Background information

# Introduction to bioinformatics (NGS data analysis)

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## Got your sequencing data - now, what to do with it?

```
File size: several Gb
```

Background information

Number of lines: >1.000.000

```
@MO2443:17:000000000-ABPBW:1:1101:12675:1533 1:N:0:1
TCGATAATTCTTACTTTCTCTCTGGTCTGAGCGTTTCACATCAACGACAAGCTCGA
TTTTTTTTTTTTTTTTT
8B6-@-,CFFED9CFAE@@C6;@,CFEEF9<@6FGGF9F<CC,,CB,@::8CF,6+
,,3733>>00,,,3880,,8*,773333,3,333738,*,,,,,76,,2,,2
0*).1.))(0*)***
@MO2443:17:000000000-ABPBW:1:1101:18658:1535 1:N:0:1
TCCCTAATTCTCTGTCTTCAAATTTTCCTTCTAAATCGTCCCTCGTTTCTACCT
TTTTCTTCTTTTTCT
-<<9-@CCEF9CE-<,,,,,;,,<C,=,6,C9,C<=C,,,;,86C,6:C,,,;<;,,
,,,,5,5:,,9++4,,,:,,,,,,,,38,853,5,,3,,7,,,6,,,,,7,,,,
+0.()+++)11.*)*
                                    4 D > 4 B > 4 B > 4 B > B
```

# Before library preparation

What you need to know to steer your way through the analysis

- Research question
  - Identify adaptive genes
  - *De novo* genome assembly
  - Population genetic structure
  - Phylogenetic relation
- Experimental design
  - Number of individuals
  - Treatment of samples (e.g. heat stress)
- Sample collection
  - Samples degraded (e.g. stored in Formalin)
  - Tissue (reproductive, vegetative)

## Library preparation

- DNA-seq, RNA-seq, Bis-Seq, Chip-Seq...
  - RNA reads (which lack introns) require splice-aware mappers.
  - Bis-seq changes GC ratio (bisulphite converts cytosine to uracil, but leaves 5-methylcytosine unaffected)
  - Chip-Seq enriches binding-sites of DNA-associated proteins
- Pooled samples?
  - Demultiplexing
  - Remove barcodes
- Adapter sequences that have to be trimmed off?
- Targeted coverage

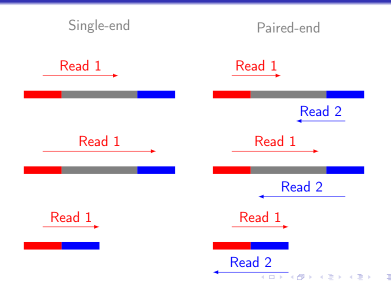
## Single- or Paired end sequencing, read length

#### Library fragment

Adapter
Flowcell/bead binding sequences
Amplification primers
Sequencing primers
Barcodes

Adapter
Flowcell/bead binding sequences
Amplification primers
Sequencing primers
Barcodes

# Single- or paired-end sequencing, read length - why does it matter



Platform	Max. length	Reads/run	Consideration
Illumina	2×150	5 billion	
HiSeq series			
Illumina	2×300	25 million	
MiSeq series			
Illumina			
NextSeq series	2×150	400 million	
Roche 454	700	0.7 million	High error rate

# Primary analysis

- Demultiplexing
- Adapter trimming
- Quality control

# Demultiplexing of pooled samples (if barcoded inline)

AATTANNNNNNNNNNNNNN File 1

Background information

AGTCGNNNNNNNNNNNNNNN File 2

AGTCGNNNNNNNNNNNNNN File 2

AATTANNNNNNNNNNNNNN File 1

AGTCGNNNNNNNNNNNNNN File 2

References

## Trimmig: Adapter removal

Mostly 3'adapters disturb assembly and alignment

GATTTGGGGTTCAANNNNNNNNNNNATTAGTATCGAT

GATTTGGGGTTCAANNNNNNNATTAGTATCGAT

TTGGGGTTCAANNNNNNNATTAGTATCGAT

GATTTGGGGTTCAANNNNNNNNNNNATTAGTATCGAT

**ATTTGGGGTTCAANNNNNNNNNATTAGTATCGAT** 

**GATTTGGGGTTCAANNNNNNNNNNTTAGTATCGAT** 

## Fastq file - 4 lines for each read

```
@HWI-ST141_0365:2:1101:2983:2114#TTAGGC/1
GATTTGGGGTTCAAATTAGTATCGATCAAATAGTAAATCCATTTGTTCAACTC
+
!''*((((***+))%%%++)(%%%%).1***-+*''))**55CCF>>>>>CC
```

- sequence id (specifications can differ slightly between sequencing platforms)
  - =@=instrument name : flowcell lane : tile number: flowcell x coordinate : flowcell y coordinates : #barcode sequence: pair number for paired-end sequencing
- 2 sequence
- 3 + optionally followed by sequence identifier again
- 4 quality scores

# Trimmig of low-quality bases

Background information

- Trim bases with a Phred quality score <20
- $Quality = -10 * log_{10}P$

Phred Score	Probability of incorrect base	Base call accuracy
10	1 in 10	90%
20	1 in 100	99%
30	1 in 1000	99.9%

# Fastq file contains both sequence reads and base quality scores

```
Fastq file

@SEQ_ID

GATTTGGGGTTCAAATTAGTATCGATCAAATAGTAAATCCATTTGTTCAACTC
+
!''**((((***+))%%%++)(%%%%).1***-+*''))**55CCF>>>>>CC

Fasta file

>SEQ_ID

GATTTGGGGTTCAAATTAGTATCGATCAAATAGTAAATCCATTTGTTCAACTC
```

## Base qualities are encoded in ascii format

ASCII stands for American Standard Code for Information Interchange. An ASCII code is the numerical representation for a character.

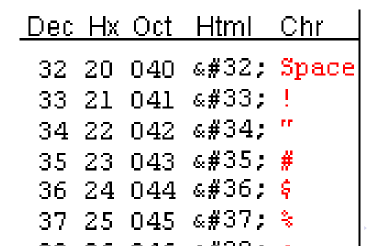
```
Dec Hx Oct Html Chrl Dec Hx Oct Html Chr
Dec Hx Oct Char
                                      Dec Hx Oct Html Chr
    0 000 NUL (null)
                                                             64 40 100 6#64; 8
                                       32 20 040   Space
                                                                                96 60 140 6#96:
   1 001 SOH (start of heading)
                                       33 21 041 4#33; !
                                                             65 41 101 6#65; A
                                                                                97 61 141 6#97; 8
    2 002 STX (start of text)
                                       34 22 042 4#34; "
                                                             66 42 102 a#66; B
                                                                                98 62 142 6#98; b
    3 003 ETX (end of text)
                                       35 23 043 4#35; #
                                                             67 43 103 a#67; C
                                                                                99 63 143 4#99; 0
                                                                                100 64 144 @#100; 4
    4 004 EOT (end of transmission)
                                       36 24 044 4#36; $
                                                             68 44 104 D D
                                       37 25 045 6#37; %
                                                                               101 65 145 @#101; e
    5 005 ENQ (enquiry)
                                                             69 45 105 E E
    6 006 ACK (acknowledge)
                                       38 26 046 6#38; 6
                                                             70 46 106 6#70; F
                                                                               102 66 146 @#102; f
   7 007 BEL (bell)
                                                             71 47 107 6#71; G 103 67 147 6#103; g
                                       39 27 047 6#39; 1
    8 010 BS
              (backspace)
                                       40 28 050 6#40; (
                                                             72 48 110 6#72; H
                                                                               104 68 150 6#104; h
                                                             73 49 111 6#73; I 105 69 151 6#105; i
   9 011 TAB (horizontal tab)
                                       41 29 051 6#41; )
              (NL line feed, new line) 42 2A 052 6#42; *
                                                             74 4A 112 6#74; J 106 6A 152 6#106; j
   A 012 LF
                                                             75 4B 113 4#75; K 107 6B 153 4#107; k
11 B 013 VT
              (vertical tab)
                                       43 2B 053 6#43; +
              (NP form feed, new page) 44 2C 054 4#44;
                                                             76 4C 114 4#76; L
12 C 014 FF
                                                                               108 6C 154 6#108; 1
13 D 015 CR
              (carriage return)
                                       45 2D 055 6#45; -
                                                             77 4D 115 6#77; M 109 6D 155 6#109; M
14 E 016 SO
                                       46 2E 056 6#46; .
                                                             78 4E 116 6#78; N 110 6E 156 6#110; n
              (shift out)
                                                             79 4F 117 6#79; 0 111 6F 157 6#111; 0
15 F 017 SI
              (shift in)
                                       47 2F 057 6#47; /
16 10 020 DLE (data link escape)
                                       48 30 060 4#48; 0
                                                             80 50 120 6#80; P 112 70 160 6#112; P
                                                             81 51 121 6#81; Q 113 71 161 6#113; Q
17 11 021 DC1 (device control 1)
                                       49 31 061 4#49; 1
18 12 022 DC2 (device control 2)
                                       50 32 062 4#50; 2
                                                             82 52 122 6#82; R 114 72 162 6#114; r
                                       51 33 063 4#51; 3
                                                             83 53 123 4#83; $ 115 73 163 4#115; $
19 13 023 DC3 (device control 3)
20 14 024 DC4 (device control 4)
                                       52 34 064 4#52; 4
                                                             84 54 124 6#84; T 116 74 164 6#116; t
21 15 025 NAK (negative acknowledge)
                                       53 35 065 4#53; 5
                                                             85 55 125 6#85; U 117 75 165 6#117; U
                                       54 36 066 6#54; 6
                                                             86 56 126 6#86; V 118 76 166 6#118; V
22 16 026 SYN (synchronous idle)
                                       55 37 067 6#55; 7
                                                             87 57 127 6#87; ₩ 119 77 167 6#119; ₩
23 17 027 ETB (end of trans. block)
24 18 030 CAN (cancel)
                                       56 38 070 4#56; 8
                                                             88 58 130 6#88; X 120 78 170 6#120; X
25 19 031 EM
              (end of medium)
                                       57 39 071 6#57; 9
                                                             89 59 131 6#89; Y
                                                                               121 79 171 6#121; Y
                                       58 3A 072 6#58; :
                                                             90 5A 132 6#90; Z 122 7A 172 6#122; Z
26 1A 032 SUB (substitute)
27 1B 033 ESC (escape)
                                                                              123 7B 173 6#123;
                                       59 3B 073 4#59;;
                                                             91 5B 133 6#91; [
                                                             92 5C 134 6#92; \
                                                                               124 7C 174 6#124;
28 1C 034 FS
              (file separator)
                                       60 3C 074 4#60; <
                                                             93 5D 135 6#93; ] 125 7D 175 6#125;
29 1D 035 GS
              (group separator)
                                       61 3D 075 4#61; =
30 1E 036 RS
              (record separator)
                                       62 3E 076 4#62; >
                                                             94 5E 136 6#94; A 126 7E 176 6#126;
                                                             95 5F 137 6#95; _ 127 7F 177 6#127; DEL
31 1F 037 US
              (unit separator)
                                       63 3F 077 4#63; ?
```



## Base qualities are encoded in ascii format

Background information

ASCII stands for American Standard Code for Information Interchange. An ASCII code is the numerical representation for a character.



# ASCII encodings of sequencing platforms

```
!"#$%&'()*+,-./0123456789:;<=>?@ABCDEFGHIJKLMNOPQRSTUVWXYZ[\]^ `abcdefghijk
33
              59
                                       104
0.2......41
        Phred+33, raw reads typically (0, 40)
S - Sanger
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)
  (Note: See discussion above).
L - Illumina 1.8+ Phred+33, raw reads typically (0, 41)
```

Figure: Quality score encodings

# Quality control tool: FastQC

#### Informs on:

- Base quality
- Duplication
- Overrepresentation of sequences
  - contamination?
  - adapters?
- GC content (should be around 50%, in Bis-Seq lower)

## Quality before trimming

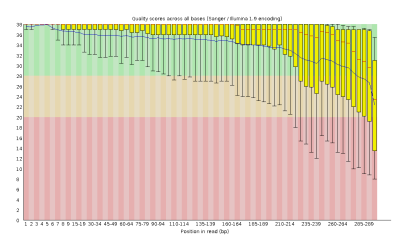


Figure: Base-quality generally decreases with increasing sequencing length

# Quality after trimming

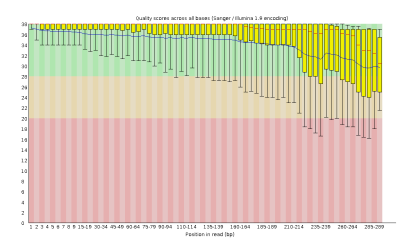
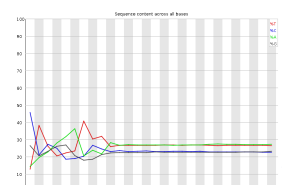


Figure: Quality after trimming

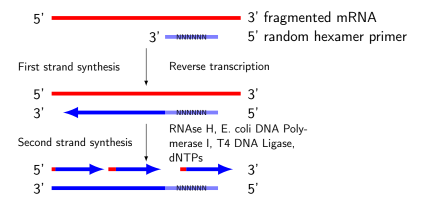
## Sequence bias

#### For example in:

- First bases of Illumina RNAseq due to 'random' hexamer primers for reverse transcription
- RADseq fragments (cutting sites)



## Hexamer primers for cDNA synthesis cause sequence bias



## **PCR Duplicates**

Duplicates are generally removed in quantitative analyses (e.g. RNA-seq)

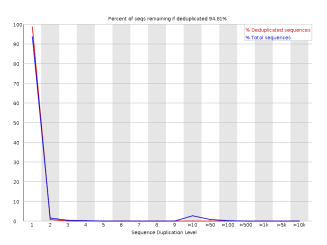


Figure: Duplication levels (FastQC output)

# De novo assembly

Task: Look for overlapping regions and create contigs (contiguous sequences)

- Genome assembly software
  - SOAP de NOVO
  - Velvet
  - MIRA (we use this one in the course)
- Transcriptome assembly software
  - Review: Martin and Wang, (2011)
  - Trinity
  - MIRA

# De novo assembly: Step by step



## De novo assembly: The N50 metric

 ${\sf N50}$  is a single measure of the contig length size distribution in an assembly

- Sort contigs in descending length order
- Size of contig above which the assembly contains at least 50% of the total length of all contigs

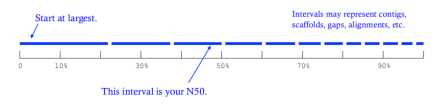


Figure: From Kane, N.C.

## Mapping against reference genome/transcriptome

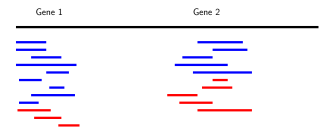
- Main purposes:
  - Identify variants (SNPs, InDels)



# Mapping against reference genome/transcriptome

■ Main purposes:

Quantify gene expression



Population 1

Population 2

# Mapping: global alignment

- Implemented in e.g. BWA, Bowtie2
- Needleman-Wunsch algorithm
- Aligns sequences in their full length
- Used for multiple sequence alignment when sequences are similar

Figure: Global alignment from rosalind.info

# Mapping: local alignment

- Smith-Waterman algorithm
- Clipping of terminal unmatched bases
- Only aligned bases contribute to the alignment's score
- Used to target smaller portions of genes with high similarity

```
\label{tccAGTTATGTCAGgggacacgagcatgcagagac} \texttt{|||||||||||} \texttt{aattgccgccgtcgttttcagCAGTTATGTCAGatc}
```

Figure: Local alignment from rosalind.info

## Splice-aware alignment of RNAseq reads to the genome

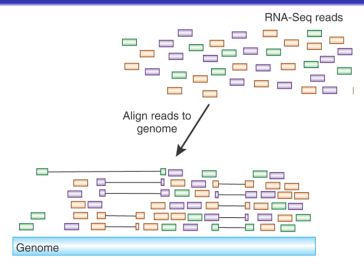


Figure: Adapted from Haas and Zody, (2010)

# Mapping: SAM/BAM files example

#### Output format of most alignment programs

- Header lines preceded by @
- One tab-delimited line per read

#### Figure: Example from http://samtools.sourceforge.net/SAM1.pdf

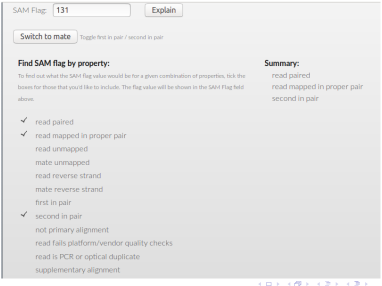
- SAM files are large
- BAM: Compressed binary versions, not human-readable

## Mapping: Mandatory fields in SAM files

Col	Field	Type	Regexp/Range	Brief description
1	QNAME	String	[!-?A-~]{1,255}	Query template NAME
2	FLAG	Int	[0,2 <sup>16</sup> -1]	bitwise FLAG
3	RNAME	String	\* [!-()+-<>-~][!-~]*	Reference sequence NAME
4	POS	Int	[0,2 <sup>31</sup> -1]	1-based leftmost mapping POSition
5	MAPQ	Int	[0,2 <sup>8</sup> -1]	MAPping Quality
6	CIGAR	String	\* ([0-9]+[MIDNSHPX=])+	CIGAR string
7	RNEXT	String	\* = [!-()+-<>-~][!-~]*	Ref. name of the mate/next read
8	PNEXT	Int	[0,2 <sup>31</sup> -1]	Position of the mate/next read
9	TLEN	Int	[-2 <sup>31</sup> +1,2 <sup>31</sup> -1]	observed Template LENgth
10	SEQ	String	\* [A-Za-z=.]+	segment SEQuence
11	QUAL	String	[!-~]+	ASCII of Phred-scaled base QUALity+33

Explanation of the flag field (click here: Link1, Link2)

# Mapping: Easy decoding of SAM flags



# Mapping: CIGAR string in SAM files

Op	BAM	Description
M	0	alignment match (can be a sequence match or mismatch)
I	1	insertion to the reference
D	2	deletion from the reference
N	3	skipped region from the reference
S	4	soft clipping (clipped sequences present in SEQ)
H	5	hard clipping (clipped sequences NOT present in SEQ)
P	6	padding (silent deletion from padded reference)
=	7	sequence match
X	8	sequence mismatch

# Mapping: CIGAR string example

```
RefPos: 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16
Ref: C C A T A C T G A A C T G A C T
Read: A C T A G A A T G G C T
```

CIGAR: 3M1I3M1D5M

## Variant calling

Consistent mismatches in the alignment indicate:

- Single Nucleotide Polymorphisms (SNPs)
- Insertions/Deletions (InDels)

## VCF file format

#### Variant call format

- described in http://www.1000genomes.org/node/101
- informs on location and quality of each SNP

#### VCF file information

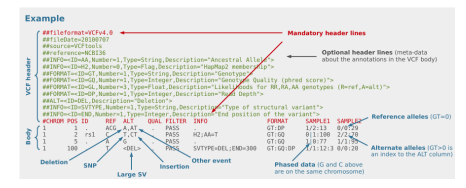


Figure: VCF file info from http://vcftools.sourceforge.net/VCF-poster.pdf

Phased alleles are on the same chromosome strand



#### VCF file information

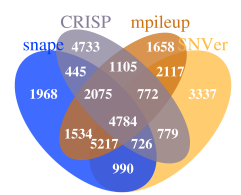


Figure: VCF file info from http://vcftools.sourceforge.net/VCF-poster.pdf

Phased alleles are on the same chromosome strand

# Identified SNPs vary between programs/algorithms

Venn diagram of the number of SNPs (coverage >400) called with four programs from the same alignment file (ddRAD tags mapped against the genome of Guppy).



Background information Primary analysis Secondary analysis Tertiary analysis Plan References

# Differential gene expression analysis

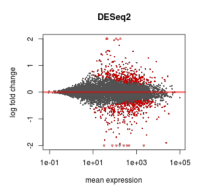


Figure: Log2 fold-change of expression over the mean of counts normalized by size factors. Differentially expressed genes (p<0.1) are red.

From the DESeq2 R package documentation



Background information Primary analysis Secondary analysis Tertiary analysis Plan References

# Clustering

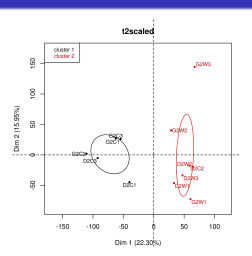


Figure: Multivariate grouping of stressed (W) and control (C) seagrass samples. Most variation is explained by the first principle component



# Visualizing differential expression

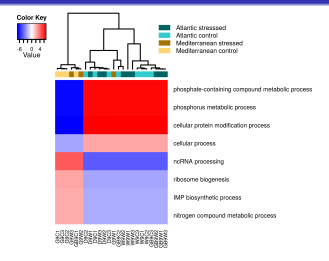
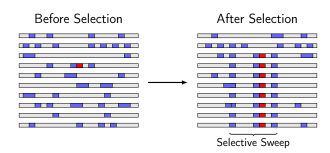


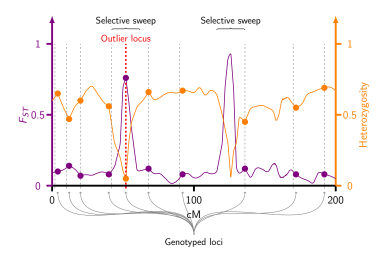
Figure: Heatmap of functions that were differentially expressed between Atlantic and Mediterranean seagrass samples.

# Outlier analysis



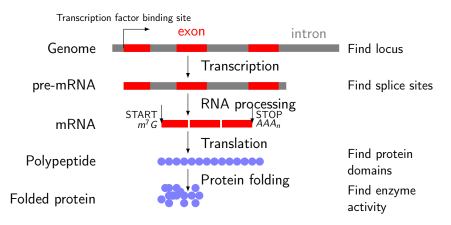
Based on Vitti et al., (2012)

#### Outlier detection



# Eukaryote genome annotation

#### Identify the strcuture and functional role



## Gene ontologies

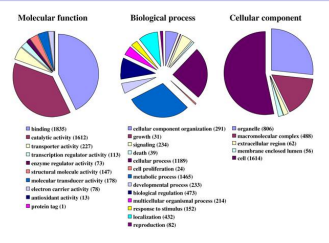


Figure: GO terms of unigenes in a moth genome

(Jacquin-Joly et al., 2012)



#### Cloud of GO term enrichments

Intercond memory operation
established of point indicates the control of the cont

# response to stimulus

cell wall organization or biogenesis cell wall modification callular carbohydrate biosynthetic proces. Proceedings of the control of the cont

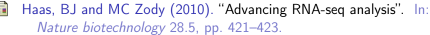
Figure: Term cloud of heat-responsive functions in seagrass



#### Bioinformatics-Practical

- Unix Tools (Martin)
- Trimming and Quality Control (Martin)
- Genome Assembly (Alexander)
- Mapping and Variant Calling (Martin)

# Background information



- Hansen, KD, SE Brenner, and S Dudoit (2010). "Biases in Illumina transcriptome sequencing caused by random hexamer priming". In: *Nucleic acids research* 38.12, e131–e131.
- Jacquin-Joly, E, F Legeai, N Montagné, C Monsempes, MC François, J Poulain, et al. (2012). "Candidate chemosensory genes in female antennae of the noctuid moth Spodoptera littoralis". In: *International journal of biological sciences* 8.7, p. 1036.
- Martin, J and Z Wang (2011). "Next-generation transcriptome assembly". In: *Nature Reviews Genetics*.
- Vitti, JJ, MK Cho, SA Tishkoff, and PC Sabeti (2012). "Human evolutionary genomics: ethical and interpretive issues". In: *Trends in Genetics* 28.3, pp. 137–145.

References