

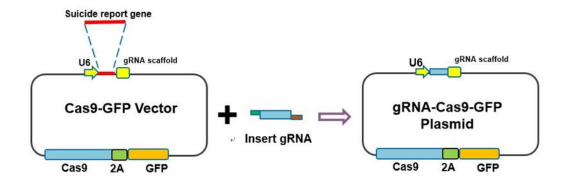
### High-Efficiency gRNA-Cas9-GFP Plasmid Assembly Kit

Cat. No.: L00692 Version 2018-07-16

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### I Description

The High-Efficiency gRNA-Cas9-GFP plasmid Assembly Kit is used for easier and highly efficient cloning of gRNA fragments into Cas9 vectors. This vector has been constructed by integrating a suicide reporter gene into the gRNA insert site. Under the selection of this suicide report gene, almost 100% of clones are positive clones without screening. The High-Efficiency gRNA-Cas9-GFP plasmid is provided in both the circular and linear formats. The GFP fluorescence reporter is linked by 2A, a self-cleavage peptide, and is expressed separately from Cas9, keeping the maximum activity of Cas9.



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### **II** Contents

Components	10-reaction	25-reaction	Concentration
Cas9-GFP Vector	20 μL	50 μL	50 ng/μL
Control gRNA insert	10 μL	25 µL	0.05 μM
Control gRNA primer F	10 μL	25 µL	10 μΜ
Control Screening primer R	50 μL	125 μL	10 μΜ

### **III Key Features**

- ♦ **Simplicity** –Only a pair of primers is needed. The digestion and ligation can be done in a single reaction.
- ♦ Efficiency –Up to 100% positive clones.
- ♦ Fidelity –No mutations.
- ♦ **Time-saving** The construction reaction can be completed within 30 minutes.

### **IV** Quality Control Analysis

- ♦ Positive clone rate: up to 100%.
- $\diamond$  None colonies could be grown in DH5 $\alpha$  competent cells without the insert.

# V Product Usage

Rapid construction of gRNA-Cas9 plasmid for the subsequent gene editing research.

### VI Storage

Please store at -20°C.



#### VII Protocol

#### 1 Design specific gRNA primers

Forward primer gRNA-F: CACC+N<sub>19-21</sub> (gRNA target sequences)
Reverse primer gRNA-R: AAAC+ N<sub>19-21</sub> (gRNA Reverse complementary sequence)
e.g: If the target sequence of gRNA is CATCATTGGAAGACGTTCTT,
the forward primer is CACCCATCATTGGAAGACGTTCTT,
and the reverse primer is AAACAAGAACGTCTTCCAATGATG.

#### 2 Anneal the primers

- Dissolve specific gRNA primers F and R to 10 μM.
- Dilute to 0.1 μM.
- ⋄ Mix the primers F and R 1:1 by volume.
- ♦ Denature at 95°C for 5 minutes.
- Place the mixture at room temperature for another 10 minutes, so that the primers cool down to room temperature gradually.

#### 3 Ligation reaction

	<b>Experimental reaction</b>	Positive control reaction
10×T4 DNA ligase buffer	1 μL	1 µL
Cas9-GFP Vector	1 μL	1 μL
annealed specific gRNA insert	0.5 μL	-
Control gRNA insert	-	0.5 μL
Bbs I(10 U/μL)	0.25µL	0.25μL
T4 DNA ligase(400U/μL)	0.25 μL	0.25 μL
$H_2O$	7 μL	7 μL
Total reaction volume	10 μL	10 μL

#### Reaction conditions:

Temperature	Time	Cycle
37°C	5 minutes	3x
16°C	5 minutes	37
4°C	Hold	-



#### 4 Transformation

- $\diamond~$  Add 5 µL of the above reaction mixture to 50 µL DH5 $\alpha$  competent cells.
- Place the cells on ice for 30 minutes.
- ♦ Heat shock at 42°C for 1 min followed by ice bath for 2 minutes.
- Add 500 μL LB Broth and shake for 30 min at 37°C under 180 rpm.
- Add 100 μL to LB Agar plate with 100 μg/ml Ampicillin and incubate overnight at 37°C.

### 5 Colony PCR screening

	Experimental reaction	Positive control reaction
10×Taq buffer	2.5 µL	0.25 μL
10mM dNTP	0.5 μL	0.5 μL
Specific gRNA-F (10μM)	0.5 μL	-
Control gRNA primer F (10µM)	-	0.5 µL
Control Screening primer R (10µM)	0.5 μL	0.5 µL
Taq DNA polymerase (5U/µL)	0.25 μL	0.25 μL
Colony	a little	a little
H <sub>2</sub> O	20.75 μL	20.75 μL
Total reaction volume	25 μL	25 μL

Note: Less than 10 clones is enough for screening in each experimental plate.

#### Reaction conditions:

Temperature	Time	Cycle
94°C	5 minutes	1
94°C	10 seconds	
55°C	10 seconds	25×
72°C	10 seconds	
72°C	5 minutes	1
4°C	Hold	-

PCR products are detected using agarose gel electrophoresis. The positive control band runs at 270bp. The negative control should remain blank.



Note:This is a basic protocol. The reagent concentrations, conditions, and parameters may need to be optimized.

#### VIII FAQ

- Which kinds of competent cells can be used for transformation?
   Answer: Suitable competent cells include, but are not limited to DH5α, Stbl2, Stbl3, EPI300, EPI400, JM108, ER2925 and DH10B-N. Inappropriate competent cells include, but are not limited to Top10, XL1-Blue, and SURE2.
- 2. What is the size of the plasmid before and after transformation? Answer: The size of Cas9-GFP vector provided in the kit is 9629bp, and the size of the target plasmid after transformation is about 9290-9292 bp depending on the inserted gRNA.
- Are there any other versions of gRNA-Cas9 constructs?
   Answer: GenScript carries a series of different versions of gRNA-Cas9 constructs, including "High-Efficiency gRNA-Cas9-Puro Plasmid (linear) Assembly Kit (Cat. No. L00691)", "High-Efficiency gRNA-Cas9-GFP Plasmid Assembly Kit (Cat. No. L00692)", and "High-Efficiency gRNA-Cas9-Puro Plasmid Assembly Kit (Cat. No. L00693)".

## IX Ordering Information

Product Name	Cat. No.
GenCrispr Cas9 Nuclease	Z03386
GenCrispr Mutation Detection Kit	L00688
GenCrispr Cas9 Antibody, pAb, Rabbit	A01885

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