

Cloning gRNA expression vectors with pCFD1

Oligo design for pCFD1:

PCFD1-dU61gRNA U6:1 promoter Bbs1 spacer gRNA core gRNA core GCCCACTTGAAGCCCCAGAAGCTCTTCTGGACAAAATCTCG

Sense: 5' – TTC**G**-N19/20

Anti-sense: 5' – AAAC-N19/20 reverse complement

Note that the G at position 4 in the sense oligo is the first base that is transcribed. If your protospacer sequence starts with a G then N will be 19. If it does not start with a G enter all 20 nucleotides behind TTCG.

Resuspend oligos to 100uM.

Set up the following phosphorylation and annealing reaction:

1ul sense oligo (100uM)

1ul anti-sense oligo (100uM)

1ul 10X T4 Ligation Buffer (NEB)

6.5ul ddH2O

0.5ul T4 PNK (NEB)

Incubate and anneal in a thermocycler:

37°C 30min

95°C 5min

ramp down to 25°C at 5°C/min

Set up the following ligation reaction:

Xul Bbsl digested pCFD1 (use 50ng)

1ul annealed oligos diluted 1:200

1.5ul 10X T4 Ligation Buffer (NEB)

Xul ddH2O

1ul T4 DNA ligase

total volume 15ul

Ligate 30min at room temperature.

Transform into competent bacteria. Plate on Ampicillin plates.