# **Prepare Gibson reaction**

# 1) pMel-10 + pGLN1

	pMel-10 PCR	gRNA pGLN1	Mastermix	Sterile MQ water
Conc fragment (ng/μL)	31,9	2,5	Х	X
Volume (μL)	3	4	10	3

# 2) Positive control

Prepare positive control: 10  $\mu$ L of control + 10  $\mu$ L of Gibson mastermix

## **Gibson reaction**

2 hours 50 degrees

# Transform to E. coli (5 tubes of E. coli)

- Transforms 2 µL of Gibson assembly to E. coli: construct and positive control.
- Also transform 2 μL of 4X diluted Gibson assembly: construct and positive control (4X dilution: mix 5 μL of Gibson assembly mix with 15 μL of sterile MQ water).
- Do a negative control: Transform 10 ng (100/ 20) of cut plasmid to *E. coli* (dilute beforehand -> transform with 2  $\mu$ L)
- Plate on LB + amp (construct