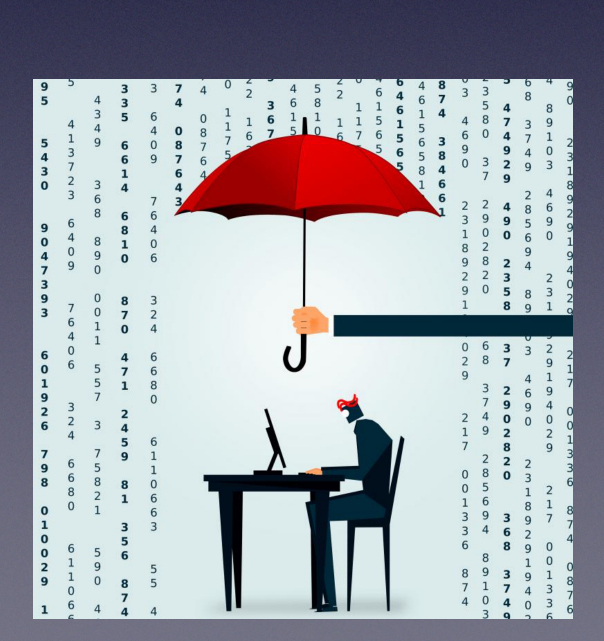
Sequence format and preprocessing

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Illumina short reads

- We receive files in fastq format from the sequencing center
- Two files per sample, 1 forward (R1) and 1 reverse (R2)
- Normally, reads overlap
- Depending on the library preparation, all reads are in the same direction (5'-3') or their directions are mixed in both R1 and R2
- We normally work with gzipped fastq files to save space

How files look like when they are received

18S_BL060704_30-MSTAReuk_R1.fastq.gz 18S_BL060704_30-MSTAReuk_R2.fastq.gz 18S_BL060801_022-MSTAReuk_R1.fastq.qz 18S_BL060801_022-MSTAReuk_R2.fastq.gz

18S_BL091222_30-MSTAReuk_R1.fastq.gz 18S_BL091222_30-MSTAReuk_R2.fastq.gz 18S_BL100120_022-MSTAReuk_R1.fastq.qz 18S_BL100120_022-MSTAReuk_R2.fastq.gz

18S_BL130417_30-MSTAReuk_R2.fastq.qz 18S_BL060613_30r3_MSTAReuk_R1.fastq.gz 18S_BL091105_30r3_MSTAReuk_R1.fastq.gz 18S_BL130417_30r3_MSTAReuk_R1.fastq.gz 18S_BL060613_30r3_MSTAReuk_R2.fastq.gz 18S_BL091105_30r3_MSTAReuk_R2.fastq.gz 18S_BL130417_30r3_MSTAReuk_R2.fastq.gz 18S_BL130507_022-MSTAReuk_R1.fastq.qz 18S_BL130507_022-MSTAReuk_R2.fastq.gz 18S_BL130507_30-MSTAReuk_R1.fastq.gz 18S_BL130507_30-MSTAReuk_R2.fastq.gz 18S_BL130507_30r3_MSTAReuk_R1.fastq.gz 18S_BL130507_30r3_MSTAReuk_R2.fastq.gz 18S_BL130604_022-MSTAReuk_R1.fastq.qz 18S_BL130604_022-MSTAReuk_R2.fastq.gz 18S_BL130604_30-MSTAReuk_R1.fastq.gz 18S_BL130604_30-MSTAReuk_R2.fastq.gz 18S_BL130604_30r3_MSTAReuk_R1.fastq.qz 18S_BL130604_30r3_MSTAReuk_R2.fastq.gz 18S_BL130709_022-MSTAReuk_R1.fastq.gz 18S_BL130709_022-MSTAReuk_R2.fastq.qz

fastq format

- Four sequences per line
 - 1. @sequence.ID
 - 2. ACTGACTGACTG # nucleotide sequence
 - 3. + (separator)
 - 4. Quality scores (Phred +33: normally 0-41)



Quite a bit of information here

@M02696:67:000000000-B44VG:1:1101:11781:1257 1:N:0:57

+

@M02696:67:000000000-B44VG:1:1101:8695:1347 1:N:0:57

+

@M02696:67:000000000-B44VG:1:1101:22691:1423 1:N:0:57

+

+

@M02696:67:000000000-B44VG:1:1101:17122:1672 1:N:0:57

+

@M02696:67:000000000-B44VG:1:1101:21438:1779 1:N:0:57

+

@M02696:67:000000000-B44VG:1:1101:11781:1257 1:N:0:57

The first line, identifying the sequence, contains the following elements.

@<instrument>:<run number>:<flowcell ID>:<lane>:<tile>:<x-pos>:<UMI> <read>:<is filtered>:<control number>:<index>

Table 1 FASTQ File Elements

Element	Requirements	Description	
@	@	Each sequence identifier line starts with @.	
<instrument></instrument>	Characters allowed:	Instrument ID.	
	a-z, A-Z, 0-9 and underscore		
<run number=""></run>	Numerical	Run number on instrument.	
<flowcell id=""></flowcell>	Characters allowed:		
	a-z, A-Z, 0-9		
<lane></lane>	Numerical	Lane number.	
<tile></tile>	Numerical	Tile number.	
<x_pos></x_pos>	Numerical	X coordinate of cluster.	
<y_pos></y_pos>	Numerical	Y coordinate of cluster.	
<umi></umi>	Restricted characters: A/T/G/C/N	Optional, appears when UMI is specified in sample sheet. UMI sequences for Read 1 and Read 2, seperated by a plus [+].	
<read></read>	Numerical	Read number. 1 can be single read or Read 2 of paired-end.	
<is filtered=""></is>	Y or N	Y if the read is filtered (did not pass), N otherwise.	
<control< td=""><td>Numerical</td><td colspan="2">0 when none of the control bits are on, otherwise it is an even number.</td></control<>	Numerical	0 when none of the control bits are on, otherwise it is an even number.	
number>		On HiSeq X and NextSeq systems, control specification is not performed and this number is always 0.	
<index></index>	Restricted characters: A/T/G/C/N	Index of the read.	

Sanger Phred quality scores

Phred Quality Score	Probability of incorrect base call	Base call accuracy
10	1 in 10	90%
20	1 in 100	99%
30	1 in 1000	99.9%
40	1 in 10000	99.99%
50	1 in 100000	99.999%

 $Q = -10 \log_{10} P$

Q = Phred quality scores

P = base calling error probability

Calculating Phred scores

- To determine quality scores, Phred first calculates several parameters related to peak shape and peak resolution at each base.
- Phred then uses these parameters to look up a corresponding quality score in huge lookup tables.
- These lookup tables were generated from sequence traces where the correct sequence was known, and are hard coded in Phred; different lookup tables are used for different sequencing chemistries and machines.

- Quality scores are encoded in ASCII
 (American Standard Code for Information Interchange)
- The start in character 33 (Phred+33)

CCCCCGGGGFG@CGG; FDEFFGEFGGGG9E@CFGCGGGEFG<EFGFEFGGGGFGGGEG<FC@@@6@F8@FCGAFFFF, 6C6EC@FCFGGGGGGGGGGGGGGFGGGFF; CFFFAFF, BCE<CFFEFF7F8?, CF<EBCF, AFDGFAFF<
9@BEFEG?FC9, CE<FD?A7CGEG:FDFG, 3A;, CDFGGGFF, =CF, 6, 6BFGF, 6+4@EEGGGG7>EC?
FGGF@FCGED8CFFGG79D9CCF<?C4713?FFFCDE

convert ascii33 to error probability $Q_{\text{PHRED}} = -10 \times \log_{10}(P_e)$ 1-error in # # errors in %correct probability 2.85Gb bases 2,858,034,764 33 1.00E+00 0.000% 20.567% 2,270,217,709 7.94E-01 1,803,298,025 6.31E-01 36.904% 1,432,410,537 36 5.01E-01 49.881% 37 3.98E-01 60.189% 3 1,137,804,133 38 3.16E-01 68.377% 903,789,949 717,905,874 39 2.51E-01 74.881% 40 2.00E-01 5 570,252,906 80.047% 452,967,984 41 1.58E-01 84.151% 42 359,805,259 1.26E-01 87.411% 285,803,476 1.00E-01 90.000% 7.94E-02 92.057% 13 227,021,771 6.31E-02 16 180, 329, 803 93.690% 5.01E-02 94.988% 20 143,241,054 113,780,413 3.98E-02 25 96.019% 90,378,995 32 3.16E-02 96.838% 2.51E-02 71,790,587 97.488% 50 2.00E-02 98.005% 57,025,291 45,296,798 1.58E-02 98.415% 35,980,526 79 1.26E-02 98.741% 53 1.00E-02 99.000% 100 28,580,348 7.94E-03 126 22,702,177 99.206% 158 18,032,980 55 22 6.31E-03 99.369% 14,324,105 23 5.01E-03 200 99.499% 251 3.98E-03 99.602% 11,378,041 24 316 58 3.16E-03 25 99.684% 9,037,899 398 59 2.51E-03 99.749% 7,179,059 26 501 2.00E-03 99.800% 5,702,529 631 4,529,680 1.58E-03 99.842% 61 1.26E-03 99.874% 794 3,598,053 29 1,000 2,858,035 1.00E-03 99.900% 7.94E-04 99.921% 1,259 2,270,218 1,585 32 6.31E-04 99.937% 1,803,298 1,995 5.01E-04 99.950% 1,432,411 2,512 67 34 3.98E-04 99.960% 1,137,804 99.968% 3,162 903,790 3.16E-04 2.51E-04 99.975% 3,981 717,906 2.00E-04 99.980% 5,012 570,253 71 | 38 1.58E-04 99.984% 6,310 452,968 H 72 39 1.26E-04 99.987% 359,805 I 73 40 1.00E-04 99.990% 10,000 285,803 J 74 41 7.94E-05 227,022 99.992% 12,589 K 75 42 6.31E-05 15,849 180,330 99.994% L 76 43 5.01E-05 99.995% 19,953 143,241 M 77 44 3.98E-05 99.996% 25,119 113,780 N 78 45 3.16E-05 99.997% 31,623 90,379 O 79 46 2.51E-05 99.997% 39,811 71,791

Phred encoding in different sequencers

```
!"#$%&'()*+,-./0123456789:;<=>?@ABCDEFGHIJKLMNOPQRSTUVWXYZ[\]^_`abcdefghijklmnopqrstuvwxyz{|}~
33
                            104
                                     126
Phred+33, raw reads typically (0, 40)
       Solexa+64, raw reads typically (-5, 40)
I - Illumina 1.3+ Phred+64, raw reads typically (0, 40)
J - Illumina 1.5+ Phred+64, raw reads typically (3, 40)
 with 0=unused, 1=unused, 2=Read Segment Quality Control Indicator (bold)
 (Note: See discussion above).
L - Illumina 1.8+ Phred+33, raw reads typically (0, 41)
```

Removing primers

- The sequences received from the sequencing center may contain primers used to amplify them
- Primers need to be removed as they normally contain ambiguous positions that can interfere with DADA2
- DADA2 assumes primers have been removed

Checking if sequences have primers in unix

- We know the primer sequence expected in the reads
- Fw:CCAGCA[ACGT]C[ACGT]GCGGTAATTCC
- Rv: ACTTCGTTCTTGAT[AGCT][AGCT]
 - Ambiguities are included to match sequences
- We use zgrep to match the primer against the gzipped sequences

```
[rlogares@marbits raw]$ ls
BL100525E-MSTAReuk_R1.fastq.gz BL100706E-MSTAReuk_R1.fastq.gz BL100914E-MSTAReuk_R1.fastq.gz primers_R1_in_reads
BL100525E-MSTAReuk_R2.fastq.gz BL100706E-MSTAReuk_R2.fastq.gz BL100914E-MSTAReuk_R2.fastq.gz primers_R2_in_reads
BL100622E-MSTAReuk_R1.fastq.gz BL100803E-MSTAReuk_R1.fastq.gz clipping_primers.sh
BL100622E-MSTAReuk_R2.fastq.gz BL100803E-MSTAReuk_R2.fastq.gz cutadapt.o40252
[rlogares@marbits raw]$ zgrep -c --color CCAGCA[ACGT]C[ACGT]GCGGTAATTCC BL100525E-MSTAReuk_R1.fastq.gz
23843
[rlogares@marbits raw]$
[rlogares@marbits raw]$ zgrep -c --color ACTTTCGTTCTTGAT[AGCT][AGCT][AGCT] BL100525E-MSTAReuk_R2.fastq.gz
23856
[rlogares@marbits raw]$
```

Forward primer

Reverse primer

- As several counts are given, we inspect the sequences visually

[rlogares@marbits raw]\$ zgrep --color CCAGCA[ACGT]C[ACGT]GCGGTAATTCC BL100525E-MSTAReuk_R1.fastq.gz

We use cutadapt to remove primers

- Program: https://cutadapt.readthedocs.io/en/stable/
- Runs in unix
- Cutadapt will search for primers in R1 and R2 sequences and remove them
- It can also remove all sequences where primers have not been found

```
# Running cutadapt in a loop (NB: use arrays if you have a cluster)
for i in $(ls *fastq.gz | cut -f 1 -d - | uniq); \
   do cutadapt -g CCAGCASCYGCGGTAATTCC -G ACTTTCGTTCTTGATYRR \
   -m 100 -M 350 --match-read-wildcards --pair-filter=both -q 10 \
   -o $i-MSTAReuk_R1.clipped.fastq.gz -p $i-MSTAReuk_R2.clipped.fastq.gz \
   $i-MSTAReuk R1.fastq.gz $i-MSTAReuk R2.fastq.gz; done
      -g ADAPTER, --front=ADAPTER
                        Sequence of an adapter ligated to the 5' end (paired
                        data: of the first read). The adapter and any
                        preceding bases are trimmed. Partial matches at the 5'
                        end are allowed. If a '^' character is prepended
                         ('anchoring'), the adapter is only found if it is a
                        prefix of the read.
       Paired-end options:
       The -A/-G/-B/-U options work like their -a/-b/-g/-u counterparts, but
       are applied to the second read in each pair.
                            5' adapter to be removed from second read in a pair.
         -G ADAPTER
         -m LENGTH, --minimum-length=LENGTH
                        Discard reads shorter than LENGTH. Default: 0
         -M LENGTH, --maximum-length=LENGTH
                        Discard reads longer than LENGTH. Default: no limit
         --match-read-wildcards
                        Interpret IUPAC wildcards in reads. Default: False
         --pair-filter=(any|both)
                        Which of the reads in a paired-end read have to match
                         the filtering criterion in order for the pair to be
                         filtered. Default: any
         -q [5'CUTOFF,]3'CUTOFF, --quality-cutoff=[5'CUTOFF,]3'CUTOFF
                        Trim low-quality bases from 5' and/or 3' ends of each
                         read before adapter removal. Applied to both reads if
                         data is paired. If one value is given, only the 3' end
                        is trimmed. If two comma-separated cutoffs are given,
                         the 5' end is trimmed with the first cutoff, the 3'
                        end with the second.
         -o output file R1
         -p FILE, --paired-output=FILE
                        Write second read in a pair to FILE.
```

ambiguities are interpreted

There are several additional options

- After cutadap, sequences are ready to be used in dada2
- It is good to double check primers are gone using the same zgrep command used before
- We don't analyse the overall quality of the sequences, as this will be done later with dada2
- We only remove entire sequences that look very wrong with cutadapt
- It is important to consider whether sequences come from sequencers with 4- or 2- color chemistries, as this will change cutadapt parameters
- There are alternative tools, such as Trimmomatic

THEEND