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# HiFi Gibson Assembly (Protocol for the NEBuilder® HiFi DNA Assembly Master Mix)

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1 Works for me dx.doi.org/10.17504/protocols.io.7kchksw



#### ABSTRACT

This is the protocol for DNA Assembly using the NEBuilder® HiFi DNA Assembly Master Mix.

#### **GUIDELINES**

### **Optimal Quantities**

NEB recommends a total of 0.03–0.2 pmols of DNA fragments when 1 or 2 fragments are being assembled into a vector, and 0.2–0.5 pmols of DNA fragments when 4–6 fragments are being assembled. Efficiency of assembly decreases as the number or length of fragments increases. To calculate the number of pmols of each fragment for optimal assembly, based on fragment length and weight, we recommend the following formula, or using the tool, <u>NEBiocalculator</u>.

pmols = (weight in ng) x 1,000 / (base pairs x 650 daltons) 50 ng of 5000 bp dsDNA is about 0.015 pmols 50 ng of 500 bp dsDNA is about 0.15 pmols

The mass of each fragment can be measured using the NanoDrop instrument, absorbance at 260 nm or estimated from agarose gel electrophoresis followed by ethicium bromide staining.

MATERIALS

NAME V CATALOG # V VENDOR V

NEBuilder HiFi DNA Assembly Master Mix - 10 rxns E2621S New England Biolabs

1 Set up the following reaction on ice (to 20µl total volume):

	2-3 Fragment Assembly	4-6 Fragment Assembly
DNA Ratio	Vector:Insert = 1:2	Vector:Insert = 1:1
Total amount of Fragments	0.03-0.2 pmols	0.2-0.5 pmols
NEBuilder HiFi DNA Assembly Master	10 μL	10 μL
Mix		
Deionized H2O	10 - X μL	10 - Χ μL
Total Volume	20 μL	20 μL

- 2 Incubate samples in a thermocycler at 50°C for 15 minutes (when 2 or 3 fragments are being assembled) or 60 minutes (when 4–6 fragments are being assembled).
- 3 Following incubation, store samples on ice or at -20°C for subsequent transformation.

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Transform into chemically competent cells (1-5 $\mu$ L) or in electrocompetent cells (1  $\mu$ L, diluted 10x or 5  $\mu$ L, after purification to remove salts).

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