

# Exercise sheet 9: Data Driven Life Sciences

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## Exercise 1

**Question 1A** Arrange the following terms into their correct order in the Illumina sequencing method and describe each of them briefly:

- bridge amplification
- deblocking
- library preparation
- annealing of template strands to flow cell
- fluorescence detection

**Solution 1. Library preparation:**

A sequencing *library* gets *prepared* from a sample by fragmenting the original DNA and adding Illumina-specific adapter sequences to both ends of the fragments. The *library* is what gets read during sequencing.

**2. Template strand annealing**

The single-stranded library fragments are used as *template strands* in the sequencing and are *annealed* to primer sequences, which are bound to the *flow cell* and are complementary to the adapter sequences of the fragments.

**3. Bridge amplification**

After complementary strands have been synthesized and the templates been washed off, the now flow cell-bound fragments are *amplified* in several cycles of so-called *bridge-amplification* to form fragment colonies, or *clusters* on the flow cell to guarantee a detectable fluorescence signal during sequencing.

**4. Fluorescence detection**

Illumina-sequencing is a form of *sequencing-by-synthesis* in which the nucleotides incorporated into the growing strand are detected via attached *fluorophores*. After the first 3 steps, the following steps are iterated to sequence the entire read:

Modified nucleotides, containing a fluorescent group, are used to extend the strand, their blocking groups are cleaved from their 3'-OH groups.

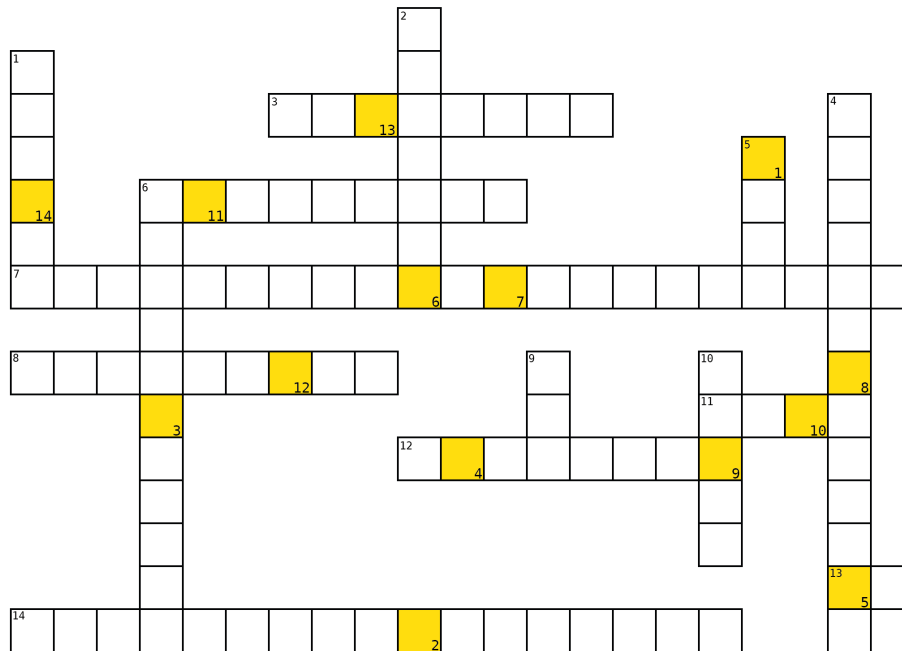
**5. Deblocking**

*Deblocking* is the removal of the fluorophore (blocking group). It is necessary before a new round of elongation by one nucleotide can begin.

More information about this topic can be found on the Illumina Webpage.

## Exercise 2

This exercise will be modified by end of today 30.06.22 to better fit the lecture recordings. Sorry for any inconvenience this may cause!



Solution:

1	2	3	4	5	6	7	8	9	10	11	12	13	14
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### Question 2A Solve the crossword puzzle!

Horizontal:

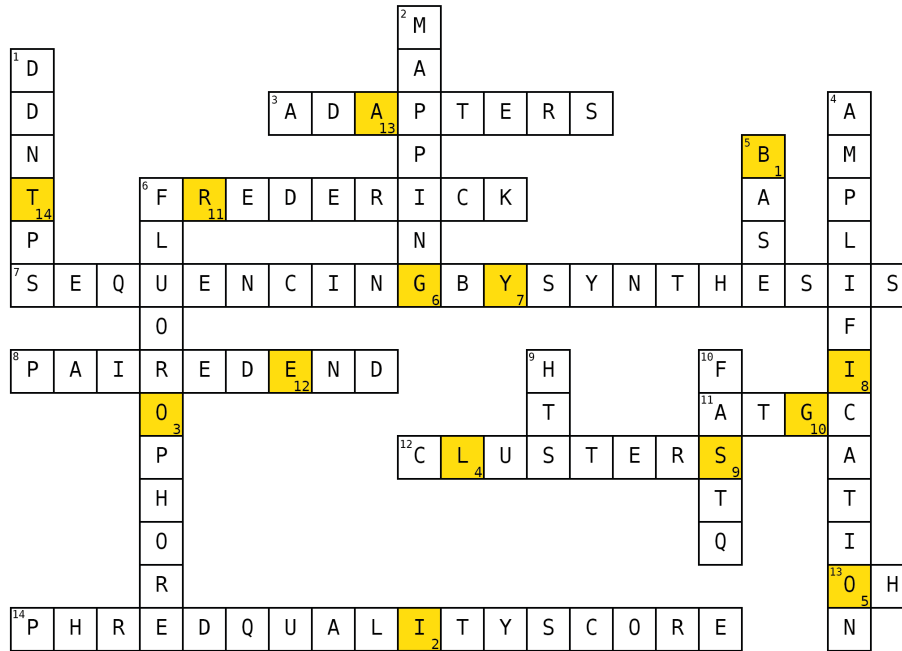
- 3. Added to DNA fragments during library preparation.
- 6. Has a sequencing method named after himself (first name)
- 7. Illumina/Pacbio way of determining the order of nucleotides in a DNA strand. (3 words)
- 8. More than just single-end. (2 words)
- 11. The alphabet of life.
- 12. Formed by bridge-amplification on Illumina flow-cells.
- 13. Keeps nucleic acid synthesis going.
- 14. Measure to assess the quality of the identification of nucleobases generated by automated DNA sequencing. (3 words)

Vertical:

- 1. Dideoxynucleosidetriphosphates (abbrev.)
- 2. Process of determining positions of reads on the reference genome.

- 4. The process of making many copies of a piece of DNA.
- 5. Found in pairs in DNA.
- 6. Chemical group attached to nucleotides to monitor incorporation into DNA.
- 9. Breakthrough sequencing method (abbrev.)
- 10. File format used to store sequence information.

### Solution



Solution:

B I O L O G Y I S G R E A T

## Exercise 3

**Question 3A** You want to determine how many reads  $N$  are needed to achieve a coverage depth  $C$  of 20X when sequencing reads for *Escherichia coli*.

The length of the reads  $L$  is 30nt and the *E. coli* genome  $G$  is approximately 4.6 million bases long.

**Formula**

$$N = \frac{C \times G}{L}$$

**Formula**

$$N = \frac{20 \times 4600000}{30} \approx 3066667 \text{ reads}$$