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Construction of U6-based sgRNA expression vectors

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Genetics



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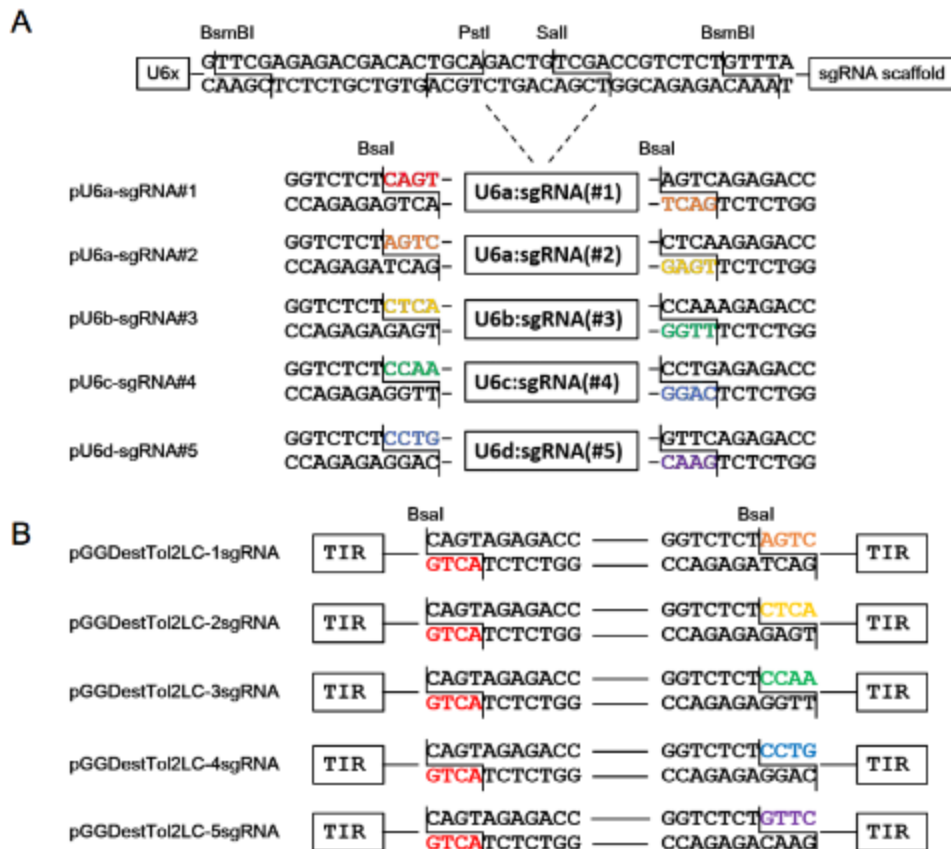
Abstract

This protocols is from:

Linlin Yin, et al. (2015) **Multiplex Conditional Mutagenesis Using Transgenic Expression of Cas9 and sgRNAs**. *Genetics* 200:431-441; doi:10.1534/genetics.115.176917

Please see the **full manuscript** for additional details.

Guidelines



List of plasmids deposited at Addgene (Deposit 71794)

Plasmid ID Plasmid Name

64237 pME-Cas9
64239 pGGDestTol2LC-1sgRNA
64240 pGGDestTol2LC-2sgRNA
64241 pGGDestTol2LC-3sgRNA
64242 pGGDestTol2LC-4sgRNA
64243 pGGDestTol2LC-5sgRNA
64245 pU6a:sgRNA#1
64246 pU6a:sgRNA#2
64247 pU6b:sgRNA#3
64248 pU6c:sgRNA#4
64249 pU6d:sgRNA#5
64250 pU6a:sgRNA(try)

Reference:

JAO, L. E., S. R. WENTE and W. CHEN, 2013 **Efficient multiplex biallelic zebrafish genome editing using a CRISPR nuclease system.** Proc Natl Acad Sci U S A 110: 13904- 13909.

Materials

STEP MATERIALS

 NEBuffer 2 - 5.0 ml **New England Biolabs Catalog #B7002S**

 NEBuffer 2 - 5.0 ml **New England Biolabs Catalog #B7002S**

Protocol materials

 NEBuffer 2 - 5.0 ml **New England Biolabs Catalog #B7002S**



- 1 Order a pair of U6 vector-specific, 23nt targeting primers for each target For each target (GN19), order a forward primer TTCGN19 for the U6 promoter vectors; order a reverse primer AAACN19 Note that the N19 in the reverse primers is reverse complementary to that in the forward primer

To order primers compatible with both the pU6x-sgRNA vectors and pT7-gRNA (Addgene plasmid #4675; (JAO et al 2013)), order degenerate primers WTMGGN18 (forward) and AMMCN18C (reverse), where M=A or C, W=A or T Again, the N18 in the reverse primers is reverse complementary to that in the forward primer

Annealing the primers

- 2 Mix 1µl 100µM stock each in a 20µl 1x NEB buffer 2

 NEBuffer 2 - 5.0 ml **New England Biolabs Catalog #B7002S**

- 3 Incubate the mixture as follows:

95 °C for 5 min,
ramp down to 50 °C at 0.1 °C/sec,
50 °C for 10 min
chill to 4 °C at normal ramp speed

Ligating the annealed oligos

- 4 Mix the following components to ligate the annealed oligos to the U6 vector of choice (pU6x-sgRNA #1-#5, see diagrams and plasmid list in the Guidelines)

Protocol

NAME

U6-based sgRNA ligation mixture

CREATED BY

Tracey Depellegrin

Preview

- 4.1 1 µl 10x CutSmart buffer

- 4.2 1 µl T4 DNA ligase buffer



4.3 0.25µl U6 plasmid (about 100ng)

4.4 1µl annealed oligos

4.5 0.3 µl T4 DNA ligase

4.6 0.3 µl BsmBI

4.7 0.2µl PstI (optional)

4.8 0.2 µl Sall (optional)

4.9 5µl H₂O

5 Incubate at 37 °C for 20 min

 00:20:00

6 Incubate at 16 °C for 15 min

 00:15:00

7 Incubate at 37 °C for 10 min

 00:10:00

8 Incubate at 55 °C for 15 min

 00:15:00

9 Incubate at 80 °C for 15 min (optional)

 00:15:00

10 The reaction is ready for transformation (use 2 µl of the ligation and plate 10% of the transformants). Transform and spread onto spectinomycin (50µg/ml) plates.

