

Strain ID	T051940	Strain Type	CKO(Cas9)	Genetic Background	C57BL/6JGp
Designer	Ya'nan Xu	Gene Name	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Mapk13	$^{\circ}$
Strategy of (Genotyping	1 7 C	m.	C That	
	F1 → ①5'a	rm ▲	F2	23'arm	22
//		->-	Flox	- >	_//_

Heterozygote: ①PCR reaction obtains a WT band and a Targeted band; ②PCR reaction obtains a WT band and a Targeted band;

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.

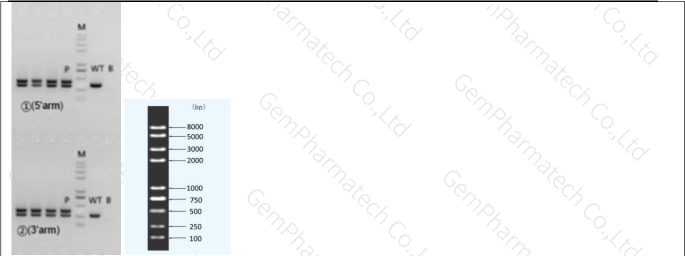
2.	Primer	Information

PCR No.	Primer No.	Primer Name	Sequence	Band Size
(1)(5'arm)	F1	T051940(P2)-F1	GTTCTGTGAGGACTCCAGTCCGA	WT: 309bp
①(5'arm)	R1 T051940(P2)-R1 AAATGGACCCTAGCCCTTCTG		AAATGGACCCTAGCCCTTCTGCCT	Targeted: 414bp
	F2	T051940(P2)-F2	GGGTTTGGCTGTTACAAAGTGTCA	WT: 270bp
2(3'arm)	R2	T051940(P2)-R2	CAGAGGCAGGAGGATCAGAAGTTC	Targeted: 376bp

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

(Generally recommend to use Vazyme P222; If the sequences contain special structures such as $GC\% \ge 60\%$ or $GC\% \le 40\%$, recommend to use Vazyme P515.)

PCR Reaction Component

Seg.		reaction component	Volume (μl)		
1 Cons,	193 a	2 × Rapid Taq Master Mix(Vazyme P222) or 2 × Phanta Max Master Mix (Vazyme P515)			
2 7		ddH2O	9.5		
3		Primer A(10pmol/µl)			
400		Primer B(10pmol/µl)			
5 202	1°C (Template(20~80ng/µl)			
PCR program I	priority selection	$\gamma_{\mathcal{S}}$	An isla		
Seg.	Temp.	Time	Cycle		
1	95°C	5min			
2 6	98°C	30s	20×		
3 70	65℃*(-0.5℃/cycle)	30s	γ_{λ}		
4	72℃	45s*			
5	98°C	30s	15×		
6 0	55°C*	30s	Con The		
7 2	72°C	45s*	$\overline{\gamma}_{2}$		
8	72℃ ·	5min	20. 36		



9 ~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	10°C	a la com	hold	1	
PCR program	II the second choice	nax.	$\langle \varphi \rangle$		912 3/x
Seg.	Temp.		Time		Cycle
1 000	95℃	6	5min	Con .	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
2 75	98°C	na.	30s	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	35×
3	58°C*	n n n n n n n n n n n n n n n n n n n	30s		
4	72°C		45s*	S	
5 702	72°C	C _C	5min	n,	20
6 Pr	10°C		hold	2	
	2	Q.	· · · · · · · · · · · · · · · · · · ·		D. C

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