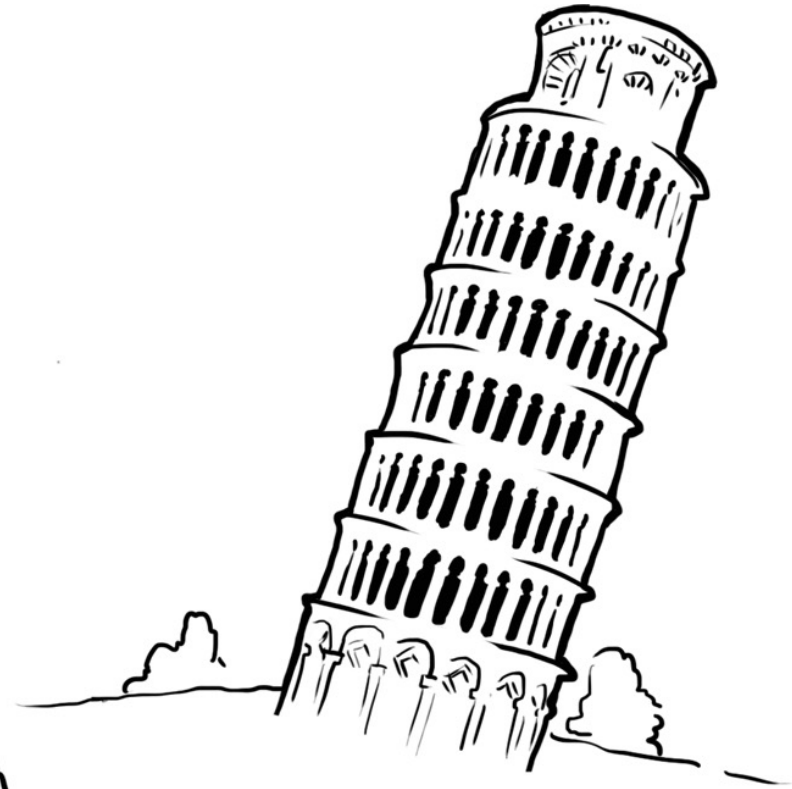


# Quality control

THAT'S GOOD ENOUGH.



QUALITY CONTROL

# Overview of this talk

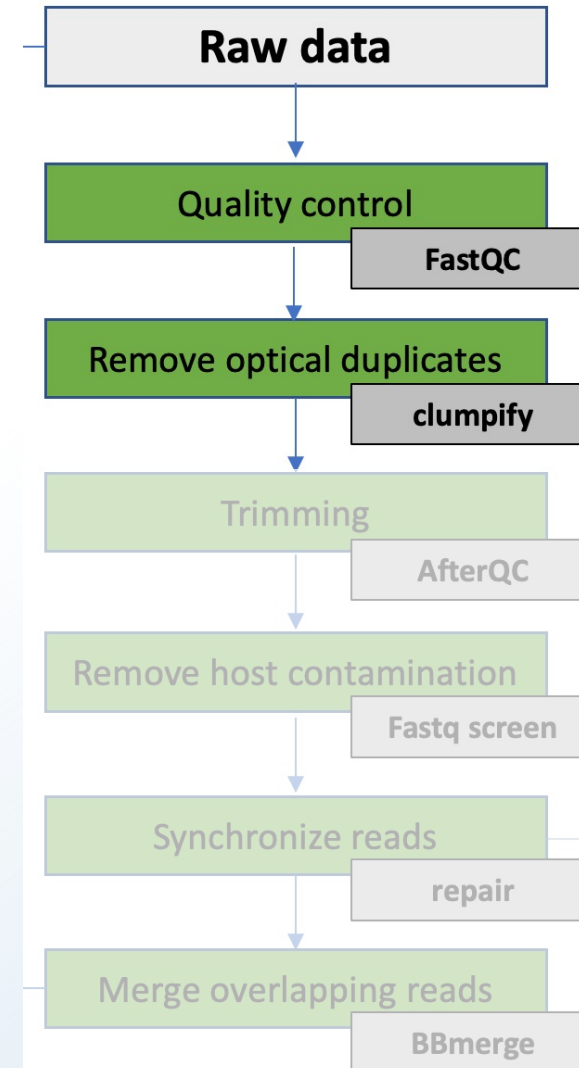
---

Sequencing recap

Data integrity

Quality control of sequence data

Optical duplicates in data



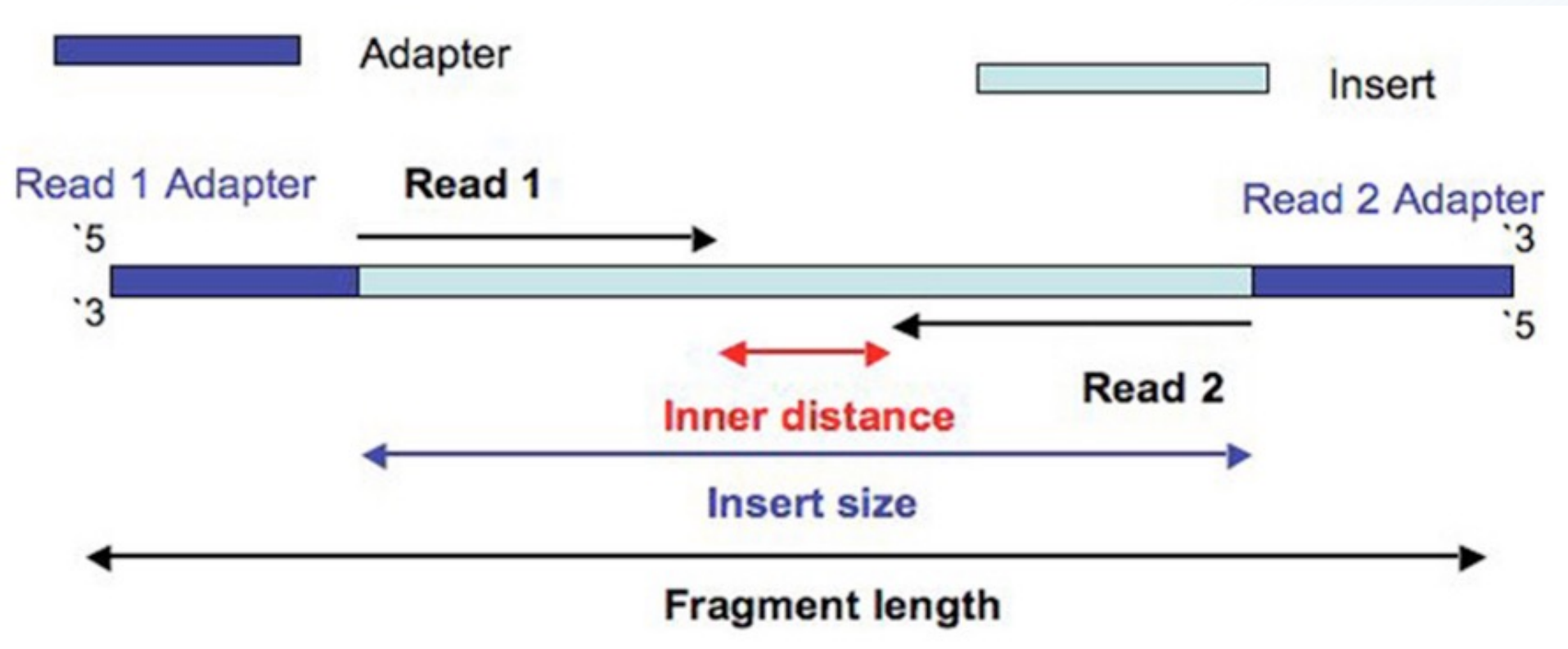
# Wet lab QC - DNA quality is always checked prior to library preparation and sequencing

Methods for quantification and evaluation of NGS libraries:

	Agarose Gel Electrophoresis	Spectrophotometry	qPCR	Digital PCR (dPCR)
Quantification	Relative	Absolute (*)	Relative	Absolute (**)
Sensitivity	+	++	++++	+++++
Quality Assessment	+	+	+++	+++++
Cost	+	++	+++	+++

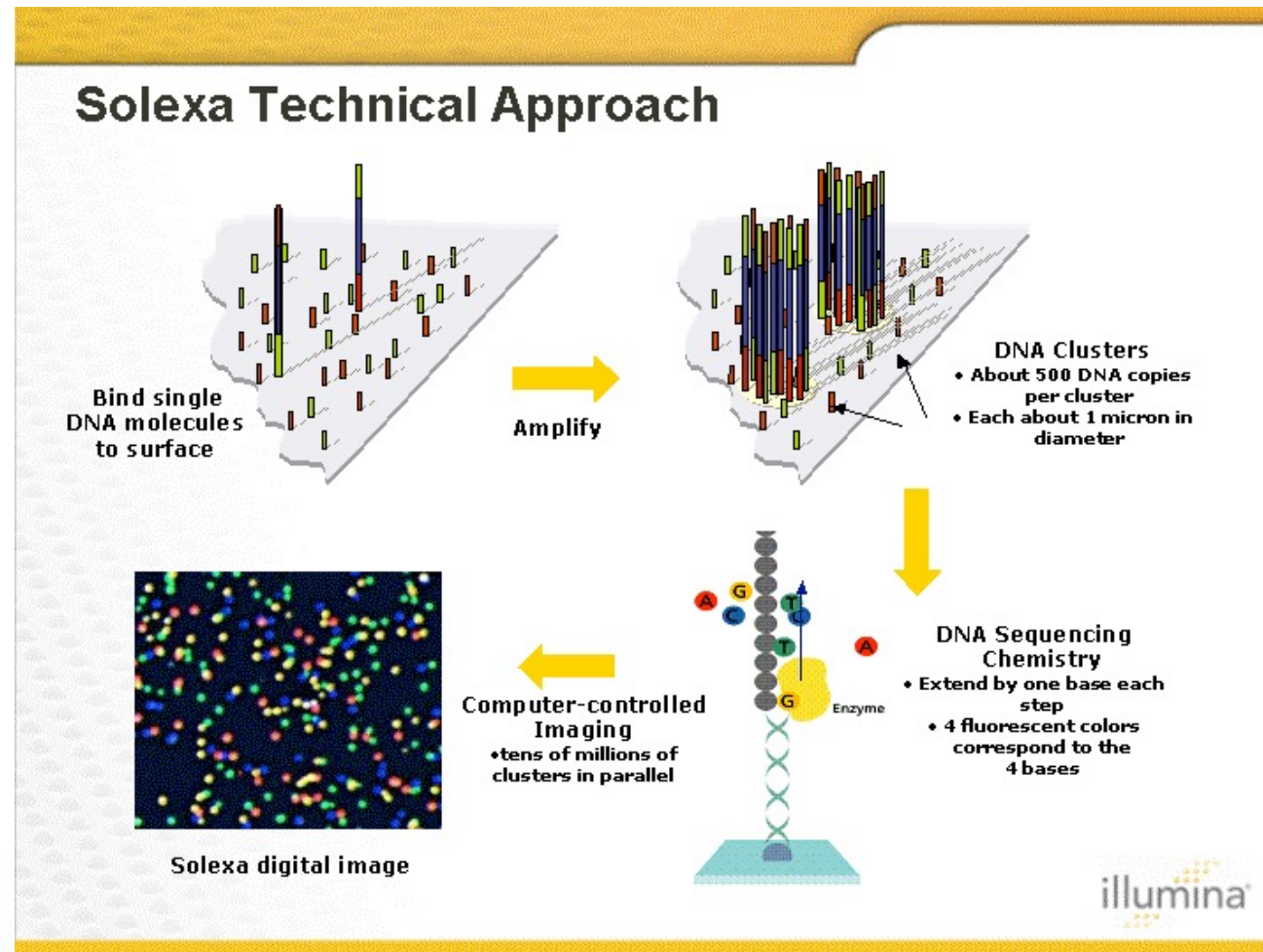
# DNA sequencing recap – Illumina technology

Paired-end (PE) Illumina data



# Sequencing quality scores measure the probability that a base is called incorrectly

Base calls are made directly from signal intensity measurements



## Sequencing technologies are not perfect and do produce errors

---

QC ensures that the data used for downstream analysis does not contain (too many) errors and poor quality sequences

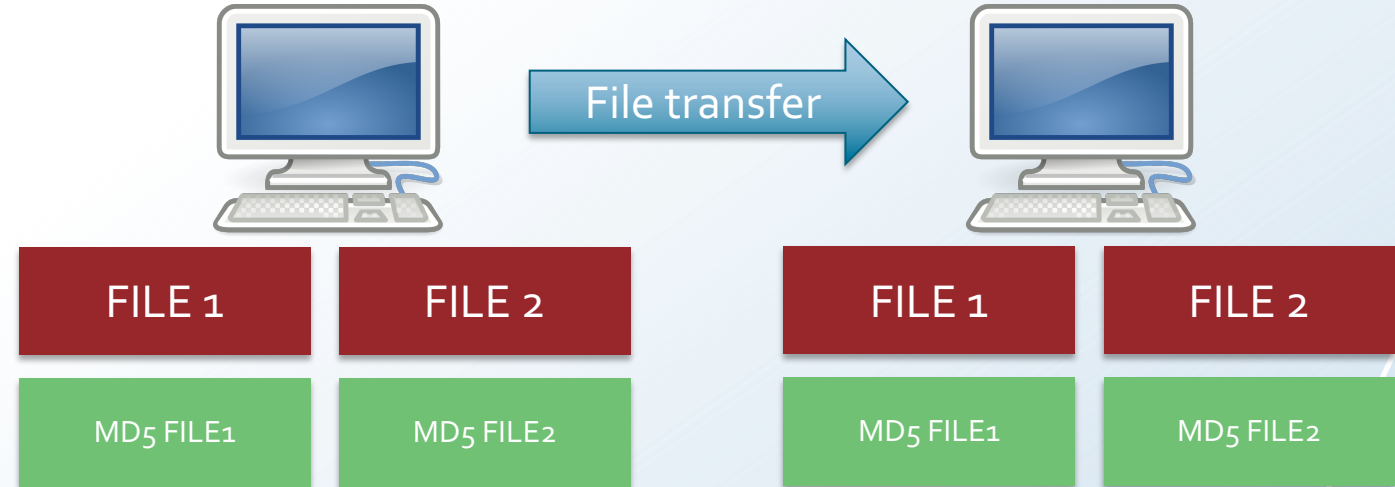
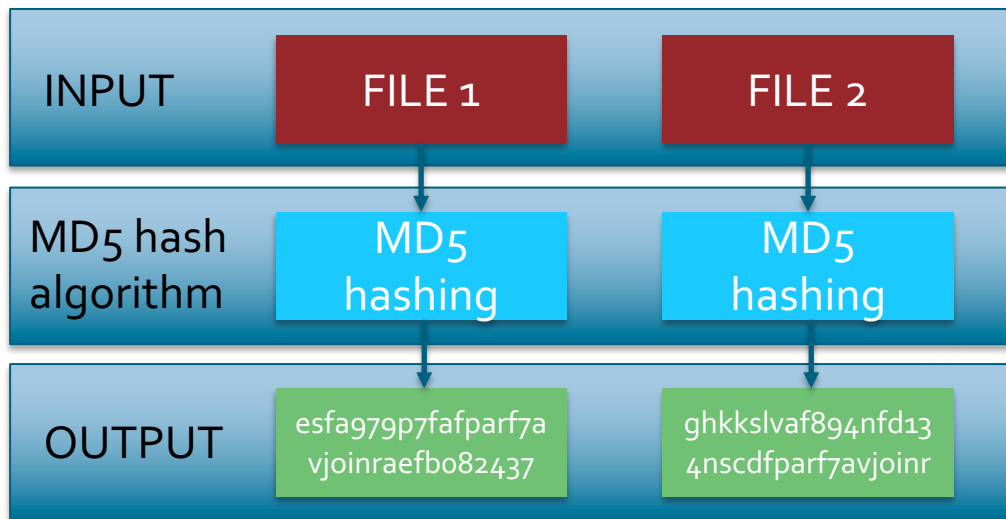


# Check that the data is complete

File checksums ensure data integrity, eg. MD5

Calculate file checksums before transfer

After the transfer, checksum the transferred file



## Check the data quantity

---

Generate simple statistics of FASTA/Q files using eg. SeqKit

```
$ seqkit stat *.fq
```

file	format	type	num_seqs	sum_len	min_len	avg_len	max_len
dataset_A.fq	FASTQ	DNA	2,675,748	244,807,643	50	99	100
dataset_B.fq	FASTQ	DNA	3,789,194	388,099,750	100	100	100
dataset_C.fq	FASTQ	DNA	9,186,045	918,604,500	100	100	100





# Check the format - The Illumina output are FASTQ files

---

## The FASTQ header

```
@EAS139:136:FC706VJ:2:2104:15343:197393 1:Y:18:ATCACG
```

<b>EAS139</b>	the unique instrument name
<b>136</b>	the run id
<b>FC706VJ</b>	the flowcell id
<b>2</b>	flowcell lane
<b>2104</b>	tile number within the flowcell lane
<b>15343</b>	'x'-coordinate of the cluster within the tile
<b>197393</b>	'y'-coordinate of the cluster within the tile
<b>1</b>	the member of a pair, 1 or 2 ( <i>paired-end or mate-pair reads only</i> )
<b>Y</b>	Y if the read is filtered, N otherwise
<b>18</b>	0 when none of the control bits are on, otherwise it is an even number
<b>ATCACG</b>	index sequence

# A range of sequence quality scores depending on technology and the base caller

Most modern sequencing machine, such as MiSeq use Illumina 1.8+



# The FASTQ quality scores derives from using ASCII encoding

ASCII codes represent text in computers, originally based on the English alphabet

Dec	Hex	Name	Char	Ctrl-char	Dec	Hex	Char	Dec	Hex	Char	Dec	Hex	Char
0	0	Null	NUL	CTRL-@	32	20	Space	64	40	@	96	60	`
1	1	Start of heading	SOH	CTRL-A	33	21	!	65	41	A	97	61	a
2	2	Start of text	STX	CTRL-B	34	22	"	66	42	B	98	62	b
3	3	End of text	ETX	CTRL-C	35	23	#	67	43	C	99	63	c
4	4	End of xmit	EOT	CTRL-D	36	24	\$	68	44	D	100	64	d
5	5	Enquiry	ENQ	CTRL-E	37	25	%	69	45	E	101	65	e
6	6	Acknowledge	ACK	CTRL-F	38	26	&	70	46	F	102	66	f
7	7	Bell	BEL	CTRL-G	39	27	'	71	47	G	103	67	g
8	8	Backspace	BS	CTRL-H	40	28	(	72	48	H	104	68	h
9	9	Horizontal tab	HT	CTRL-I	41	29	)	73	49	I	105	69	i
10	0A	Line feed	LF	CTRL-J	42	2A	*	74	4A	J	106	6A	j
11	0B	Vertical tab	VT	CTRL-K	43	2B	+	75	4B	K	107	6B	k
12	0C	Form feed	FF	CTRL-L	44	2C	,	76	4C	L	108	6C	l
13	0D	Carriage feed	CR	CTRL-M	45	2D	-	77	4D	M	109	6D	m
14	0E	Shift out	SO	CTRL-N	46	2E	.	78	4E	N	110	6E	n
15	0F	Shift in	SI	CTRL-O	47	2F	/	79	4F	O	111	6F	o
16	10	Data line escape	DLE	CTRL-P	48	30	0	80	50	P	112	70	p
17	11	Device control 1	DC1	CTRL-Q	49	31	1	81	51	Q	113	71	q
18	12	Device control 2	DC2	CTRL-R	50	32	2	82	52	R	114	72	r
19	13	Device control 3	DC3	CTRL-S	51	33	3	83	53	S	115	73	s
20	14	Device control 4	DC4	CTRL-T	52	34	4	84	54	T	116	74	t
21	15	Neg acknowledge	NAK	CTRL-U	53	35	5	85	55	U	117	75	u
22	16	Synchronous idle	SYN	CTRL-V	54	36	6	86	56	V	118	76	v
23	17	End of xmit block	ETB	CTRL-W	55	37	7	87	57	W	119	77	w
24	18	Cancel	CAN	CTRL-X	56	38	8	88	58	X	120	78	x
25	19	End of medium	EM	CTRL-Y	57	39	9	89	59	Y	121	79	y
26	1A	Substitute	SUB	CTRL-Z	58	3A	:	90	5A	Z	122	7A	z
27	1B	Escape	ESC	CTRL-[	59	3B	;	91	5B	[	123	7B	{
28	1C	File separator	FS	CTRL-\	60	3C	<	92	5C	\	124	7C	
29	1D	Group separator	GS	CTRL-]	61	3D	=	93	5D	]	125	7D	}
30	1E	Record separator	RS	CTRL-^	62	3E	>	94	5E	^	126	7E	~
31	1F	Unit separator	US	CTRL-`	63	3F	?	95	5F	_	127	7F	DEL

# Q<sub>30</sub> is considered a benchmark for sequence quality in next-generation sequencing

A quality score of 99% (Q<sub>30</sub>) will have an incorrect base call probability of 1 in 1000

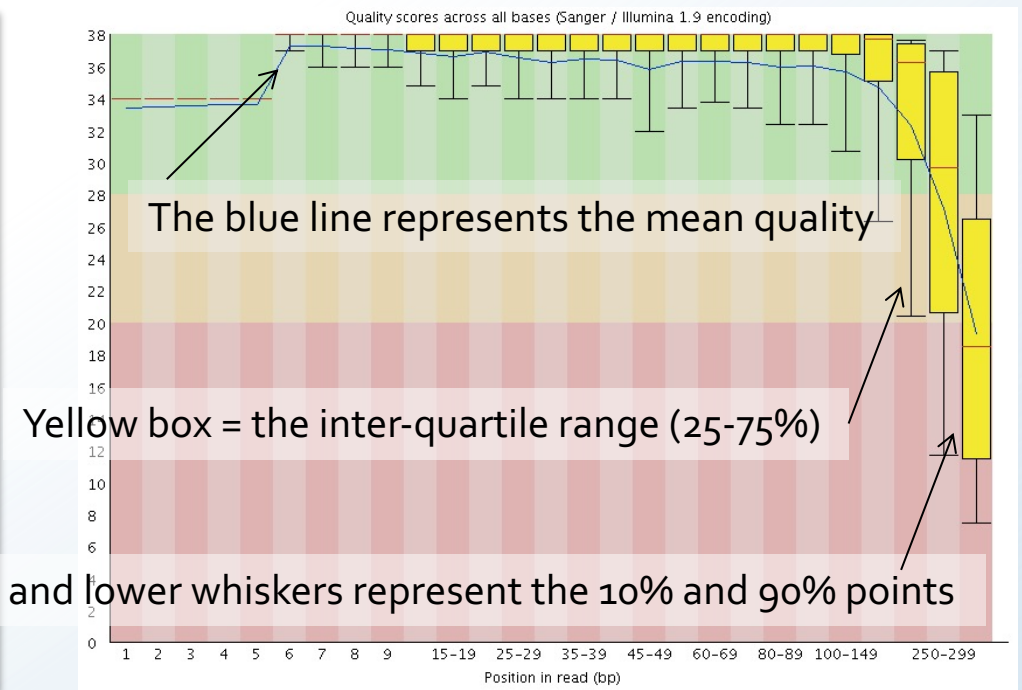
## Relationship Between Sequencing Quality Score and Base Call Accuracy:

Quality Score	Probability of Incorrect Base Call	Inferred Base Call Accuracy
10 (Q <sub>10</sub> )	1 in 10	90%
20 (Q <sub>20</sub> )	1 in 100	99%
30 (Q <sub>30</sub> )	1 in 1000	99.9%

# When performing Quality Control (QC) you generate a general summary of the input data

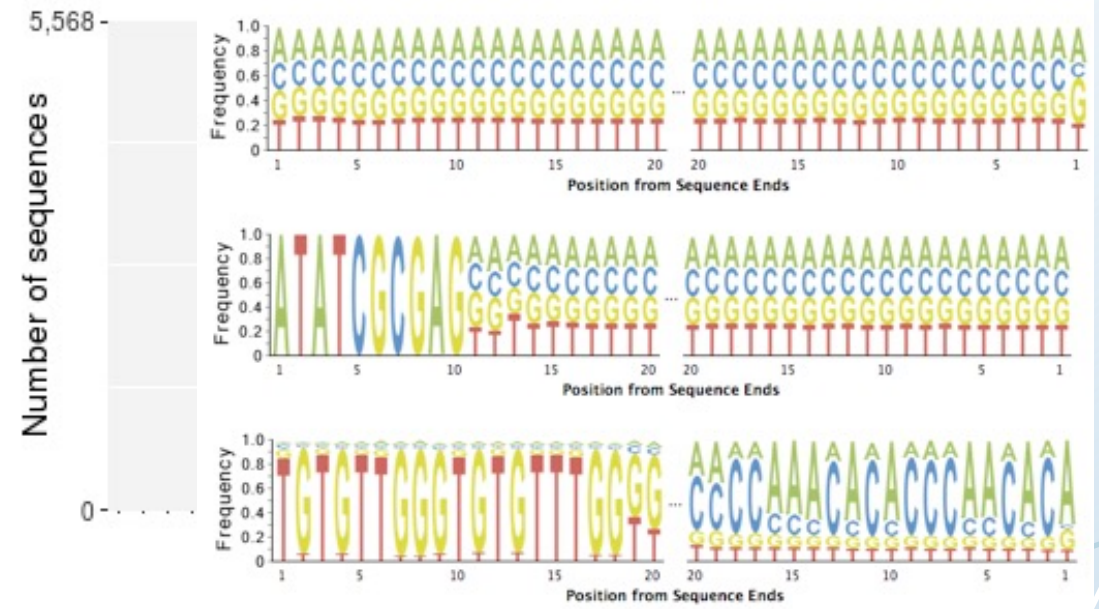
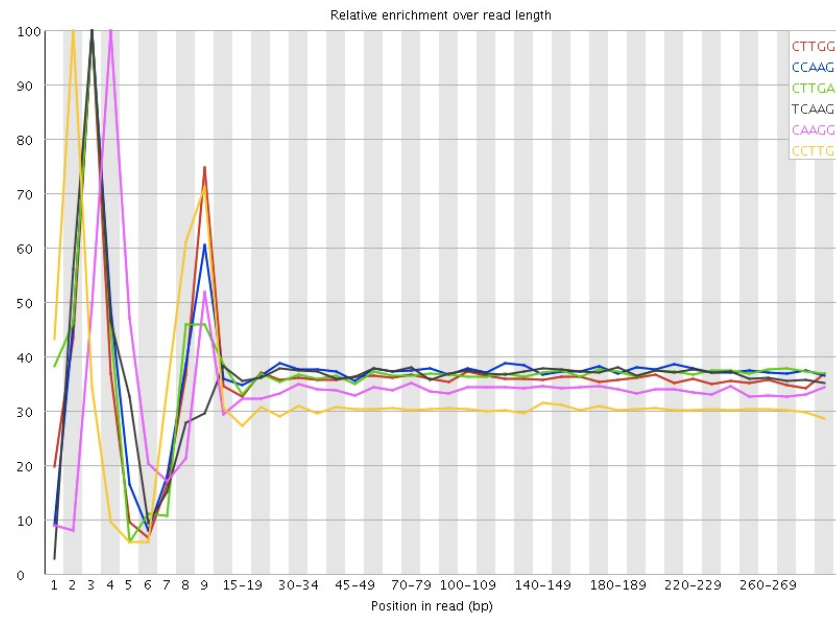
The range of quality values across all bases at each position in the FASTQ file

Measure	Value
Filename	CAV3_S1_L001_R1_001.fastq.gz
File type	Conventional base calls
Encoding	Sanger / Illumina 1.9
Total Sequences	2172538
Filtered Sequences	0
Sequence length	35-301
%GC	41



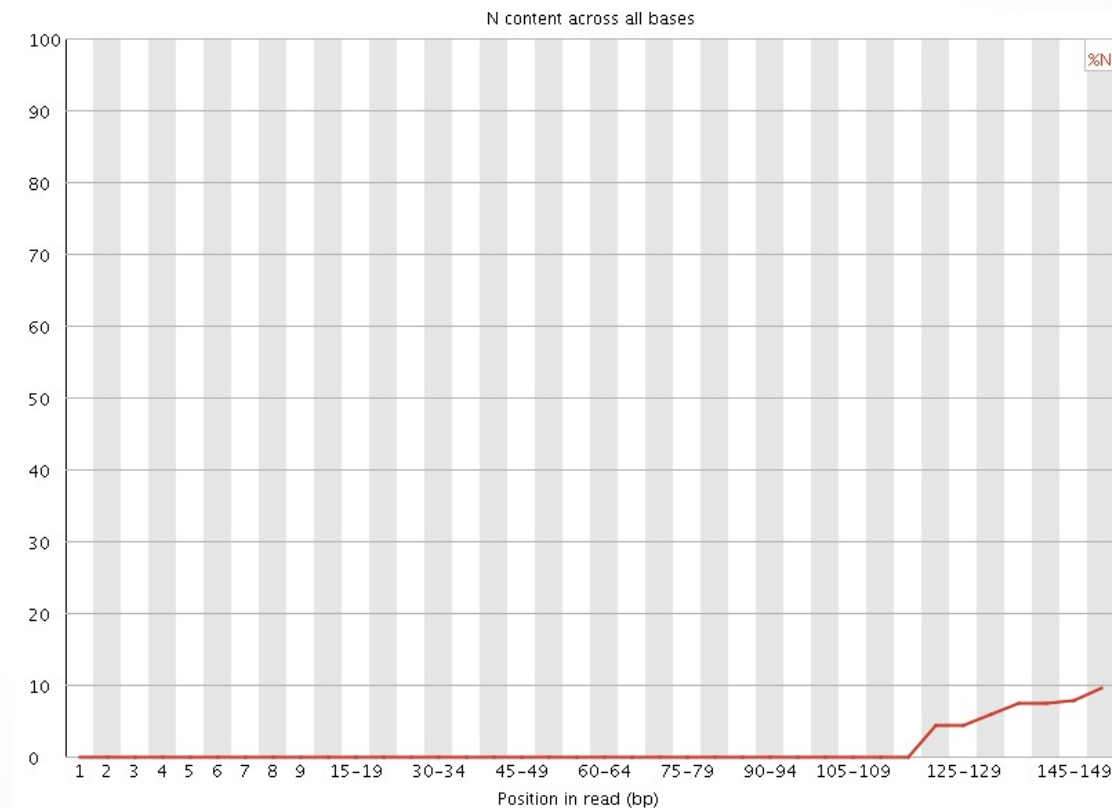
# Low quality sequences can cause problems during downstream analysis

Analyzing K-mer content may identify artifacts in sequence reads, eg. multiplex identifiers, adapters, and primer sequences



# Ambiguous bases (Ns) arises when a sequencer is unable to make a base call with sufficient confidence

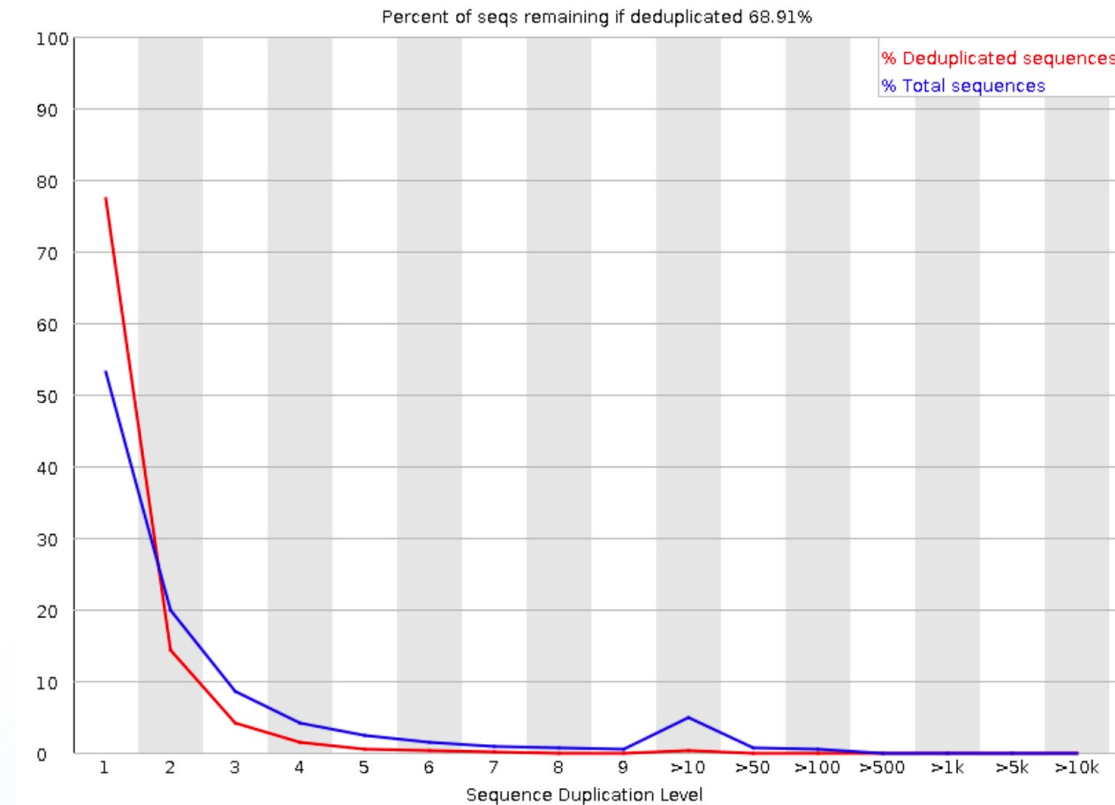
A high number of Ns can be a sign for a low quality sequence and can cause problems during downstream analysis





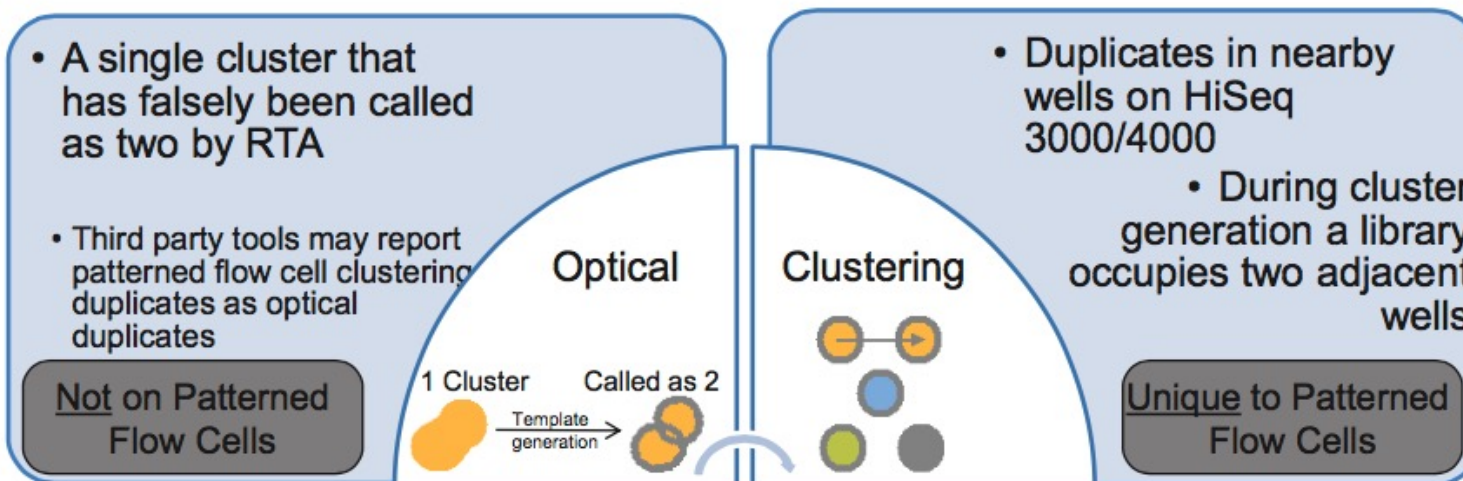
# Identification of sequence duplication since ideally, no reads should not start at the same position and have the same errors

Most likely to indicate some kind of enrichment bias



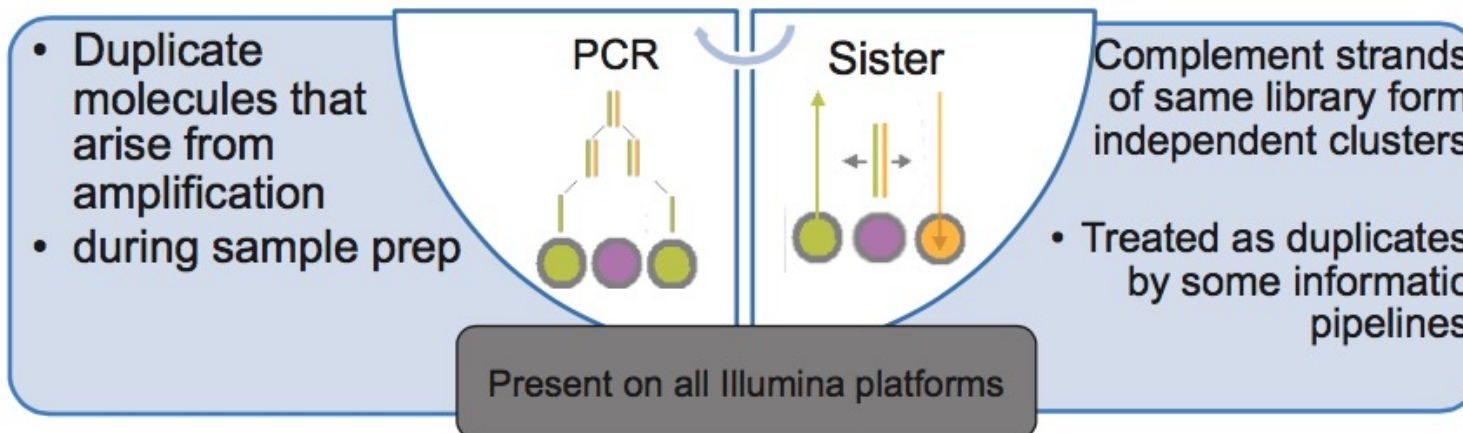
# Duplicates in NGS data

Optical duplicates



Local re-clustering

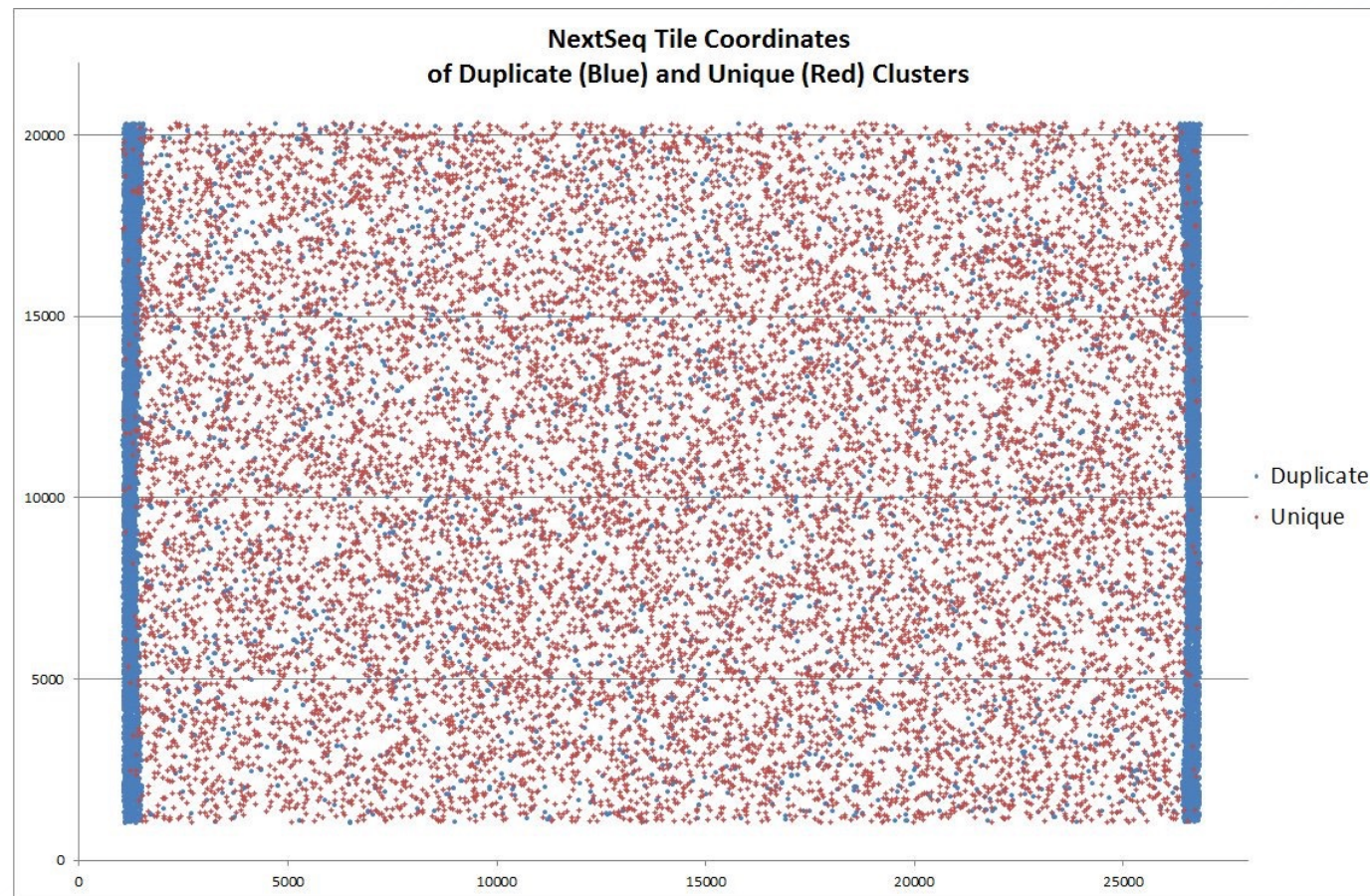
PCR and enrichment duplicates



Exclusion Amplification duplicates

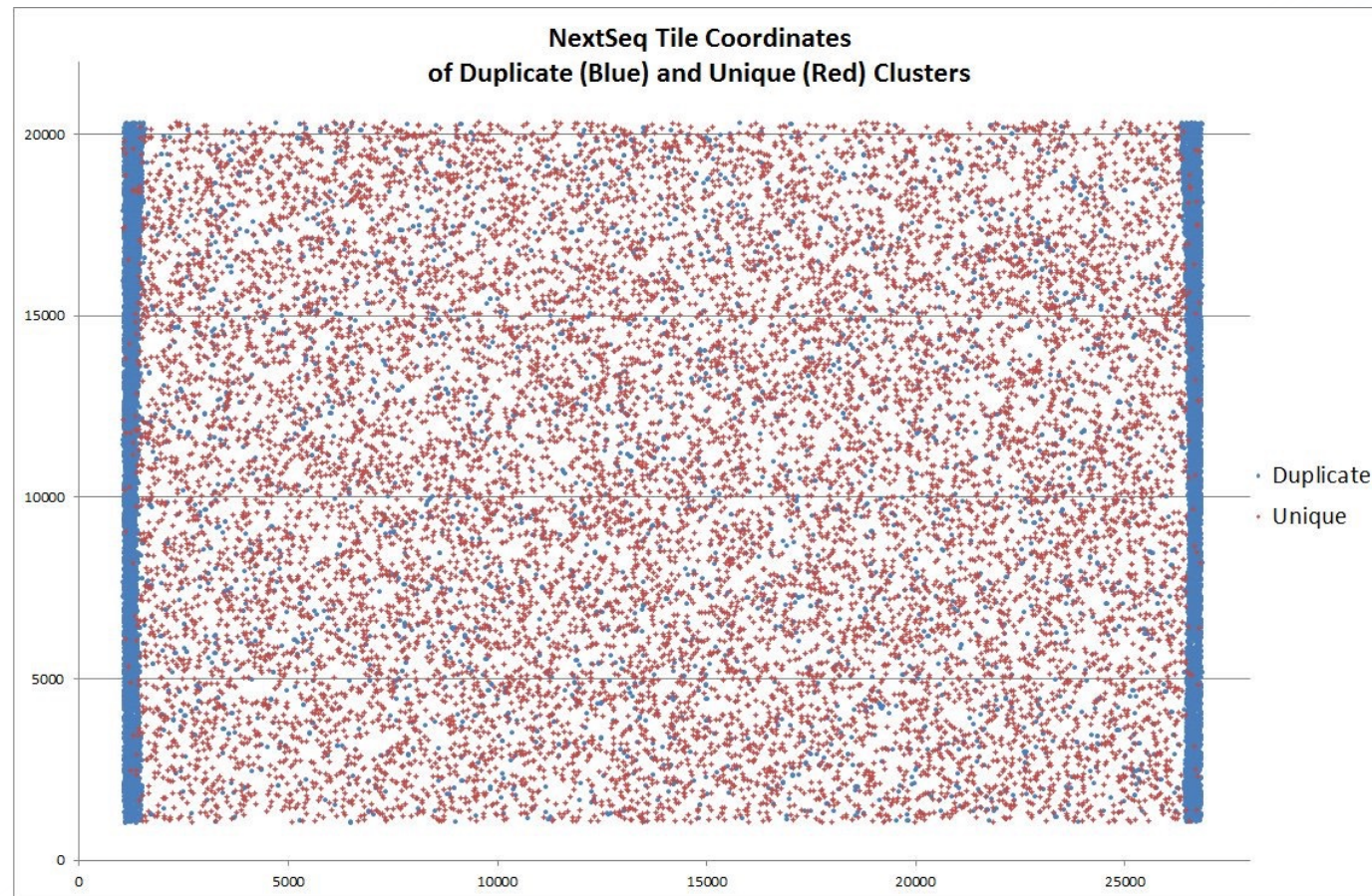
# Duplicates in NGS data

Tile-edge duplicates may account for >80% of the duplicates



# Remove duplicates from NGS data

Clumpify removes duplicates and reorders reads to maximize gzip compression



# Simple FASTQ/FASTA manipulations using SeqKit

---

SeqKit is a cross-platform ultrafast comprehensive toolkit that can perform common manipulations of FASTA/Q files

Eg converting, searching, filtering, deduplication, splitting

Works on all major operating systems, including Windows, Linux, and Mac OS X, and can be directly used without any dependencies

Easy to use on command line:

```
$ seqkit <subcommand>
```

List of subcommands can be found here: <https://github.com/shenwei356/seqkit>