

		Genotyp	ing Report	and -	
Strain ID	T040689	Strain Type	CKO(Cas9)	Genetic Background	C57BL/6JGp
Designer	Ya'nan Xu	Gene Name	34.2	Klhl11	°C –
Strategy of C	Genotyping	<u>^</u>	mar and a second	C nate	
Strategy of C	F1	R1	F1	A R	· ·
Strategy of C	F1 → ①5'a		F2	@3'arm 🖁	2
Strategy of C	F1		F2 Flox	@3'arm ▲	<u>2</u>
Strategy of C	F1		 →	@3'arm 🚆	2 <u>-</u>
Strategy of C	F1 → ①5'a		Flox	②3'arm R → → → → → → → → → → → → → → → → → → →	2 <u>-</u> //

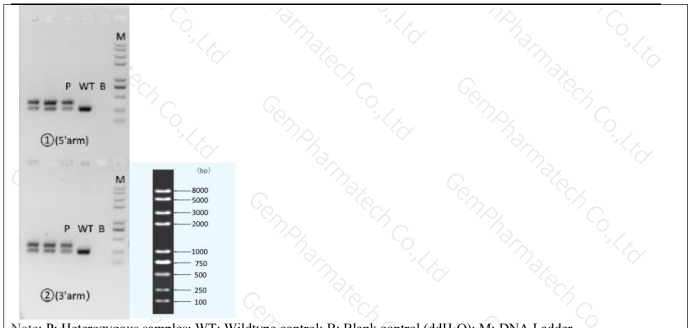
Targeted band, Homozygote: ①PCR reaction obtains a wireband and a rangeted band; ②PCR reaction obtains a single Targeted band. Note: The sizes of WT and Targeted band are shown below.

2. Primer Information

PCR No.	Primer No.	Sequence	Band Size
()(5 1)	T040689-F1	ACCCTGTCTCGAAAACAAACAAAC	WT: 264bp
()(5°arm)	(1)(5'arm) T040689-R1 TTTACTGTA	TTTACTGTAGAAGGGCAGAGGTGGC	Targeted: 369bp
2(3'arm)	T040689-F2	ACAAGTAAAGCTTTCAAATGGCAAGG	WT: 258bp
	TAAAAACCTTTTCTGCAAGATGACATTG	Targeted: 364bp	

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

PCR Reaction Con	nponent		No. St
Seg.	reaction comp	Volume (µl)	
1 7	2 × Rapid Taq Master Mix (Vazyme	P222)	12.5
2	ddH2O		9.5
3	Primer A(10pmol/µl)		13 CA
4	Primer B(10pmol/µl)	1 22	
5 70,	Template(20~80ng/µl)		1 2
PCR program I p	priority selection	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	
Seg.	Temp.	Time	Cycle
1	95°C	5min	A K
2	98°C	30s	20×
3	65℃*(-0.5℃/cycle)	30s	S. S.
4 ⁷ .	72°C	45s*	
5	98°C	30s	15×
6	55°C*	30s	
7 70/	72°C	45s*	
8 2	72°C	5min	1300



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9	10°C	hold	7.S. (C.
PCR progra	im $^{\mathrm{II}}$ the second cho	pice	an ilx
Seg.	Temp.	Time	Cycle
1 22	95°C	5min	Centra Contra
2 7	98°C	30s	35×
3	58°C*	30s	The star
4	72°C	45s*	Co Co
5 70	72°C	5min	γ_{S}
6 🖓	10°C	hold	

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