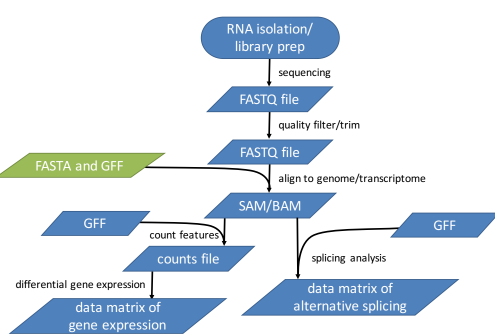
Common file formats, Part 1 FASTA/FASTQ

The FASTA file format

- Text-based format for representing
 - nucleotide sequence OR
 - peptide sequence (single letter codes)
- Each sequence in the file begins with a unique header line denoted with a ">"
- The following line(s) are are sequence, typically between 60 and 130 characters in length (may include spaces and/or numbers)



What does a FASTA file look like?

>NODE_1140748_length_208_cov_4.298077

GTATATTAGAAGGGCCGCGCGCGCATGAGATGGGTGACAGTACACTTTCCATGCAAGAACG GGCGGGTTTGTAATATTCCTTAAATTATTGTCAGAAACTCTGTGATGGAGACATTGACCT CAGTTAATTAGTCTGGGTATTGCTGTCTGATTTAATAAGACACCAAACCAGTCTCCCTGG CGACAGGAAGTGCTGCATGCCACTGCATTTGTTGCAGCGGTGTTCTAGAGCGGTCA CCCTGAGGGATGCTGC

>NODE_1140775_length_818_cov_9.328851

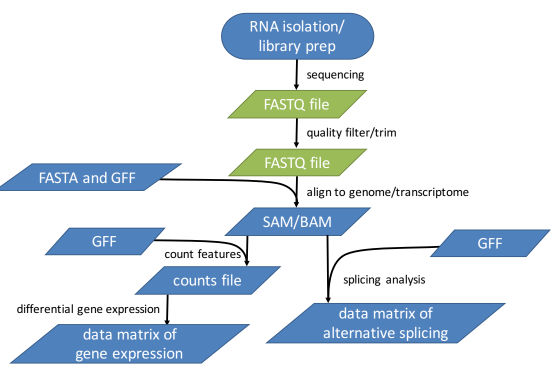
>NODE_1141553_length_172_cov_7.779070

The FASTQ file format

 FASTQ is a modified form of FASTA

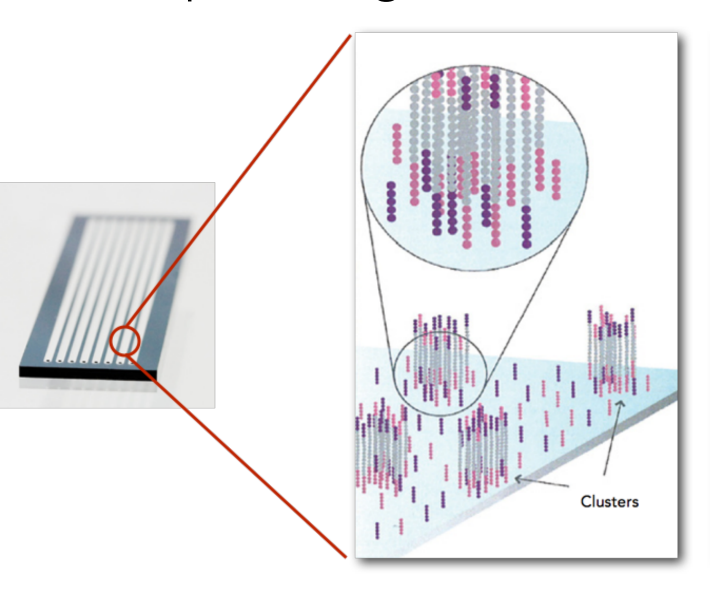
Goal: include sequencing quality scores

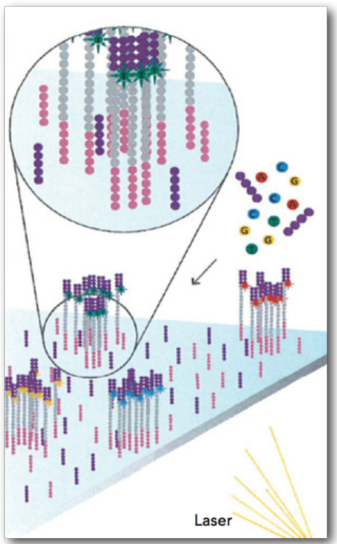
 Has become the standard for storing data generated by NGS data



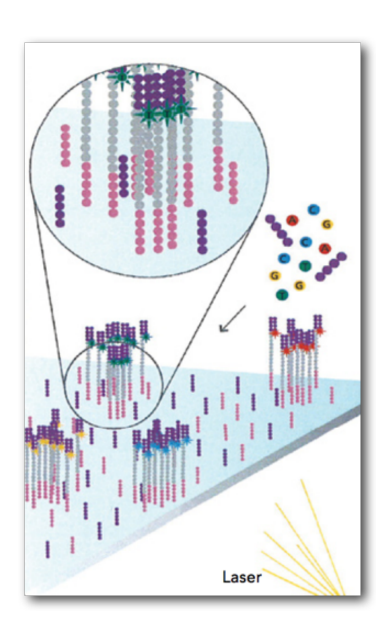
 To understand quality scores, it's helpful to understand how sequencing data are generated

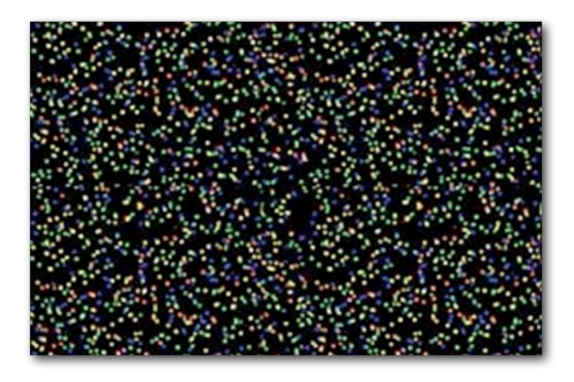
Sequencing on Illumina's flow cell





Sequencing on Illumina's flow cell

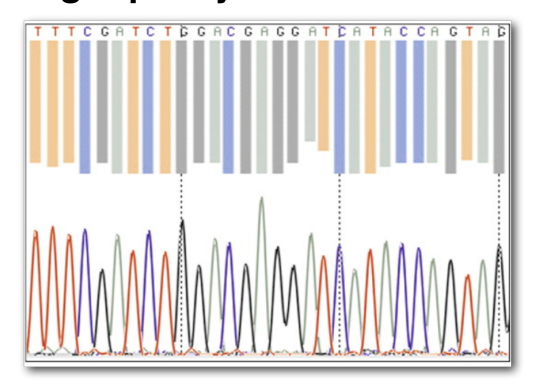




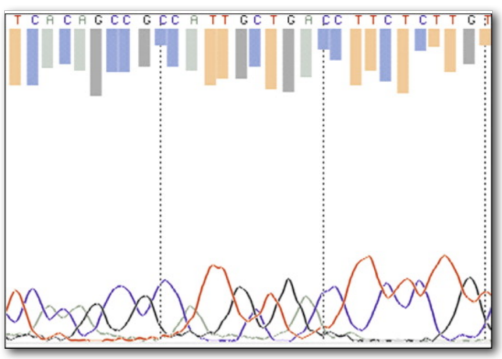


What does high/low quality data look like?

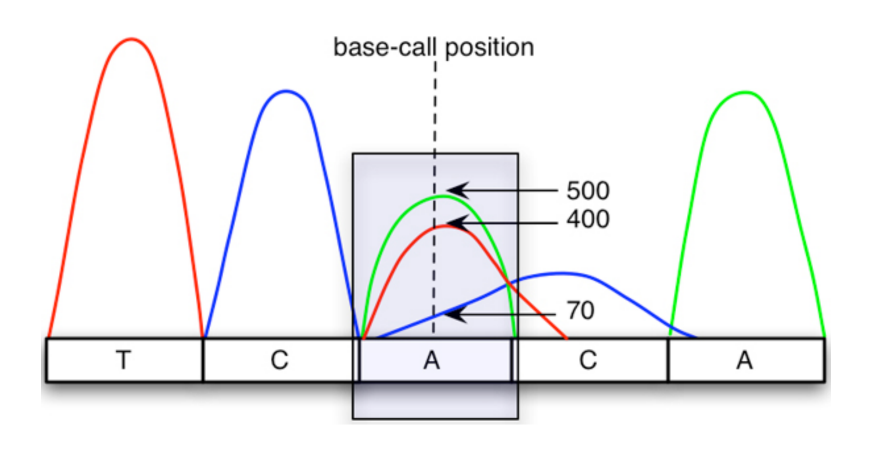
High quality data



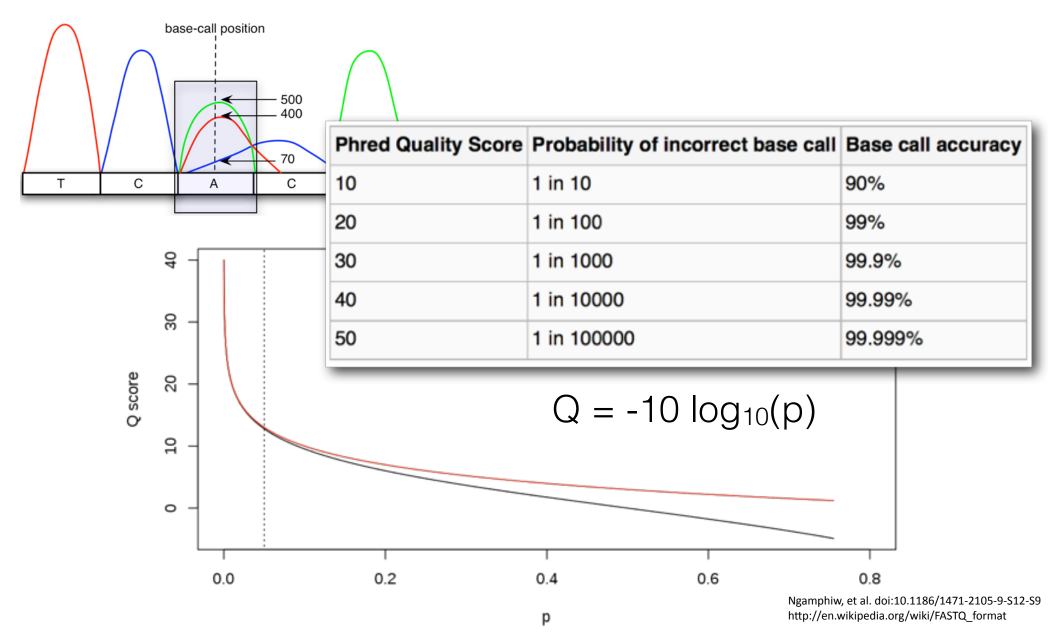
Lower quality data



Each base call is given a score: Phred quality score



Each base call is given a score: Phred quality score



What does a FASTQ file look like?

File uses four lines per read

- 1. Begins with "@" followed by unique sequence identifier
- 2. Sequence of the read
- Begins with "+" and is optionally followed by identifier again
- 4. Encodes the quality scores (must be same length as line 2)

Notice phred scores encoded by a single letter – how can we decode?

First, decode ASCII

_															
0	<nul></nul>	32	<spc></spc>	64	@	96	*	128	Ä	160	+	192	خ	224	‡
1	<soh></soh>	33	!	65	Α	97	a	129	Å	161	0	193	i	225	.
2	<stx></stx>	34	"	66	В	98	b	130	Ç	162	¢	194	\neg	226	,
3	<etx></etx>	35	#	67	C	99	С	131	É	163	£	195	\checkmark	227	,,
4	<eot></eot>	36	\$	68	D	100	d	132	Ñ	164	§	196	f	228	%0
5	<enq></enq>	37	%	69	E	101	e	133	Ö	165	•	197	≈	229	Â
6	<ack></ack>	38	&	70	F	102	f	134	Ü	166	1	198	Δ	230	Ê
7	<bel></bel>	39	'	71	G	103	g	135	á	167	ß	199	«	231	Á
8	<bs></bs>	40	(72	Н	104	h	136	à	168	®	200	>>	232	Ë
9	<tab></tab>	41)	73	I	105	i	137	â	169	©	201	***	233	
10	<lf></lf>	42	*	74	J	106	j	138	ä	170	TM	202		234	Í
11	<vt></vt>	43	+	75	K	107	k	139	ã	171	,	203	À	235	Î
12	<ff></ff>	44	,	76	L	108	1	140	å	172		204	Ã	236	Ĭ
13	<cr></cr>	45	-	77	M	109	m	141	ç	173	≠	205	Õ	237	Ì
14	<50>	46		78	N	110	n	142	é	174	Æ	206	Œ	238	Ó
15	<si></si>	47	/	79	0	111	0	143	è	175	Ø	207	œ	239	ô
16	<dle></dle>	48	0	80	Р	112	р	144	ê	176	00	208	-	240	•
17	<dc1></dc1>	49	1	81	Q	113	q	145	ë	177	±	209	_	241	Ò
18	<dc2></dc2>	50	2	82	R	114	r	146	í	178	≤	210	w	242	Ú
19	<dc3></dc3>	51	3	83	S	115	S	147	ì	179	≥	211	"	243	Û
20	<dc4></dc4>	52	4	84	T	116	t	148	î	180	¥	212	`	244	Ù
21	<nak></nak>	53	5	85	U	117	u	149	ï	181	μ	213	,	245	1
22	<syn< td=""><td>54</td><td>6</td><td>86</td><td>V</td><td>118</td><td>V</td><td>150</td><td>ñ</td><td>182</td><td>д</td><td>214</td><td>÷</td><td>246</td><td>^</td></syn<>	54	6	86	V	118	V	150	ñ	182	д	214	÷	246	^
23	<etb></etb>	55	7	87	W	119	W	151	ó	183	Σ	215	♦	247	~
24	<can></can>	56	8	88	X	120	X	152	ò	184	Π	216	ÿ	248	-
25		57	9	89	Υ	121	У	153	ô	185	п	217	Ÿ	249	·
26		58	:	90	Z	122	z	154	Ö	186	ſ	218	/	250	.
27	<esc></esc>	59	;	91]	123	{	155	õ	187	a	219	€	251	۰
28	<fs></fs>	60	<	92	\	124		156	ú	188	0	220	<	252	,
29	<gs></gs>	61	=	93]	125	}	157	ù	189	Ω	221	>	253	"
30	<rs></rs>	62	>	94	^	126	~	158	û	190	æ	222	fi	254	ι Ι
31	<us></us>	63	?	95	_	127		159	ü	191	Ø	223	fl	255	v

Each character has been assigned a numerical value that can be encoded by 8 bits 8 bits = 28 combinations = 256 possibilities

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0	<nul></nul>	32	<spc></spc>	64	@	96	•	128	Ä	160	+	192	ċ	224	#
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14	<s0></s0>	46		78	N	110	n	142	é	174	Æ	206	Œ	238	Ó
15	<si></si>	47	/	79	0	111	0	143	è	175	Ø	207	œ	239	ô
16	<dle></dle>	48	0	80	Р	112	p	144	ê	176	00	208	-	240	•
17	<dc1></dc1>	49	1	81	Q	113	q	145	ë	177	±	209	_	241	Ò
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22	<syn< td=""><td>54</td><td>6</td><td>86</td><td>V</td><td>118</td><td>V</td><td>150</td><td>ñ</td><td>182</td><td>д</td><td>214</td><td>÷</td><td>246</td><td>^</td></syn<>	54	6	86	V	118	V	150	ñ	182	д	214	÷	246	^
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25		57	9	89	Υ	121	У	153	ô	185	П	217	Ÿ	249	·
26		58	:	90	Z	122	z	154	Ö	186	ſ	218	/	250	.
27	<esc></esc>	59	;	91]	123	{	155	õ	187	a	219	€	251	۰
28	<fs></fs>	60	<	92	\	124	1	156	ú	188	0	220	<	252	,
29	<gs></gs>	61	=	93]	125	}	157	ù	189	Ω	221	>	253	"
30	<rs></rs>	62	>	94	^	126	~	158	û	190	æ	222	fi	254	.
31	<us></us>	63	?	95	_	127		159	ü	191	Ø	223	fl	255	v

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 $\mathbf{01111001} \quad \mathbf{0} \times 2^{7} + \mathbf{1} \times 2^{6} + \mathbf{1} \times 2^{5} + \mathbf{1} \times 2^{4} + \mathbf{1} \times 2^{3} + \mathbf{0} \times 2^{2} + \mathbf{0} \times 2^{1} + \mathbf{1} \times 2^{0} = 121 = y$

Then convert quality score

```
@NS500451:154:HWKTMBGXX:1:11101:10065:1121 1:N:0:TAGAACAC
AGGTTGCTATGAAATTTTAGTTGTCGTAGTAGGCAAACAATAAGGAATGTTGATCCAATAATTACATGGAGTCCATGGAA
```

Quality scores

```
!"#$%&'()*+,-./0123456789:;<=>?@ABCDEFGHIJKLMNOPQRSTUVWXYZ[\]^ `abcdefghijklmnopqrstuvwxyz{|}~
33
                      59
                          64
                                  73
                                                            104
                                                                              126
S - Sanger
          Phred+33, raw reads typically (0, 40)
X - Solexa
               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)
L - Illumina 1.8+ Phred+33, raw reads typically (0, 41)
```

ASCII values 33 through 73 correspond to phred scores 0 through 40

$$E' = 69$$

(E' = 69) 69 - 33 = 36 $p_{error} = 0.025\%$ Base call accuracy: 99.975%