



## **Target Sequence Cloning Protocol**

(Standard de-salted oligos are sufficient)

## <u>PX330-based plasmids, including PX458-462 – SpCas9 (or SpCas9n D10A nickase) + single guide</u> RNA:

To clone the guide sequence into the sgRNA scaffold, synthesize two oligos of the form:

5' - CACCGNNNNNNNNNNNNNNNNN - 3' 3' - CNNNNNNNNNNNNNNNNNNNNNNNCAAA - 5'

## PX260 and PX334 - SpCas9 (or SpCas9n D10A nickase) + CRISPR array + tracrRNA:

To clone the guide sequence into the sgRNA scaffold, synthesize two oligos of the form:

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## Oligo annealing and cloning into backbone vectors:

1. Digest 1ug of plasmid with *Bbsl* for 30 min at 37°C:

1 ug	Plasmid
1 ul	FastDigest Bbsl (Fermentas)
1 ul	FastAP (Fermentas)
2 ul	10X FastDigest Buffer
X ul	ddH <sub>2</sub> O
20 ul	total

- 2. Gel purify digested plasmid using QIAquick Gel Extraction Kit and elute in EB.
- 3. Phosphorylate and anneal each pair of oligos:

1 ul	oligo 1 (100uM)
1 ul	oligo 2 (100uM)
1 ul	10X T4 Ligation Buffer (NEB)
6.5 ul	ddH <sub>2</sub> O
0.5 ul	T4 PNK (NEB)
10 ul	total

Anneal in a thermocycler using the following parameters:

37°C	30 min
95°C	5 min and then ramp down to
	25°C at 5°C/min

4. Set up ligation reaction and incubate at room temperature for 10 min:

X ul	Bbsl digested plasmid
	from <b>step 2</b> (50ng)
1 ul	phosphorylated and annealed
	oligo duplex from <b>step 3</b> (1:200
	dilution)
5 ul	2X Quickligation Buffer (NEB)
X ul	ddH <sub>2</sub> O
10 ul	subtotal
<u>1 ul</u>	Quick Ligase (NEB)
11 ul	total

5. (optional) Treat ligation reaction with PlasmidSafe exonuclease to prevent unwanted recombination products:

11 ul	ligation reaction from <b>step 4</b>
1.5 ul	10X PlasmidSafe Buffer
1.5 ul	10mM ATP
<u>1 ul</u>	ddH <sub>2</sub> O
15 ul	total

Incubate reaction at 37C for 30 min.

6. Transformation