

Strain ID	T025270	Strain Type	CKO(Cas9)	Genetic Background	C57BL/6JC
Designer	Ya'nan Xu	Gene Name		Kel	20
	F1 → ① ⁴	5'arm 🗲	F2	23'arm	22
	- (1):	sarm -	+	@5 arm	-

Wild type: ①PCR reaction obtains a single WT band; ②PCR reaction obtains a single WT band. Heterozygote: ①PCR reaction obtains a WT band and a Targeted band; ②PCR reaction obtains a WT band and a Targeted band.

(1) (2) PCR Reaction Number - Sequencing Region

Homozygote: ①PCR reaction obtains a single Targeted band; ②PCR reaction obtains a single Targeted band. Note: The sizes of WT and Targeted band are shown below.

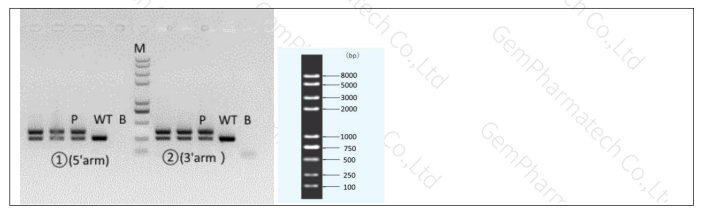
2. Primer Information

Loxp

Primer

PCR No.	Primer No.	Sequence	Band Size	
1)(5'arm)	T025270(P1)-F1	CACCTGAGTCTGGAACTCTAGTCTATC	WT: 264bp Targeted: 369bp	
	T025270(P1)-R1	TAAGAGAAGAGGCATTTTGGAGACAG		
@(3'arm)	T025270(P1)-F2	GGATATAAGGTTGAGATAGATGAGGTG T	WT: 269bp Targeted: 375bp	
	T025270(P1)-R2	CTCCATCATCTCTATCTCTATCTCTA		

3. Gel Image & Conclusion





Note: P: Heterozygous samples; WT: Wildtype control; B: Blank control (ddH₂O); M: DNA Ladder ① Control (WT) : It is an important reference mark for whether the PCR reaction is successful and whether the 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

4. PCR Condi	tion		Party istra	
PCR Reaction Co	mponent	3. 0	24	
Seg.	reaction co	reaction component		
ı M	2 × Rapid Taq Master Mix (Vazym	ie P222)	12.5	
<u>2</u> 7.	ddH2O	0	9.5	
3	Primer A(10pmol/µl)	De la Carlo	1 7	
ļ	Primer B(10pmol/µl)	Primer B(10pmol/µl)		
; 6,	Template(20~80ng/μl)	$^{\prime}$ C (
CR program I	priority selection	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Mr. Co	
Seg.	Temp.	Time	Cycle	
L C	95°C	5min	The second	
	98°C	30s	20× 20×	
s Tr	65℃*(-0.5℃/cycle)	30s	K. G.	
1 ⁽⁴⁾	72℃	45s*		
G C	98℃	30s	15×	
; ⁵ 75.	55℃*	30s	3. [°] 30	
	72℃	45s*		
3	72℃	5min	3. 4	
jC _C	10°C	hold		
CR program II	the second choice			
Seg.	Temp.	Time	Cycle	
r ?	95°C	5min	1737 × 14	
2	98°C	30s	35×	
6	58°C*	30s	G C	
1 ⁷ 0	72°C	45s*		
;	∂72℃	5min	The second se	
;	10°C	hold	- Bar	

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



