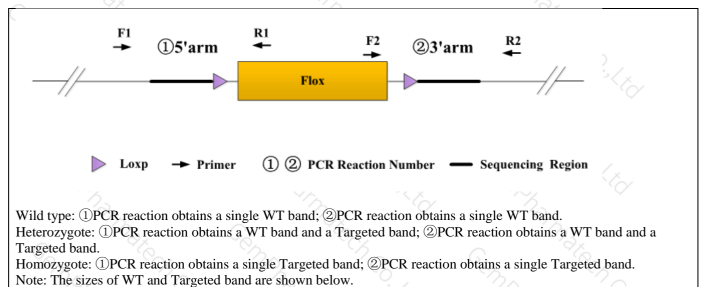


		Genotyp	oing Report		Co.
Strain ID	T053623	Strain Type	CKO(Cas9)	Genetic Background	C57BL/6JGpt
Designer	Tianjiao Wang	Gene Name	· · · · ·	Epor	C

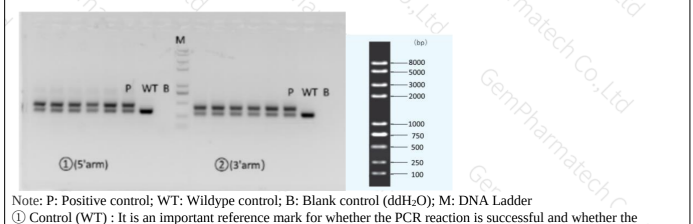
## 1. Strategy of Genotyping



### 2. Primer Information

PCR No. Primer No.		Sequence	Band Size	
	T053623(P3)-F1	CTGTCCATGGGTGTGTCAATGC	WT: 309bp	
(1)(5'arm)	T053623(P3)-R1	ACCAGCCCCAGACCTGTGCTTT	Targeted: 414bp	
2(3'arm)	T053623(P1)-F2	GGTCAAAGGACAACTTTTGTGAGC	WT: 281bp Targeted: 387bp	
	T053623(P1)-R2	TTGGCTCAAAGCCAATCAGATAG		

#### 3. Gel Image & Conclusion



product band position and size meet the theoretical requirements.

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(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 Compo	onent	C C	°°
Seg.	reaction comp	Volume (µl)	
1 3/2	2 × Rapid Taq Master Mix (Vazyme P2	22)	12.5
2 2	ddH2O	9.5	
3	Primer A(10pmol/µl)		1 3
4	Primer B(10pmol/µl)	and the second	1
5 3	Template(≈100ng/µl)	0	1
PCR program $①$ pric	ority selection		RX 4
Seg.	Temp.	Time	Cycle
1 6	95°C	5min	
2	98°C	30s	20×
3 977	65°C* (-0.5°C/cycle)	30s	12m
4	72℃	45s*	
5 3	98°C	30s	20×
6	55℃*	30s	K. C.
7 7	72°C	45s*	Str. St
8 6	72℃	5min	
9 70.	10°C	hold	
PCR program $2$ the	e second choice		5. 0.
Seg.	Temp.	Time	Cycle
	95°C	5min	ALC CA
2	98°C	30s	35×
3 77	58℃*	30s	the star
4 77	72℃	45s*	134
5	72℃	5min	Ϋ́Υ <sub>Ω</sub>
5 6	10°C	hold	G

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