

		Genotypi	ing Report		- C K.K.
Strain ID	T024657	Strain Type	CKO(Cas9)	Genetic Background	C57BL/6JGpt
Designer	Ya'nan Xu	Gene Name		Slc17a7	°C
Strategy of	Genotyping	in ha	3	armare.	
	F1 → ①5'arm	R1	F2 ②3	'arm 🐥	
_//_		≻— Flox		//	"" < x 

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. 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.

Primer

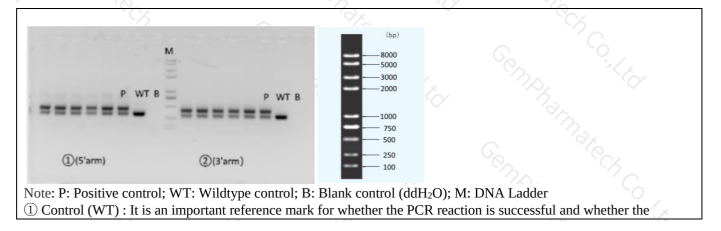
(1) (2) PCR Reaction Number - Sequencing Region

## 2. Primer Information

Loxp

PCR No.	Primer No.	Sequence	Band Size	
(1)(5'arm)	T024657-F1	GGGAATGAATGCGCCAATCT	WT: 306bp Targeted:441bp	
	T024657-R1	TTCCGAGCCATCTCGGCTATA		
@(3'arm)	T024657-F2 GCGGGTACTAGGTGGGTGATTT		WT: 281bp	
	T024657-R2	CAGAATGGGCCACACGTTTCT	- Targeted:387bp	

## 3. Gel Image & Conclusion





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product band position and size meet the theoretical requirements.

<sup>(2)</sup> 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		M. M		
Seg.	reaction co	reaction component			
175,	2 × Rapid Taq Master Mix (Vazym	2 × Rapid Taq Master Mix (Vazyme P222)			
2 3	ddH2O		9.5		
3	Primer A(10pmol/µl)	. ··· / x	1		
4	Primer B(10pmol/μl)	Primer B(10pmol/µl)			
5	Template(≈100ng/µl)	Template(≈100ng/µl)			
PCR program	① priority selection	C C	· · · · · · · · · · · · · · · · · · ·		
Seg.	Temp.	Time	Cycle		
1	95°C	5min	ann-		
2 6	98°C	30s	20×		
3 <sup>7</sup> 0,	65°C*(-0.5°C/cycle)	30s	24. 3		
4	72℃	45s*			
5	98°C	30s	20×		
6	55°C*	30s			
7 2	72°C	45s*			
8	72°C	5min	Dr. Sh		
9	10°C	hold	1 Max 1 4		
PCR program	② the second choice	Tak Con			
Seg.	Temp.	Time	Cycle		
1	95°C	5min			
2	98°C	30s	35×		
3	58℃*	30s	3		
4	72°C	45s*			
5	72℃	5min	$\gamma_{\mathcal{A}_{\mathcal{L}}}$		
6	10°C	hold	ng p.		

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