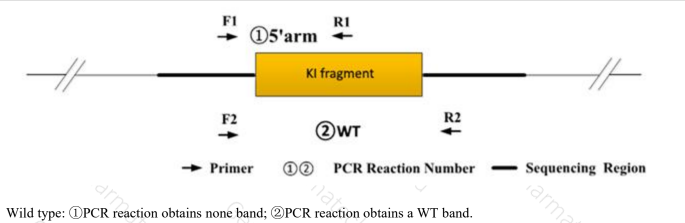


	m	Genotyp	oing Report		~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
Strain ID	T055748	Strain Type	KI(Cas9)	Genetic Background	C57BL/6JGpt
Designer	Tianjiao Wang	Gene Name		Rosa26-CAG-EGFP-polyA	°C.

1. Strategy of Genotyping

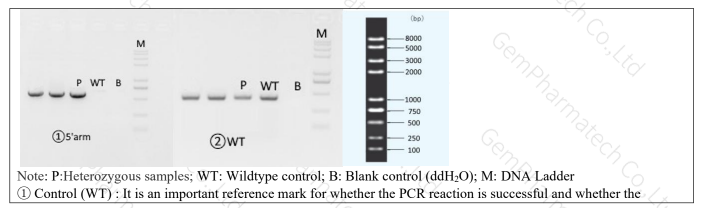


Wild type: ①PCR reaction obtains none band; ②PCR reaction obtains a WT band. Heterozygote: ①PCR reaction obtains a Targeted band; ②PCR reaction obtains a WT band. Homozygote: ①PCR reaction obtains a Targeted band; ②PCR reaction obtains none band. Note: The sizes of WT and Targeted band are shown below. For ②PCR reaction, because the WT band is much smaller than the target band, it is likely to produce dominant amplification, the reaction is only used to judge whether there is a WT allele.

2. Primer Information

PCR No.	Primer No.	Sequence	Band Size	
1)5'arm	Rosa26-S4-tF2	GTGGGATACAGAAGACCAATGCAG	WT:0bp	
	CAG-tR1	TGGCGTTACTATGGGAACATACGTC	Targeted:555bp	
2WT	Rosa26-S4-tF1	AATGTAGGGCCAGAGTTTAGCCAG	WT:460bp Targeted:0bp	
	Rosa26-S4-tR1	TGAAAGATTTCCCAACCCCAC	Targeted.00p	

3. Gel Image & Conclusion





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product band position and size meet the theoretical requirements.

2 Control (B) : PCR amplification was performed without template in the PCR reagent to monitor whether the reagent was contaminated.

4. PCR Condition

PCR Reaction Component

PCR Reaction C	omponent	$\gamma_{\mathcal{O}_{\mathcal{I}}} = \gamma_{\mathcal{O}_{\mathcal{I}}}$	·	· · · / ·	
Seg.		reaction component			
12	2 × Rapid Taq Master	2 × Rapid Taq Master Mix (Vazyme P222)			
2 75	ddH2O		9.5	50	
3 20	Primer A(10pmol/µl)	ns C	7,1	~~~ `	
4 ?	Primer B(10pmol/µl)		1	N C	
5	Template(20~80ng/µl	Template(20~80ng/µl)			
PCR program	priority selection	S. S.		2 C	
Seg.	Temp.	Time	Cycle		
1	95℃	5min			
2	98°C	30s	20×	20×	
3 6	65℃*(-0.5℃/cycle)) 30s	6		
4 ¹ N ₂	72°C	45s*			
5	98°C	30s	15×	-0,	
6	55°C*	30s	(?)		
7 %	72°C	45s*	G _e		
8	72°C	5min	70,	6	
9 9	10°C	hold	Pr.		
PCR program I	I the second choice	and the	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	< ^ ^	
Seg.	Temp.	Time	Cycle		
1	95°C	5min	12 A	6	
2 7	98°C	30s	35×		
3	58°C*	30s			
4	72°C	45s*			
5 %	72°C	5min	- Ce		
6	10°C	hold -	/.	\sim	

Note*: Annealing temperature and extension time can be determined according to the actual amplification situation and amplification enzyme efficiency.