# 04\_05 QubitFluorometer

#### **DIENSTAG. 13.7.2021**

### **Goal-Setting**

- Measure the concentration of DNA in a sample
- The fluorescence molecules can intercalade into the DNA and the Qubit 2.0 fluorometer measures the intensity

### **Terms / abbreviations**

None

#### Risk areas

None

## Required materials and / or information

- Medium-sized tubes, Eppendorf
- Qubit dsDNA BR assay kit, ThermoFisher
  - o Qubit dsDNA BR Reagent
  - o Qubit dsDNA BR buffer
  - Qubit dsDNA BR Standards #1 and #2
- Solutions in the 4 °C fridge in Extension Lab, in a plastic bag

## Templates, devices, software

• Qubit 2.0 fluorometer, ThermoFisher

## **Preliminary work**

None

### **Operation**

## **Preparing samples for measuring**

- Prepare working solution:
  - $\circ~$  The Qubit dsDNA BR Reagent is diluted with Qubit ds DNA BR buffer 1:200 because 200  $\mu L$  for one sample is needed
  - o 199 µL from Qubit dsDNA BR buffer
  - o 1 µL Qubit dsDNA BR Reagent
- Prepare two standards with Qubit dsDNA BR Standards #1 and #2 (one negative (-) and one positive (+)):
  - Use the medium-sized tubes
  - 190 μL working solution
  - 10 μL standard
- Prepare sample:
  - Use the medium-sized tubes
  - o 199 µL working solution
  - o 1 µL sample

#### Measurement

- Choose ssDNA or dsDNA
- Measure in new standards
  - o Put Qubit dsDNA BR Standard #1 (-) in the opening, close the lid and start reading
  - o Put Qubit dsDNA BR Standard #tandard 2 (+) in the opening, close the lid and start reading

- Measure the sample
  - o Put the sample in the opening, close the lid and start reading
- Adjust the settings, because only 1  $\mu L$  of the sample is used and not 10  $\mu L$  like for the standards
  - o Press the button
  - o Choose 1 µL
  - $\circ$  Adjust the units --> ng/ $\mu$ L

# **Troubleshooting**

• For each new measurement new standards should be prepared because the fluorescence solution is not that stable

# Follow-up work

None