05_03_Seamless-cloning-reaction

DIENSTAG, 1.6.2021

Goal-Setting

• Cloning of an insert and transformation into competent cells

Terms / abbreviations

- cPCR = Colony polymerase chain reaction
- RT = Room temperature

Risk areas

None

Required materials and / or information

- Chemicals:
 - o Chemical competent cells (E. coli C3040I)
 - o Desired insert
 - o pMA_217 vector
- Materials:
 - 1.5 mL tubes, Eppendorf
 - Agar plates (Ampicillin 100 μg/mL)
 - GeneArt[™] Seamless Cloning and Assembly Enzyme Mix, Thermo Fisher Scientific
 - Ice
 - o Pipettes, Eppendorf
 - o S.O.C. Medium, Thermo Fisher Scientific or self-made
 - o Timer

Templates, devices, software

• ThermoMixer C, Eppendorf

Preliminary work

- 05_01_Preparation-of-S.O.C.-medium
- 06_03_Casting-agar-plates

Operation

- 1. Pipette 5 µL enzyme mix into a cup (on ice)
- 2. The insert/vector ratio is 4:1. The amount of the insert is 200 ng. In addition, 50 ng of the vector is pipetted to the enzyme mix. The volume should not exceed $5 \mu L$.
- 3. Incubate for 20 min at RT and incubate for 5 min on ice afterwards
- 4. Add 50 µL of chemical competent cells (E. coli C3040I)
- 5. Incubate on ice for 20 min
- 6. Heat shock at 42 °C for 45 s and incubate on ice for 3 min afterwards
- 7. Add 300 µL S.O.C. medium
- 8. Incubate for 30 min at 37 °C and 300 rpm
- 9. Plating out the medium on an agar plate and incubate them for 12-16 h at 37 °C

Troubleshooting

• Expected heights of the bands after cPCR for pMA_217 vector:

Uncut vector: 763 bpRe-ligated vector: 190 bp

o Vector with control insert: 732 bp

○ Vector with AT-rich insert: 190 + ~insert length bp

Follow-up work

None