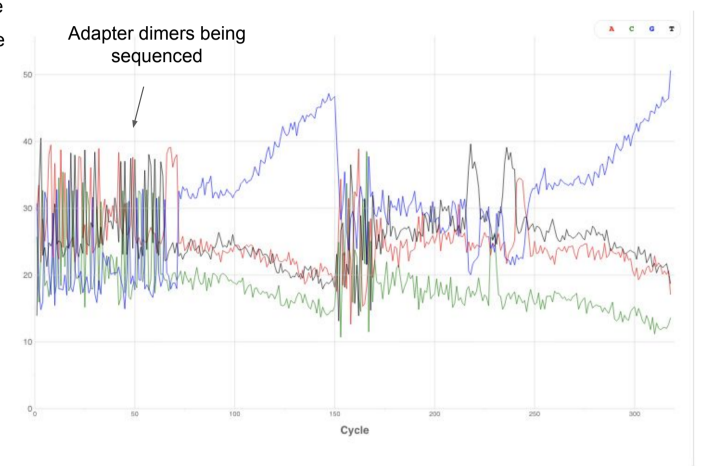






Issue 3 - adapter dimers and short reads: Large peaks for first 70 cycles followed by more stable signal and then a steady increase in G signal

- Adapter dimers will yield reads with low sequence variation (large peaks) that are about 70 bp long
- Adapter dimers are short fragments and thus get preferentially sequenced, taking up valuable sequencing space
- Crucial to remove ALL adapter dimers during library prep QC !!





Evaluate error rate based on PhiX

COVID-19 ARTIC_Batch-10

METRICS

INDEXING QC

SAMPLE SHEET

FILES



SUMMARY









Per Read Metrics

READ	CYCLES	YIELD	PROJECTED YIELD	ALIGNED (%)	ERROR RATE (%)	INTENSITY CYCLE 1	%>Q30
Read 1	151	860.56 Mbp	860.56 Mbp	2.72	0.45	259.25	90.79
Read 2 (I)	8	40.16 Mbp	40.16 Mbp	0.00	0.00	210.56	87.55
Read 3 (I)	8	40.16 Mbp	40.16 Mbp	0.00	0.00	376.44	74.09
Read 4	151	860.56 Mbp	860.56 Mbp	2.43	1.09	390.75	87.34
Non-index Reads Total	302	1.72 Gbp	1.72 Gbp	2.58	0.77	325.00	89.06
Total	318	1.80 Gbp	1.80 Gbp	2.58	0.77	309.25	88.69

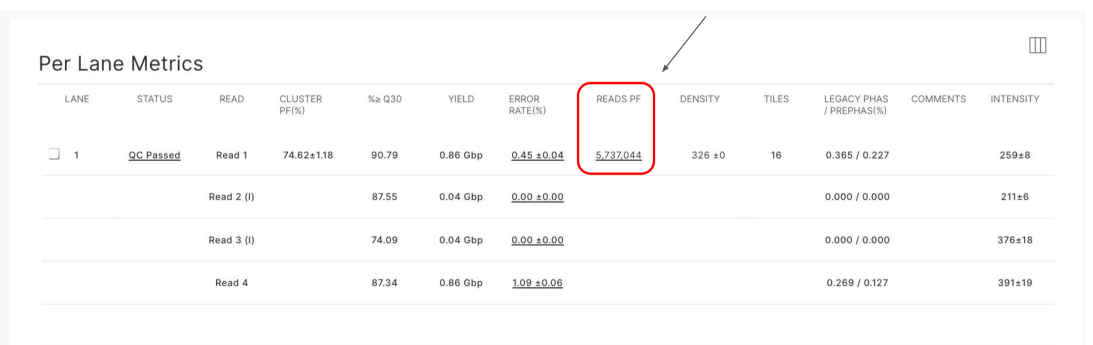
Evaluate sequencing data yield

COVID-19 ARTIC_Batch-10

SUMMARY BIOSAMPLES CHARTS METRICS INDEXING QC SAMPLE SHEET FILES

Scroll down to 'Per Lane Metrics'

Number of paired end reads that were generated



On the 'Metrics' page (if using PhiX)

- Per Read Metrics:
 - a. **% Aligned** (PhiX): should be close to % of PhiX that you spiked in.
 - i. If it's higher, PhiX outcompeted your library, meaning you put less library in than you thought (double check your pooling and quantification calculations).
 - ii. If it's lower than expected, small fragments like adapter dimers might have outcompeted PhiX AND/OR you underestimated the quantity of material in your sample (added way more than you thought, double check your quantification data).
 - b. **Error rate** % (PhiX): should be less than 1%.
- Per Lane Metrics:
 - c. **Reads PF**: number of passed filter reads that were generated in this run. This should be close to (or over) the minimum number of guaranteed reads that your sequencer is advertised to generate. If lower, you probably under- or overclustered. Check your quantification calculations and adjust the loading concentration if needed.

Evaluate demultiplexing

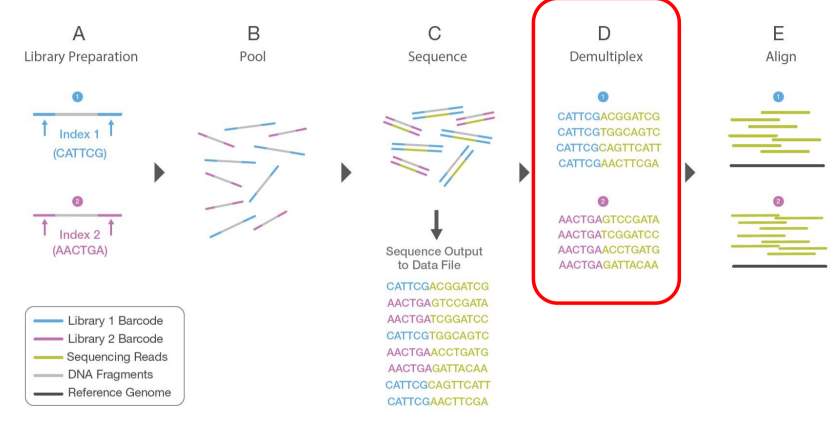
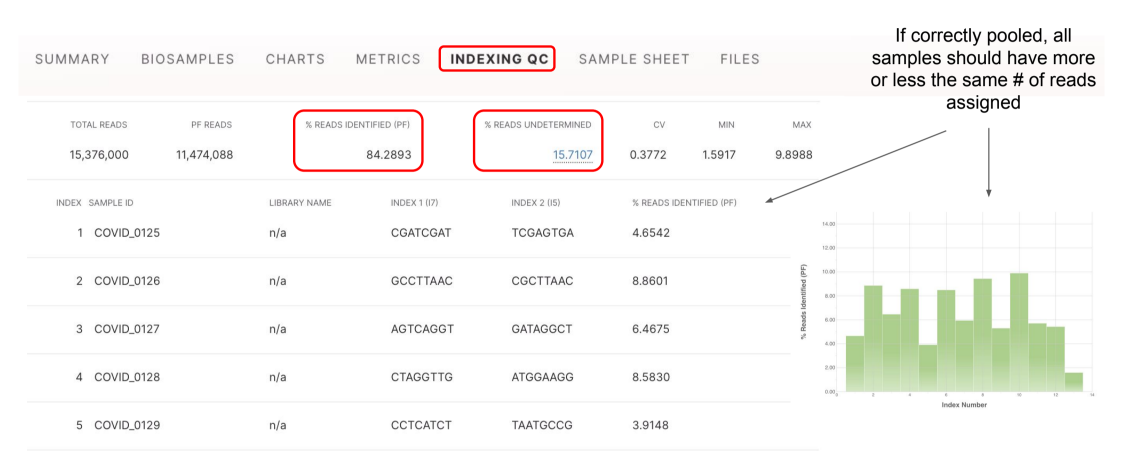


Figure 5: Library Multiplexing Overview.

- a. Two distinct libraries are attached to unique index sequences. Index sequences are attached during library preparation.
- b. Libraries are pooled together and loaded into the same flow cell lane.
- c. Libraries are sequenced together during a single instrument run. All sequences are exported to a single output file.
- d. A demultiplexing algorithm sorts the reads into different files according to their indexes.
- e. Each set of reads is aligned to the appropriate reference sequence.

Adapted from Illumina.com

Evaluate demultiplexing



...

Evaluate demultiplexing

On the 'Indexing QC' page:

- a. **% read identified** = reads with identified barcodes: should be >80% (the higher the better). If this % is low, you should check to make sure your sample sheet assigned the correct index/barcode sequences for each sample.
- b. **% reads undetermined** = reads without an associated known barcode. If this % is high, you should check to make sure your sample sheet was correct.
- c. **PF reads**: this number details the amount of (unpaired) single end reads. This number is always double than your true amount of paired-end reads shown on the 'Metrics' page.

Troubleshoot demultiplexing

- A high % of undetermined reads and missing samples (samples with no reads assigned) indicate an issue with demultiplexing!
 - Double check the sample sheet to verify if the correct indices/barcodes were assigned to each sample.
 - Look at the barcodes sequences identified for 'undetermined reads' listed in DemuxSummaryF1L1.txt, if there is barcode combination (paired end) that occured in a high number of reads, it might be an unidentified sample.

Info on finding troubleshooting and finding the demux file in BaseSpace here
How to edit sample sheet and requeue in Basespace here
Info on finding troubleshooting and finding the demux file using Local Run Manager here
Info on finding troubleshooting and finding the demux file using MiSeq Reporter here
How to edit sample sheet and requeue using MiSeq Reporter here

Summary library and sequencing run QC

Concentration

- Accurate measurements are crucial, use fluorometry-based method (e.g. Qubit),
 - do NOT use spectrophotometry (e.g. no Nanodrop)
- Make sure values are within dynamic range of assay
- If too low, amplify library if needed, or adjust # PCR cycles of indexing PCR step for future runs

Library length

- Remove all adapter dimers to prevent them taking up sequencing space
- Capillary electrophoresis is the fastest and most accurate method
- PCR + gel electrophoresis is an alternative method if capillary electroph. is not available

Sequencing run QC:

- Optimal loading concentration will yield maximum amount and quality of data
- Monitor sequencing runs and adjust loading concentration of future runs as needed