

# Klhl34 Cas9-CKO Strategy

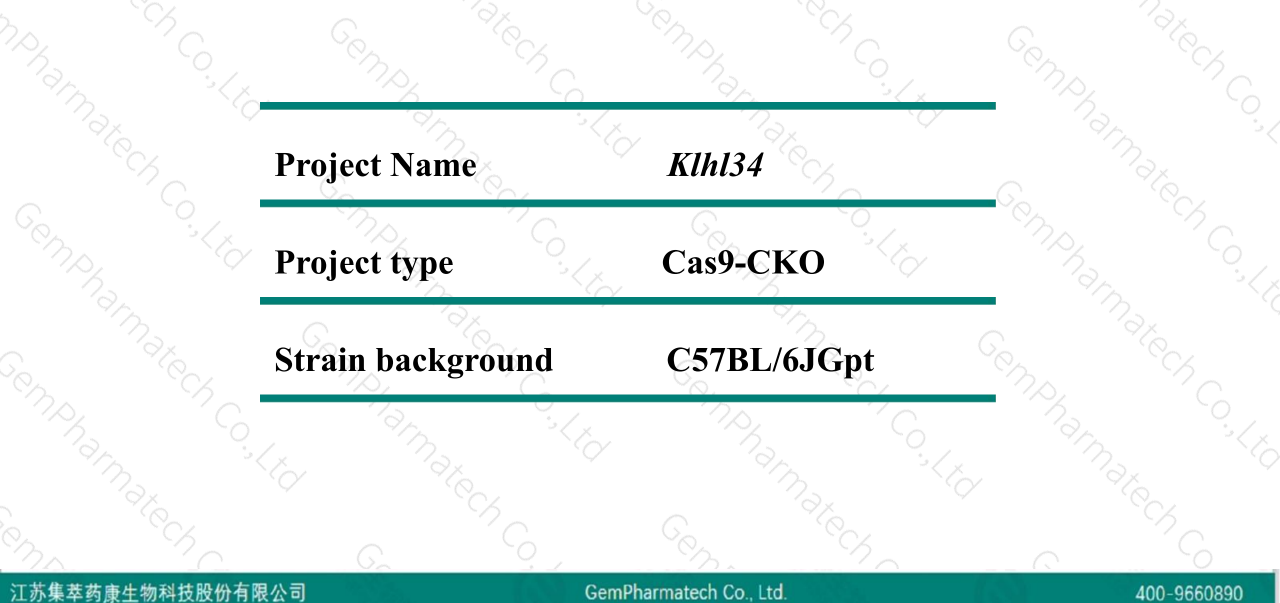
Designer: Daohua Xu

**Reviewer: Huimin Su** 

**Design Date: 2020-7-23** 

## **Project Overview**



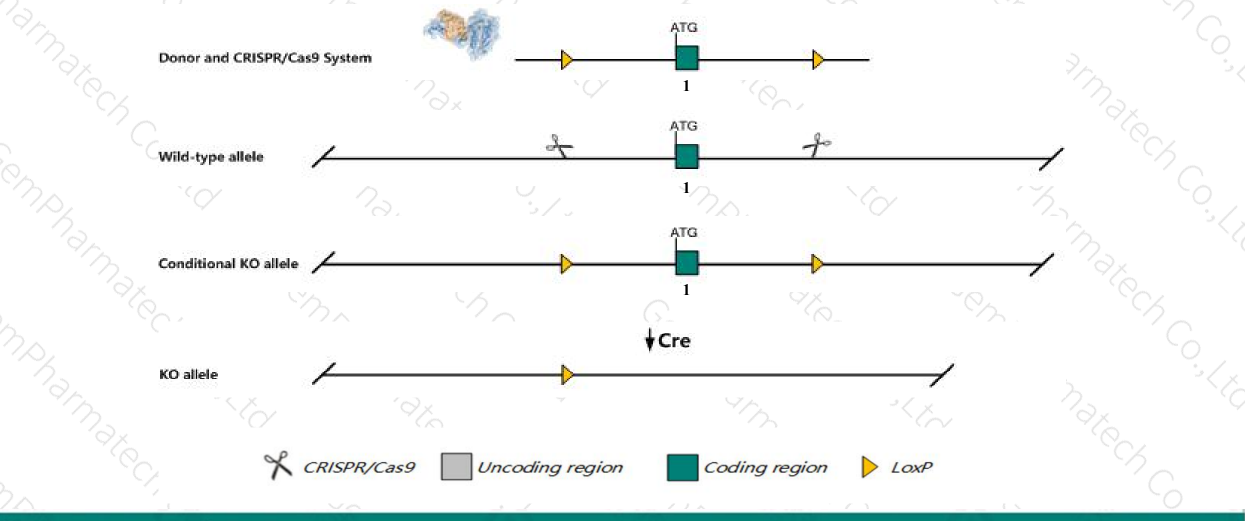


## **Conditional Knockout strategy**



400-9660890

This model will use CRISPR/Cas9 technology to edit the *Klhl34* gene. The schematic diagram is as follows:



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The Klhl34 gene has 1 transcript. According to the structure of Klhl34 gene, exon1 of Klhl34-201(ENSMUST00000087157.4) transcript is recommended as the knockout region. The region contains all of the coding sequence. Knock out the region will result in disruption of protein function.

➤ In this project we use CRISPR/Cas9 technology to modify *Klhl34* gene. The brief process is as follows:CRISPR/Cas9 system and Donor were microinjected into the fertilized eggs of C57BL/6JGpt mice.Fertilized eggs were transplanted to obtain positive F0 mice which were confirmed by PCR and sequencing. A stable F1 generation mouse model was obtained by mating positive F0 generation mice with C57BL/6JGpt mice.

> The flox mice will be knocked out after mating with mice expressing Cre recombinase, resulting in the loss of function of the target gene in specific tissues and cell types.



- > The *Klhl34* gene is located on the ChrX. If the knockout mice are crossed with other mice strains to obtain double gene positive homozygous mouse offspring, please avoid the two genes on the same chromosome.
- > This strategy is designed based on genetic information in existing databases.Due to the complexity of biological processes,all risk of loxp insertion on gene transcription, RNA splicing and protein translation cannot be predicted at existing technological level.

## **Gene information (NCBI)**



☆ ?

#### Klhl34 kelch-like 34 [Mus musculus (house mouse)]

Gene ID: 245683, updated on 13-Mar-2020

#### Summary

Official Symbol	Klhl34 provided by MGI
<b>Official Full Name</b>	kelch-like 34 provided byMGI
<b>Primary source</b>	MGI:MGI:2685234
See related	Ensembl:ENSMUSG0000047485
Gene type	protein coding
RefSeq status	VALIDATED
Organism	Mus musculus
Lineage	Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Euarchontoglires; Glires; Rodentia; Myomorpha; Muroidea; Muridae; Murinae; Mus; Mus
Also known as	Gm388
Orthologs	human all

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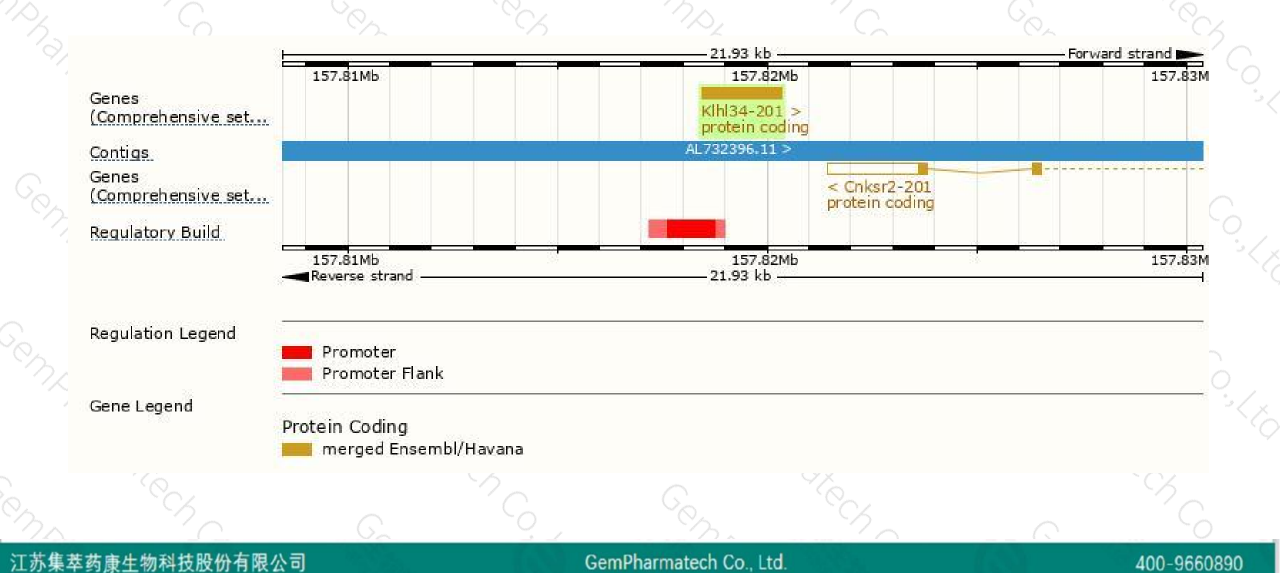


The gene has 1 transcript, and the transcript is shown below:

Name	Transcript ID	bp	Protein	Biotype	CCDS	UniProt	Flags
Klhl34-201	ENSMUST0000087157.4	1935 <u>644aa</u> Pr		Protein coding	CCD541191	A2A176	TSL:NA GENCODE basic APPRIS P1
Sen phann	Constants	53173			Semphann,		Cenphanaten C.
The strategy	is based on the design of	Klhl34	2-201 trans	script,the transc 1.94 kb	eription is sho	own below	Forward strand
Klhl34-201 > protein coding			-		∕D	- 10	
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### **Genomic location distribution**





## **Protein domain**



ENSMUSP00000084 MobiDB lite Low complexity (Seg) Superfamily	SKP1/BTB/POZ doma	ain superfamily		-	elch-type beta	propallar	8		•	-
SMART.	BTB/POZ domain		ociated		ich repeat type		6		М	
Pfam PROSITE profiles	BTB/POZ domain	-			Kelch repea	at type 1	12			
PIRSF	BTB/POZ domain BTB-kelch protein	n.								
PANTHER	PTHR45632									
Gene3D	PTHR45632:SF8 3.30.710.10	1,25,40,420	_	Ke	lch-type beta p	ropeller				
CDD	cd18264	_								
		cd18473	_							
All sequence SNPs/i	Sequence variants	(dbSNP and all oth	ner sources)		040		i T	3	11	
Variant Legend	missense varia									
Scale bar	0 60	120 180	240	300	360	420	480	540		64



If you have any questions, you are welcome to inquire. Tel: 400-9660890



