

## Introduction to NGS

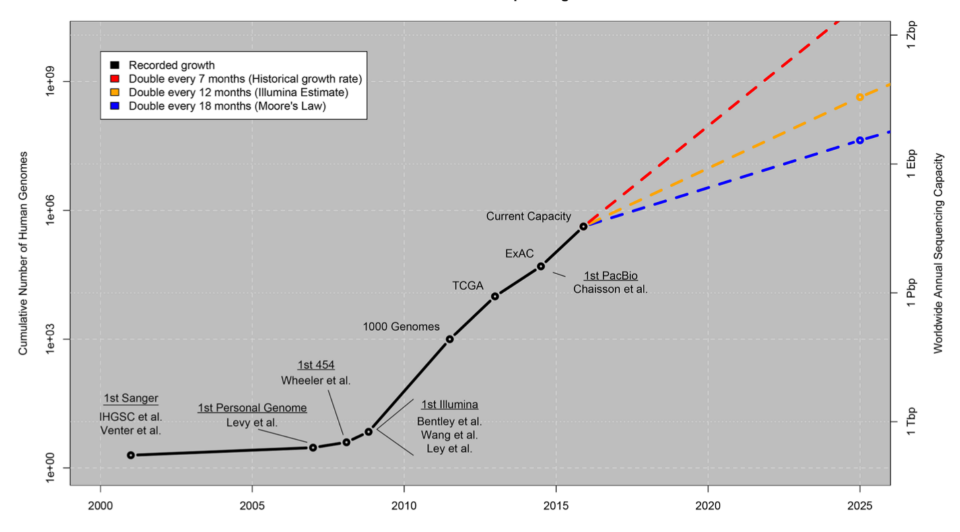
**Hubert Rehrauer** 





#### **NGS Data Increase**

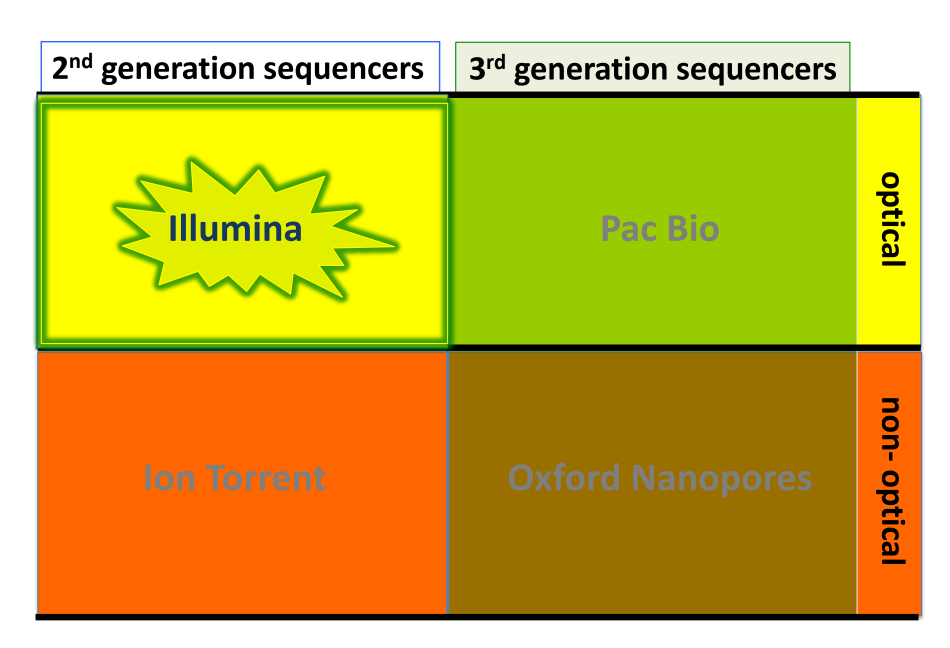
#### **Growth of DNA Sequencing**



NGS data increases faster than computer speed

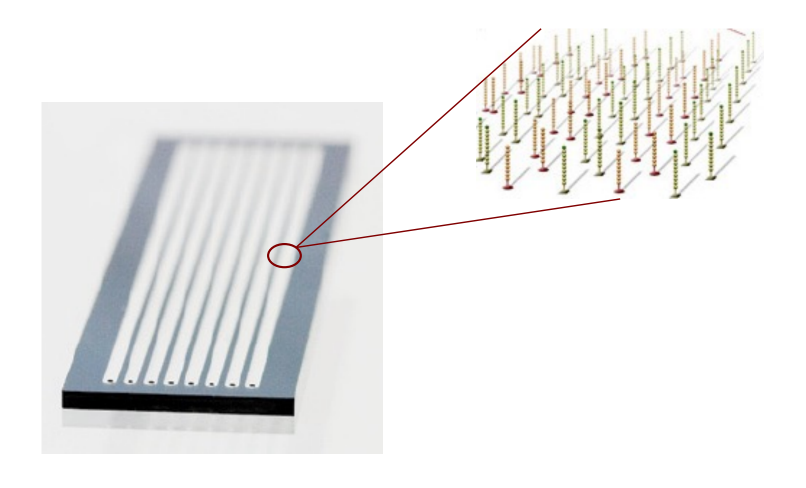
#### Ingredients for the success

- Evolution has yielded DNA and RNA molecules for information storage and transfer. They have good properties to be read (measured)
- NGS technologies rely on
  - massive parallelization
  - measurement process is done by individual molecules (cheap and fast)

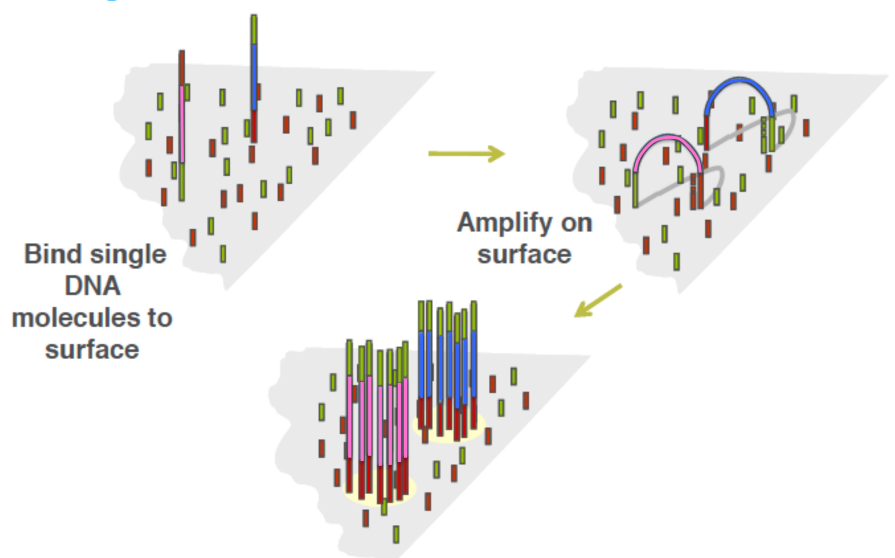


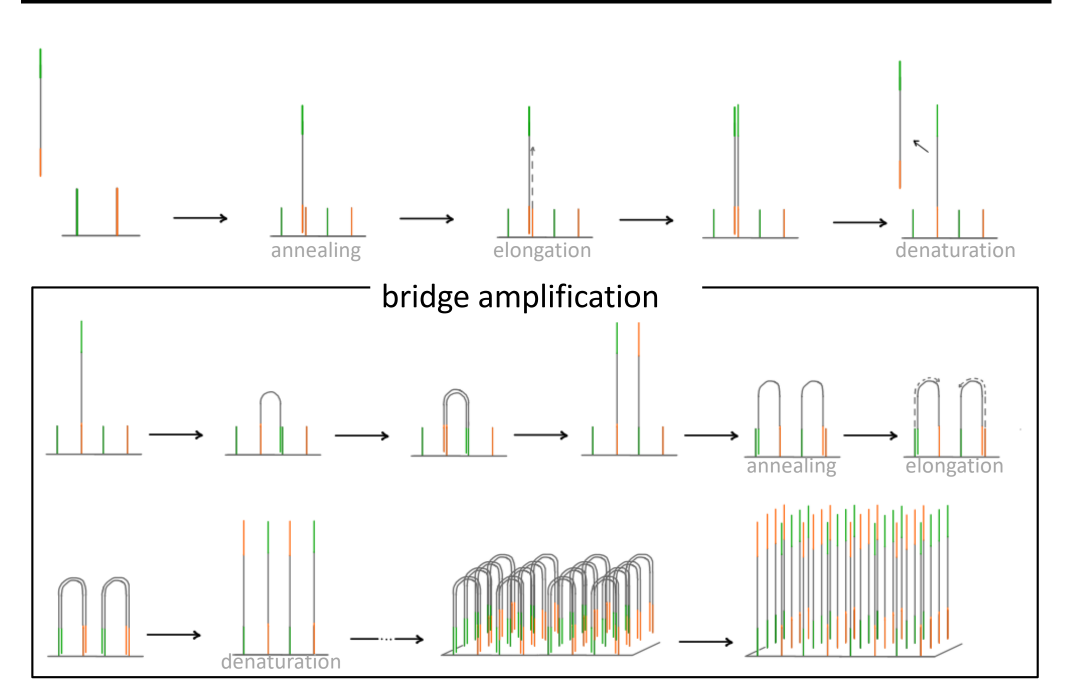


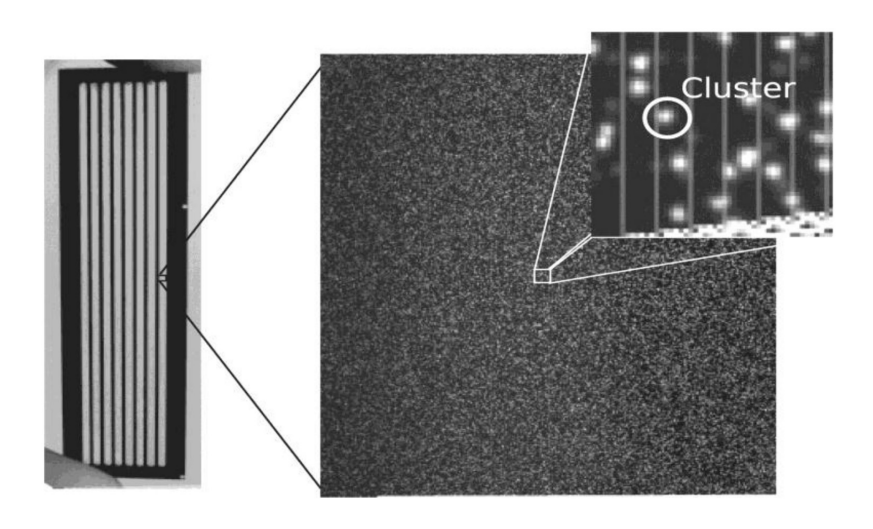
## **Illumina Flow cell**



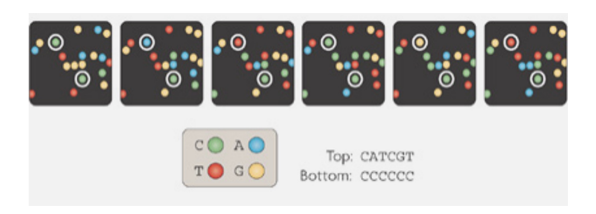
#### **Cluster generation overview**

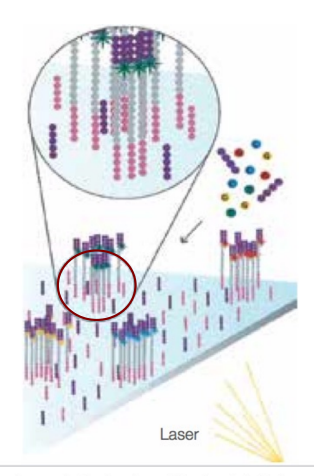






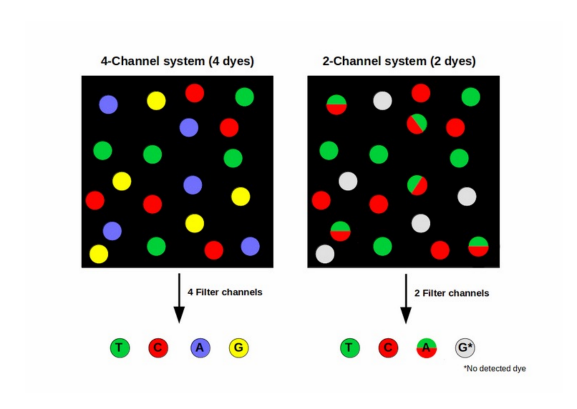
### **Illumina Sequencing**

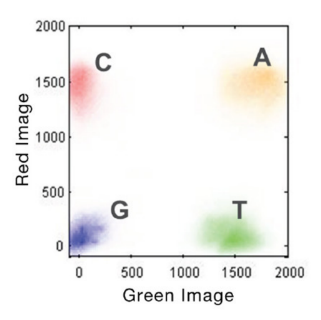




The first sequencing cycle begins by adding four labeled reversible terminators, primers, and DNA polymerase.

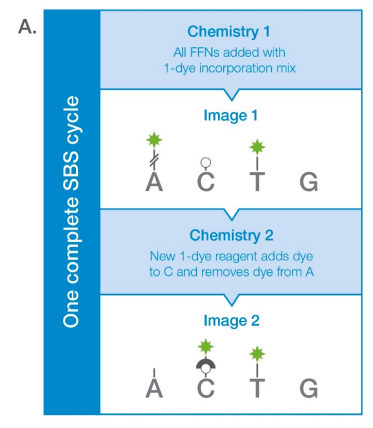
### **Color coding of bases**







### Color coding of illumina iSeq



| 3. | Image 1 | Image 2 | Result |
|----|---------|---------|--------|
|    | ON      | OFF     | А      |
|    | OFF     | ON      | С      |
|    | ON      | ON      | Т      |
|    | OFF     | OFF     | G      |

### functional genomics center zurich

## Phred scores measure base call accuracy

- P
- error probability of a given base call
- Q
- -10log<sub>10</sub>P
- Assign to each base
- Range from 0-41



| Phred Quality<br>Score | Probability of<br>Incorrect Base Call | Base Call<br>Accuracy |  |  |  |  |  |
|------------------------|---------------------------------------|-----------------------|--|--|--|--|--|
| 10                     | 1 in 10                               | 90%                   |  |  |  |  |  |
| 20                     | 1 in 100                              | 99%                   |  |  |  |  |  |
| 30                     | 1 in 1,000                            | 99.9%                 |  |  |  |  |  |
| 40                     | 1 in 10,000                           | 99.99%                |  |  |  |  |  |
| 50                     | 1 in 100,000                          | 99.999%               |  |  |  |  |  |

Ewing B, Green P. 1998. Genome Res. 8(3):186-194.

http://en.wikipedia.org/wiki/Phred\_quality\_score

## Phred scores are stored with sequences

- FASTQ
  - 4 lines:
    - 1. Header line for Read (starts with "@" and the sequence ID)
    - 2. Sequence
    - 3. Header line for Qualities (starts with "+")
    - 4. Quality score (represented in ASCII format)

#### Phred scores can be ASCII encoded

- Add an offset and convert the sum to ASCII
- Current format
  - Illumina 1.9 ( i.e. Sanger format)
  - Phred scoring: 0-41;
  - Offset: 33
  - 41+33=74 (J)
  - All current sequencers

| Dec | H) | Oct | Chai | r                        | Dec | Нх | Oct | Html   | Chr   | Dec | Нх | Oct | Html  | Chr | Dec | Нх | Oct | Html Ch | <u>ır</u> |
|-----|----|-----|------|--------------------------|-----|----|-----|--|-------|-----|----|-----|-------|-----|-----|----|-----|---------|-----------|
| 0   | 0  | 000 | NUL  | (null)                   | 32  | 20 | 040 | @#32;  | Space | 64  | 40 | 100 | a#64; | 0   | 96  | 60 | 140 | a#96;   | 8         |
| 1   | 1  | 001 | SOH  | (start of heading)       | 33  | 21 | 041 | @#33;  | 1     | 65  | 41 | 101 | @#65; | A   | 97  | 61 | 141 | a#97;   | a         |
| 2   | 2  | 002 | STX  | (start of text)          | 34  | 22 | 042 | @#3 <b>4</b> ;   | **    | 66  | 42 | 102 | B     | В   | 98  | 62 | 142 | a#98;   | b         |
| 3   | 3  | 003 | ETX  | (end of text)            | 35  | 23 | 043 | #  | #     |     |    |     | C     |     |     |    |     | c       |           |
| 4   | 4  | 004 | EOT  | (end of transmission)    |     |    |     | <b>4#36</b> ;  |       |     |    |     | 4#68; |     |     |    |     | @#100;  |           |
| 5   | 5  | 005 | ENQ  | (enquiry)                |     |    |     | %  |       |     |    |     | E     |     |     |    |     | e       |           |
| 6   |    |     |      | (acknowledge)            |     |    |     | &  |       |     |    |     | F     |     |     |    |     | f       |           |
| 7   |    |     |      | (bell)                   |     | _  |     | '  |       | 71  |    |     | G     |     |     | _  |     | g       |           |
| 8   |    | 010 |      | (backspace)              |     |    |     | &# <b>4</b> 0;   |       |     |    |     | H     |     |     |    |     | h       |           |
| 9   |    |     |      | (horizontal tab)         |     |    |     | @#41;  |       |     |    |     | 6#73; |     |     |    |     | a#105;  |           |
| 10  |    | 012 |      | (NL line feed, new line) |     |    |     | &#<b>4</b>2;</td><td></td><td></td><td></td><td></td><td>6#74;</td><td></td><td></td><td></td><td></td><td>@#106;</td><td></td></tr><tr><td>11</td><td></td><td>013</td><td></td><td>(vertical tab)</td><td></td><td></td><td></td><td>&#<b>4</b>3;</td><td></td><td></td><td></td><td></td><td>a#75;</td><td></td><td></td><td></td><td></td><td>a#107;</td><td></td></tr><tr><td>12</td><td></td><td>014</td><td></td><td>(NP form feed, new page)</td><td></td><td></td><td></td><td>,</td><td></td><td></td><td></td><td></td><td>a#76;</td><td></td><td></td><td></td><td></td><td>l</td><td></td></tr><tr><td>13</td><td></td><td>015</td><td></td><td>(carriage return)</td><td></td><td></td><td></td><td>&#<b>4</b>5;</td><td></td><td></td><td></td><td></td><td>a#77;</td><td></td><td></td><td></td><td></td><td>m</td><td></td></tr><tr><td>14</td><td></td><td>016</td><td></td><td>(shift out)</td><td></td><td></td><td></td><td>&#<b>4</b>6;</td><td></td><td></td><td></td><td></td><td>a#78;</td><td></td><td></td><td></td><td></td><td>n</td><td></td></tr><tr><td>15</td><td></td><td>017</td><td></td><td>(shift in)</td><td></td><td></td><td></td><td>6#47;</td><td>-</td><td></td><td></td><td></td><td>6#79;</td><td></td><td></td><td></td><td></td><td>o</td><td></td></tr><tr><td></td><td></td><td></td><td>DLE</td><td>(data link escape)</td><td></td><td></td><td></td><td>a#48;</td><td></td><td></td><td></td><td></td><td>6#8O;</td><td></td><td></td><td></td><td></td><td>p</td><td></td></tr><tr><td></td><td></td><td></td><td></td><td>(device control 1)</td><td></td><td></td><td></td><td>a#49;</td><td></td><td></td><td></td><td></td><td>4#81;</td><td>_</td><td></td><td></td><td></td><td>q</td><td></td></tr><tr><td></td><td></td><td></td><td></td><td>(device control 2)</td><td></td><td></td><td></td><td>2</td><td></td><td></td><td></td><td></td><td>R</td><td></td><td></td><td></td><td></td><td>r</td><td></td></tr><tr><td></td><td></td><td></td><td></td><td>(device control 3)</td><td></td><td></td><td></td><td>3</td><td></td><td></td><td></td><td></td><td>S</td><td></td><td></td><td>-</td><td></td><td>s</td><td></td></tr><tr><td></td><td></td><td></td><td></td><td>(device control 4)</td><td></td><td></td><td></td><td>4</td><td></td><td></td><td></td><td></td><td>T</td><td></td><td></td><td></td><td></td><td>t</td><td></td></tr><tr><td></td><td></td><td></td><td></td><td>(negative acknowledge)</td><td></td><td></td><td></td><td><b>6#53</b>;</td><td></td><td></td><td></td><td></td><td>6#85;</td><td></td><td></td><td></td><td></td><td>u</td><td></td></tr><tr><td></td><td></td><td></td><td></td><td>(synchronous idle)</td><td></td><td></td><td></td><td>a#54;</td><td></td><td></td><td></td><td></td><td>V</td><td></td><td></td><td></td><td></td><td>v</td><td></td></tr><tr><td></td><td></td><td></td><td></td><td>(end of trans. block)</td><td></td><td></td><td></td><td>a#55;</td><td></td><td></td><td></td><td></td><td>a#87;</td><td></td><td></td><td></td><td></td><td>a#119;</td><td></td></tr><tr><td></td><td></td><td></td><td></td><td>(cancel)</td><td></td><td></td><td></td><td>a#56;</td><td></td><td></td><td></td><td></td><td>4#88;</td><td></td><td></td><td></td><td></td><td>x</td><td></td></tr><tr><td></td><td></td><td>031</td><td></td><td>(end of medium)</td><td></td><td></td><td></td><td><u>4</u>#57;</td><td></td><td></td><td></td><td></td><td>Y</td><td></td><td></td><td></td><td></td><td>y</td><td></td></tr><tr><td></td><td></td><td>032</td><td></td><td>(substitute)</td><td></td><td></td><td></td><td><u>4#58;</u></td><td></td><td>90</td><td></td><td></td><td>Z</td><td>Z</td><td></td><td></td><td></td><td>z</td><td></td></tr><tr><td></td><td></td><td></td><td>ESC</td><td>(escape)</td><td></td><td></td><td></td><td>6#59;</td><td></td><td>91</td><td></td><td></td><td>6#91;</td><td>[</td><td></td><td></td><td></td><td>6#123;</td><td></td></tr><tr><td></td><td></td><td>034</td><td></td><td>(file separator)</td><td></td><td></td><td></td><td><b>%#60;</b></td><td></td><td></td><td></td><td></td><td>6#92;</td><td></td><td></td><td></td><td></td><td>a#124;</td><td></td></tr><tr><td></td><td></td><td>035</td><td></td><td>(group separator)</td><td></td><td></td><td></td><td>a#61;</td><td></td><td></td><td></td><td></td><td>6#93;</td><td>_</td><td></td><td></td><td></td><td>a#125;</td><td></td></tr><tr><td></td><td></td><td>036</td><td></td><td>(record separator)</td><td></td><td></td><td></td><td>></td><td></td><td></td><td></td><td></td><td>a#94;</td><td>^</td><td></td><td></td><td></td><td>~</td><td></td></tr><tr><td>31</td><td>1F</td><td>037</td><td>US</td><td>(unit separator)</td><td>63</td><td>ЗF</td><td>077</td><td>?</td><td>2</td><td>95</td><td>5F</td><td>137</td><td>_</td><td>_</td><td>127</td><td>7<b>F</b></td><td>177</td><td></td><td>DEL</td></tr></tbody></table> |       |     |    |     |       |     |     |    |     |         |           |

Source: www.LookupTables.com

## **Read Quality Control**

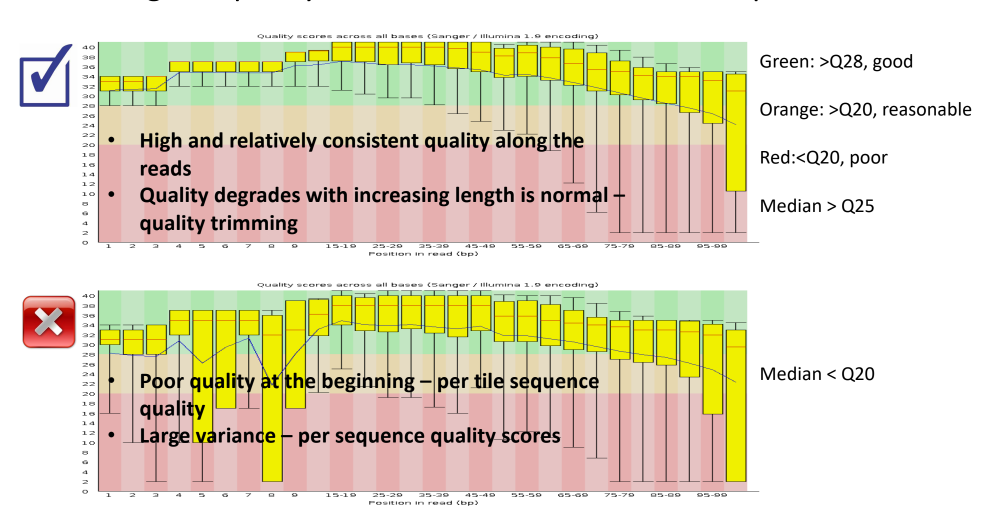
- Library construction could introduce bias
  - Fragmentation, ligation, amplification
  - GC bias
  - Over-amplification
  - Contamination

- Sequencing errors
  - Chemical, optical, computational

| Platform           | Primary<br>error | Error rate (%)    |
|--------------------|------------------|-------------------|
| Illumina           | Substitution     | 0.1               |
| PacBio             | Indel            | 12 (consensus: 1) |
| Oxford<br>Nanopore | Indel            | 3 - 20            |

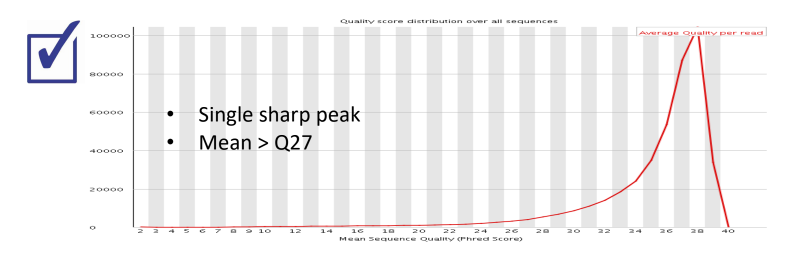
#### Per base sequence quality - FastQC

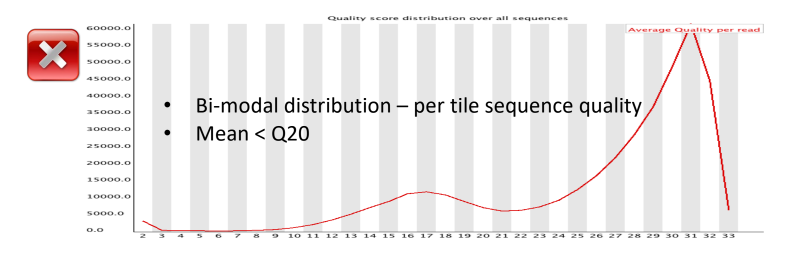
Range of quality values across all bases at each position



#### Per sequence quality scores - FastQC

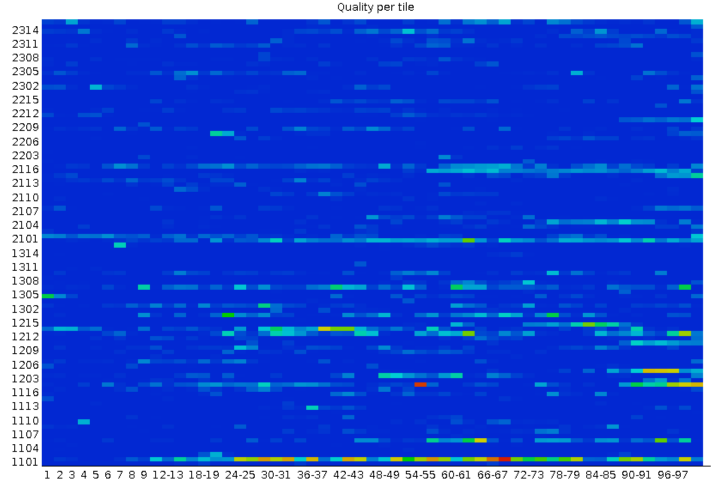
Subset of sequences with universally low quality values





### Per tile sequence quality - FastQC

 Quality scores from each tile across all bases - loss in quality associated with only one part of the flowcell



Deviation from average quality

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Cold colors: ≥ average

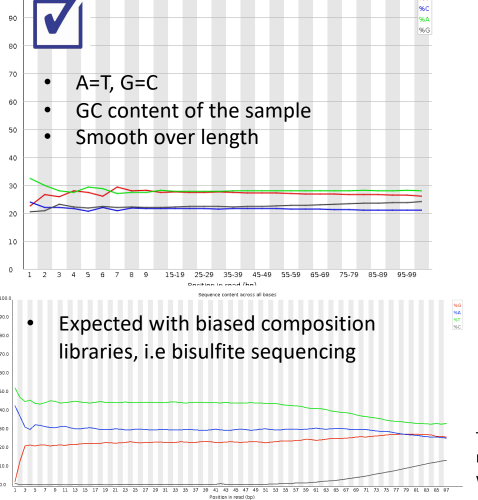
Hotter color: worse quality

Good: universal blue

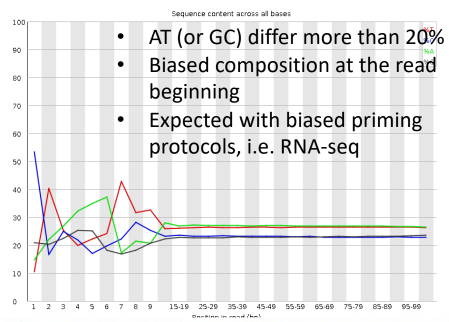
Failure: < average - 5

#### Per base sequence content - FastQC

• The portion of A, T, G, and C at each position



Sequence content across all bases



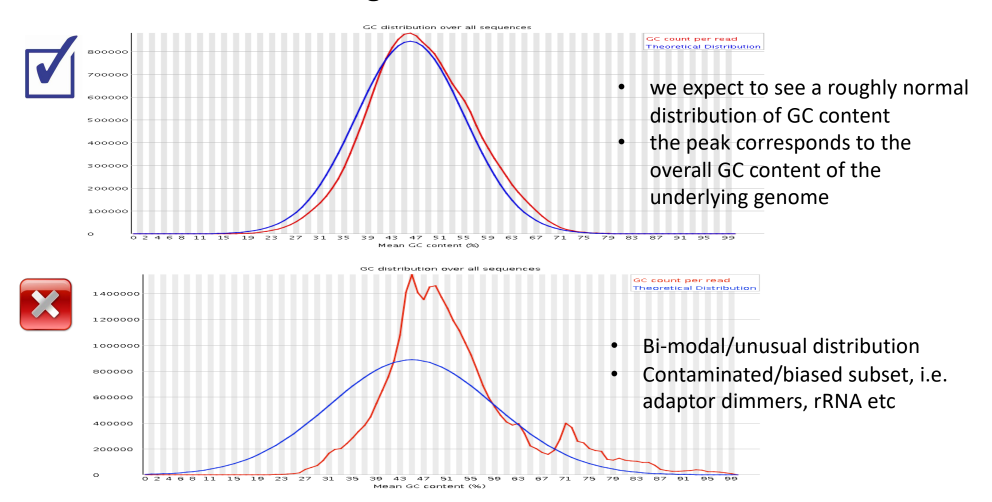
#### Biases in Illumina transcriptome sequencing caused by random hexamer priming

Kasper D. Hansen¹,\*, Steven E. Brenner² and Sandrine Dudoit¹,3

Treatment of DNA with bisulfite converts cytosine to uracil, but leaves methylated cytosine unaffected. Therefore, DNA that has been treated with bisulfite retains only methylated cytosines.

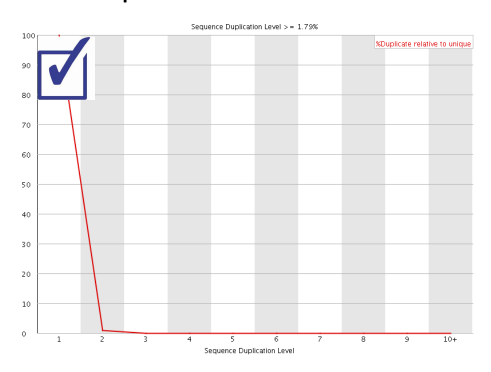
#### Per sequence GC content - FastQC

Distribution of average GC in all reads

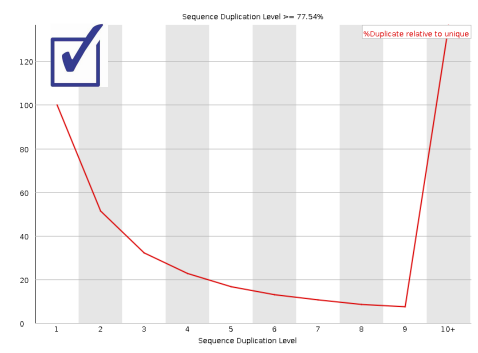


#### Sequence duplication - FastQC

 Relative number of sequences with different degrees of duplication



Essentially no duplication



#### High duplication levels:

- DNA-seq: PCR over amplification, too little input material
- Normal in RNA-seq: high expression

#### Overrepresented sequences - FastQC

- Sequences make up >0.1 % of the total
- Compare those with a contamination database for finding contamination (i.e. adaptor dimmers)

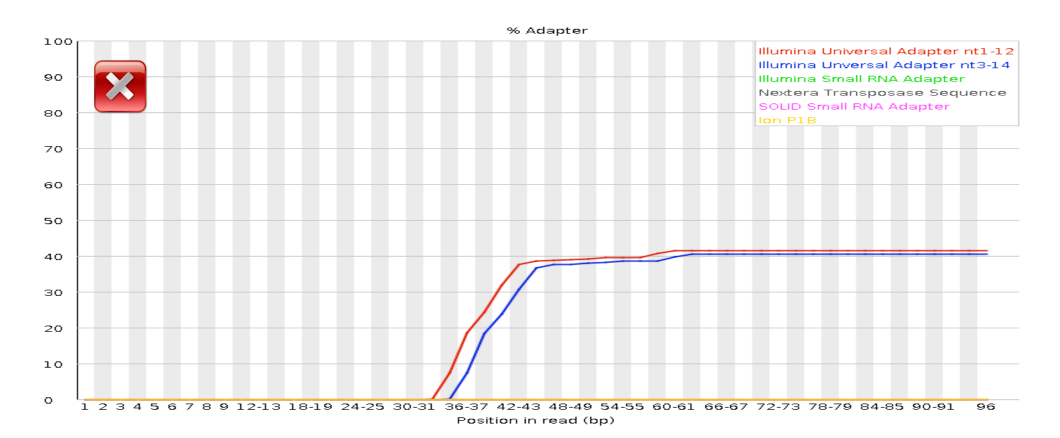
# Overrepresented sequences

| Sequence   | Count | Percentage          | Possible Source                          |
|--|-------|---------------------|--|
| GGAAGAGCACACGTCTGAACTCCAGTCACCAGATCATCTCGTATGCCGTC | 75874 | 1.5613887498682963  | TruSeq Adapter, Index 7 (100% over 50bp) |
| GGAAGAGCACACGTCTGAACTCCAGTCACCGATGTATCTCGTATGCCGTC | 7636  | 0.15713900010536297 | TruSeq Adapter, Index 2 (100% over 50bp) |
| GGAAGAGCACACGTCTGAACTCCAGTCACACAGTGATCTCGTATGCCGTC | 7539  | 0.1551428656095248  | TruSeq Adapter, Index 5 (100% over 50bp) |
| GGAAGAGCACACGTCTGAACTCCAGTCACGCCAATATCTCGTATGCCGTC | 5117  | 0.10530123933199874 | TruSeq Adapter, Index 6 (100% over 50bp) |

- Can be normal and biologically meaningful
  - highly expressed transcripts
  - high copy number repeats
  - Less diverse library (amplicons)

#### **Adapter Content - FastQC**



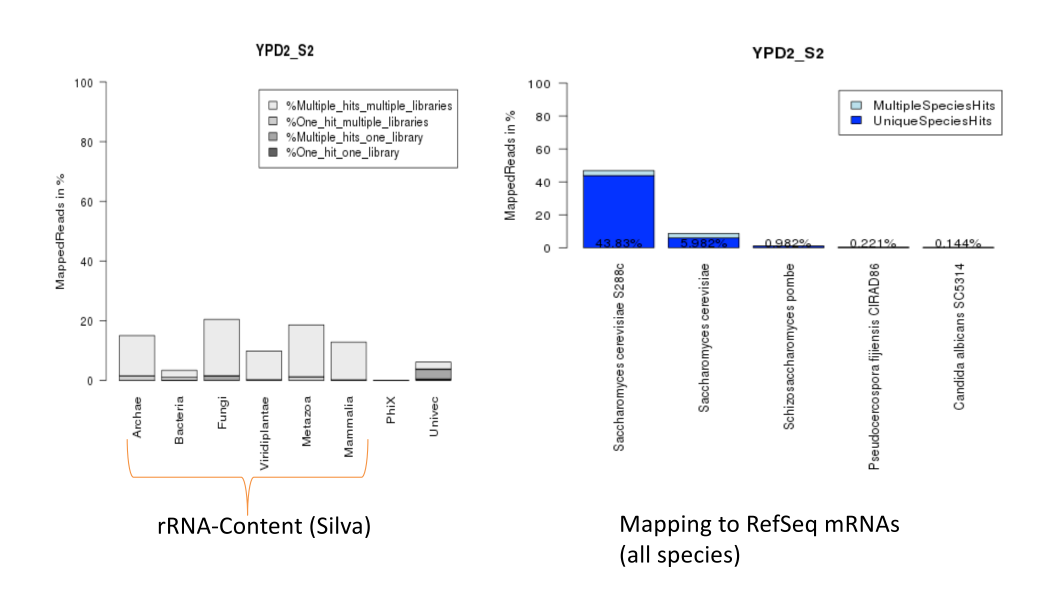


# Millions of reads with base resolution

@HWI-ST1034:40:C08PJACXX:2:1101:20681:1994 1:N:0:ATCACG  ${\tt CTCGNAGACTGGCAACTTGTTCTGGTTTACTGCACCTTCTTTTAAAGGCAGAAAGGCTTTTTGATAAAGAAGTTGTGAAAAGGCTACATGAGCTGCTTTTA$ @HWI-ST1034:40:CO8PJACXX:2:1101:1907:2005 1:N:0:ATCACG @HWI-ST1034:40:CO8PJACXX:2:1101:2155:2031 1:N:0:ATCACG  ${\tt CAATCAATTAACAATATTAGTTAGATAAGCACTTCCTTAACCACCCTCTCAAAGTTGGCAAATGAAGAACCCCCTTTCTCAATAGCTTTAACCGCGCTCTC$ @HWI-ST1034:40:CO8PJACXX:2:1101:2220:2057 1:N:0:ATCACG @HWI-ST1034:40:CO8PJACXX:2:1101:2460:2116 1:N:0:ATCACG @HWI-ST1034:40:CO8PJACXX:2:1101:2463:2168 1:N:O:ATCACG  ${\tt CGTTCATATGCAAAAGAAGCTTCTCAGTCTGCTTTACCACCTCTTAAAGGGGATCAAATGTTGAAGAACATCTTTTTTGAGGTAAAGAACAAATTTGATAT$ @HWI-ST1034:40:CO8PJACXX:2:1101:2378:2207 1:N:0:ATCACG  ${\tt CACGCGGTGTGGAAAACCCCTTCACATCCATCAATGGCGGCTCGGAGCGATTCAAAATCAAGCATATCCGCTTTGTACAGCACAAGACGATCCGATGCTCC$ 

- How accurate was the sequencing → Fastqc
- Are these reads the intented ones → FastqScreen

#### **Contamination Check - FastqScreen**



## Data preprocessing common tasks

- 1. Trimming: remove bad bases from (end(s) of) reads
  - Adapter sequence
  - Low quality bases
- 2. Filtering: remove bad reads
  - Low quality reads
  - Contaminating sequences
  - Low complexity reads (repeats)
  - Short (<20bp) reads they slow down mapping software</li>

## Data preprocessing software

- fastp
  - https://github.com/OpenG ene/fastp
  - Adapter trimming, quality trimming &filtering, ...
- Trimmomatic
  - https://github.com/usadellab/T rimmomatic
  - Adapter trimming, quality trimming &filtering, ...
- FlexBar (FAR)
  - https://github.com/seqan/flexb ar
  - Flexible barcode detection and adapter removal

#### FASTX

- <a href="http://hannonlab.cshl.edu/fast">http://hannonlab.cshl.edu/fast</a>
   x toolkit/
- Reformat, stats, collapse duplicated reads, trim, filter, reverse compliment
- TagCleaner
  - http://tagcleaner.sourceforge.n et
  - Trim MIDs or adaptors, demultiplexing
- DeconSeq
  - <a href="http://deconseq.sourceforge.">http://deconseq.sourceforge.</a>
    net
  - Remove potential contaminants

#### Recommendations

- Always generate quality control plots visualizing key characteristics for all libraries
- Trim and/or filter data if needed
- Applications where erroneous reads are of concern:
  - de novo assembly
  - low coverage variant calling
- Applications that are more tolerant to low quality bases
  - RNA-seq