## GemPharmatech Co.,Ltd

Strain ID	T012901	Strain Type	CKO(Cas9)	Genetic Back	around	C57BL/6JGr
		G	CKO(Cas)			CJ/BE/030
Designer	Tiantian Sun	Gene Name		SCN4B		6
				(Q)		
Strategy of	Genotyping				12	
	F1	R1	F2	@ <b>2!</b>	R2	
_//_	<sup>F1</sup> → ①5'ar		F2 Flox	@3'arm	R2 ◀─	_//
_//_			+	@3'arm	R2	-//

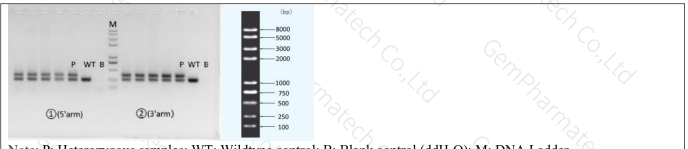
Heterozygote: ①PCR reaction obtains a WT band and a Targeted band; ②PCR reaction obtains a WT band and a Targeted band.

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	
	∕∕∕F1	T012901(P2)-F1	TCTGCATCCTGGTTCTGGGCTTA	WT: 243bp	
(1)(5'arm)	R1	T012901(P2)-R1	CTCCTTTCCCCAAGAGCCATTC	Targeted: 348bp	
@(3'arm)	F2	T012901(P2)-F2	CGAAAGAGACTCTAGAATGGAGGGA	WT: 252bp	
	R2	T012901(P2)-R2	TAGGAGGTCATGTCACCCTTTGG	Targeted: 358bp	

## 3. Gel Image & Conclusion



Note: P: Heterozygous samples; WT: Wildtype control; B: Blank control (ddH<sub>2</sub>O); 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	- 12 - 12 - 12 - 12 - 12 - 12 - 12 - 12	900	- 12	
Seg.	rea	reaction component			
1704		q Master Mix(Vazyme P222) or ax Master Mix (Vazyme P51	$\gamma_{\rm S}$ $\gamma_{\rm C}$	ò	
2	B. Ch	ddH2O	9.5	34	
3	Pri	imer A(10pmol/µl)		<i>\</i>	
4	Pri	imer B(10pmol/µl)	1 3		
5 %	Ten	nplate(20~80ng/µl)	G 1 O	_	
PCR program	I priority selection	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	no.		
Sog	Tomp	Time	Cycle		

Seg.	Temp.	Time	Cycle
1	95°C	5min	The second
2	98°C	30s	20×
3	65℃*(-0.5℃/cycle)	30s	K. C.
4 ろ、	72°C	45s*	an it
5 0	98°C	30s	15× 2
6	55℃*	30s	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
7	72°C	45s*	
8 72	72°C	5min	Mr. Mr
96.	10°C	hold	

PCR program II the second choice

Seg.		Temp.		Time		Cycle	
1	n <sub>ate</sub>	95°C	narry.	5min		nay.	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
2	C	98°C	6	30s		35× 🔗	
3	Co.	58°C*	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	30s	G.	1	
4	na,	72°C	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	45s*	~~/		
5	732	<b>72</b> ℃	- M.	5min		~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	
6		<b>10°</b> C	G. 97	hold		3	

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