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## ONA Cloning (Gibson Assembly, Transformation, Plating and Incubation)

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#### **ABSTRACT**

2023 NUS-Singapore iGEM Team followed this protocol to construct the plasmid of interest or to clone the plasmid of interest.

#### **MATERIALS**

- 1. NEBuilder HiFi DNA Assembly Master Mix
- 2. Competent cells made by 10β E. coli.
- 3. LB media.

#### DOI:

dx.doi.org/10.17504/protocol s.io.8epv5xnddg1b/v1

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**Protocol status:** Working We use this protocol and it's working

Created: Sep 27, 2023

#### SAFETY WARNINGS



- Proper laboratory PPE must be worn at all times.
- Thermal gloves shall be worn when handling competent cells from the -80°C fridge.
- Since cells like 10β E. coli are used in this protocol, a Biosafety Cabinet (BSC) is required to ensure safety.

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**PROTOCOL** integer ID:

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**Keywords:** DNA Cloning, Gibson Assembly, Assembly, Plasmid Assembly, Transformation, Plating, E coli.

Gibson	<b>Assem</b>	bly
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- Calculate the volumes of respective DNA fragments to assemble based on their length and concentration. (The maximum final volume of the mixed fragments is  $\Xi$  5  $\mu$ L for each reaction.)
- 2 Add the DNA fragments into a PCR tube according to the volumes obtained from the calculation.
- 3 Add  $\ \underline{\ \ }$  5  $\mu L$  of NEBuilder HiFi DNA Assembly Master Mix into the same PCR tube.
- 4 Vortex to mix the solution and centrifuge the PCR tube to spin down the remaining solution on the wall.
- 5 Incubate the PCR tube at 50°C for 01:00:00

#### 11

### **Transformation**

6 Switch on the water bath and set the temperature to 42°C.

- 7 Prepare a box of ice.
- Take an Eppendorf tube that contains Sample from the -80°C fridge.
- 9 Immediately place the Eppendorf tube with competent cells into the ice box for 00:05:00
- 11 Tap the bottom of the Eppendorf tube to mix the solution.
- Leave the Eppendorf tube in ice for 00:10:00
- 13 Place the Eppendoft tube into a foam floating.
- Place them into the water bath for 00:00:45 at 42°C for heat shock.

10m

- 15 Place the Eppendorf tube into the ice immediately.
- Add I 1 mL of the LB media into the Eppendorf tube.
- Place the Eppendoft tube into the incubator at 37°C for 01:00:00 for recovery.
- 18 Centrifuge the Eppendorf tube to form a cell pellet (no specific speed and time).

# **Plating and Incubation**

1h

- 19 Prepare an LB agar plate with the correct antibiotics.
- Remove 950  $\mu$ L of the LB solution from the Eppendorf tube that contains the cell pellet, leaving about 100  $\mu$ L in the Eppendorf tube.
- 21 Resuspend the cells by pipetting the solution.
- 22 Spread the cells onto the agar with the L-spreader.

1h

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