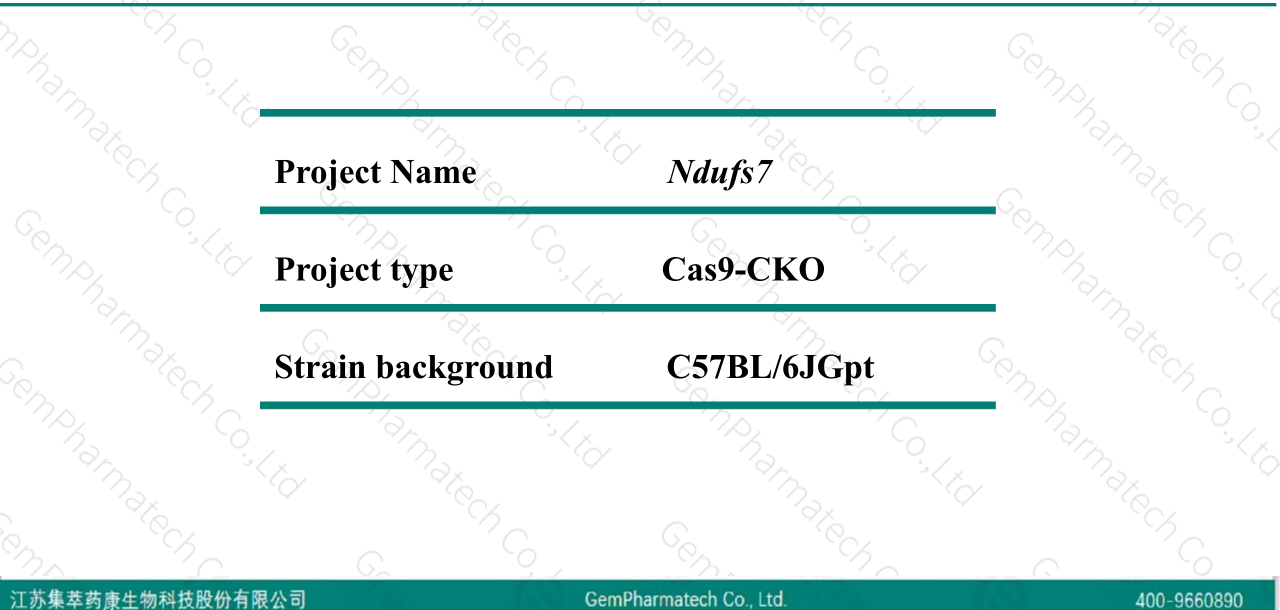


Ndufs7 Cas9-CKO Strategy

Designer:Huan FanDesign Date:2019-7-25

Project Overview

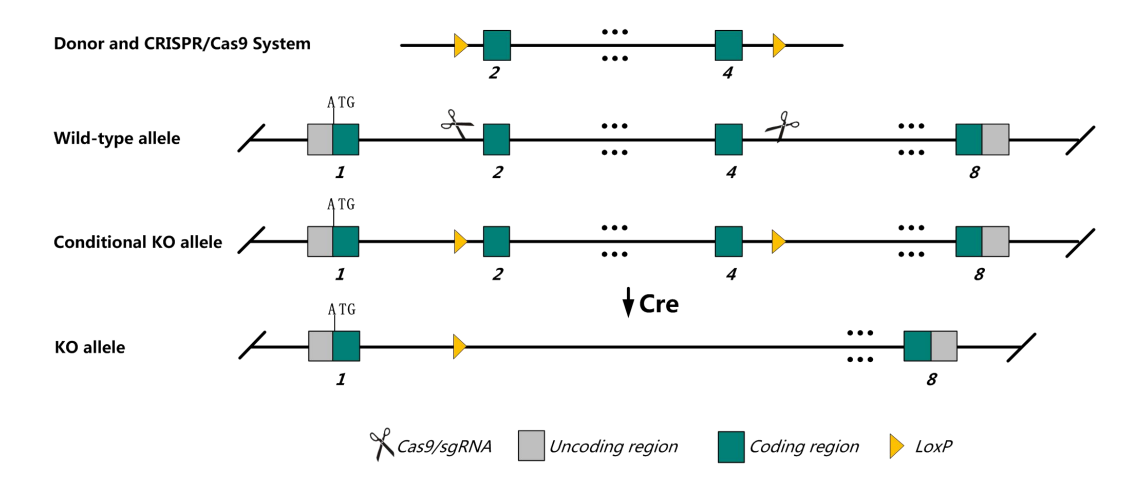




Conditional Knockout strategy



This model will use CRISPR/Cas9 technology to edit the Ndufs7 gene. The schematic diagram is as follows:





The Ndufs7 gene has 6 transcripts. According to the structure of Ndufs7 gene, exon2-4 of Ndufs7-201 (ENSMUST0000020361.6) transcript is recommended as the knockout region. The region contains 245bp coding sequence. Knock out the region will result in disruption of protein function.

In this project we use CRISPR/Cas9 technology to modify *Ndufs7* 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 Ndufs7 gene is located on the Chr10. 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)



\$?

Ndufs7 NADH:ubiquinone oxidoreductase core subunit S7 [Mus musculus (house mouse)]

Gene ID: 75406, updated on 7-Apr-2019

Summary

Official Symbol	Ndufs7 provided by MGI
Official Full Name	NADH:ubiguinone oxidoreductase core subunit S7 provided byMGI
Primary source	MGI:MGI:1922656
See related	Ensembl:ENSMUSG0000020153
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	1010001M04Rik, CI-20kD
Expression	Ubiquitous expression in heart adult (RPKM 322.2), duodenum adult (RPKM 296.5) and 28 other tissues See more
Orthologs	human all

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Transcript information (Ensembl)



The gene has 6 transcripts, all transcripts are shown below:

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Name	Transcript ID	bp	Protein	Biotype	CCDS	UniProt	Flags
Ndufs7-202	ENSMUST00000105364.7	983	<u>224aa</u>	Protein coding	CCDS24015	Q9DC70	TSL:5 GENCODE basic APPRIS P1
Ndufs7-201	ENSMUST00000020361.6	758	<u>224aa</u>	Protein coding	CCDS24015	Q9DC70	TSL:1 GENCODE basic APPRIS P1
Ndufs7-206	ENSMUST00000157063.1	722	No protein	Processed transcript	-	-	TSL:5
Ndufs7-204	ENSMUST00000155336.1	770	No protein	Retained intron	22	-	TSL:2
Ndufs7-205	ENSMUST00000155523.7	435	No protein	Retained intron		-	TSL:2
Ndufs7-203	ENSMUST00000152939.7	420	No protein	Retained intron	-	-	TSL:2

The strategy is based on the design of Ndufs7-201 transcript, The transcription is shown below

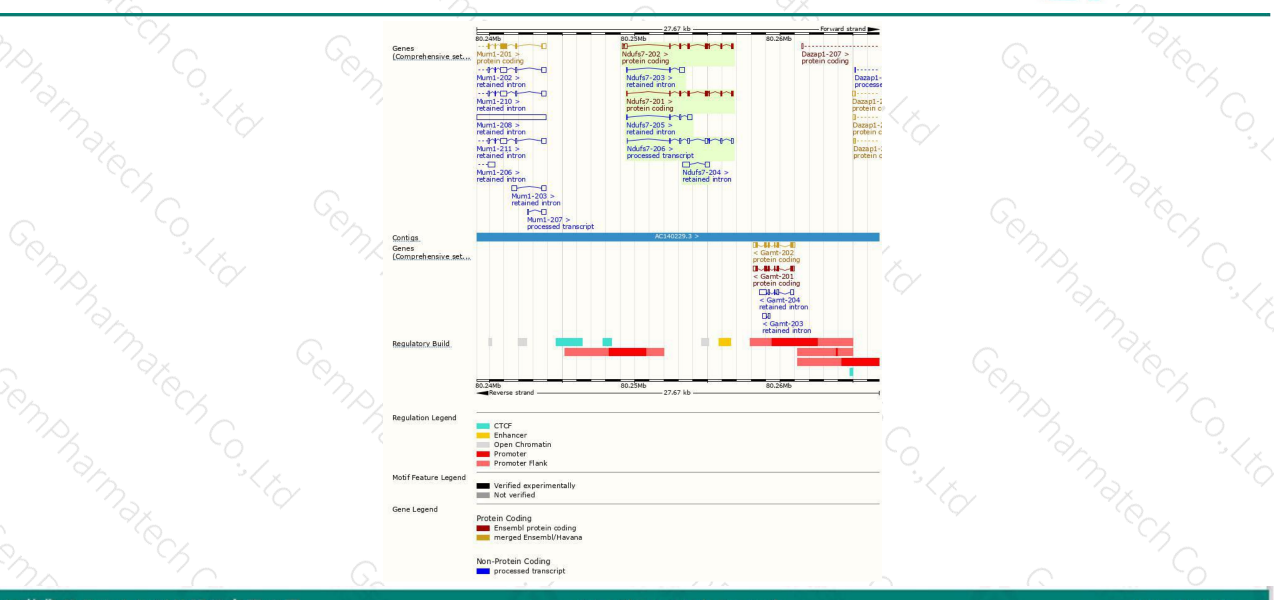
			-7.34 kb		Forward strand
Ndufs7-201 > protein coding					
	°С	~	í C	S.	6

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Genomic location distribution



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Protein domain



		NC4	1A.		1		
ENSMUSP00000020 MobiDB lite Low complexity (Seg)	1					-	20
hmmpanther		PTHR11995					
		PTHR11995:SF19					
TIGRFAM domain		NA	DH-ubiquinone oxi	doreductase,	20 Kd subunit		
Superfamily domains		SSF	56770				
Pfam domain			NADH:	u biquinone ox	doreductase-like	, 20kDa subunit	
PROSITE patterns						NADH-u big	uinone oxidored
HAMAP		NADH-ubiquinone	oxidoreductase, 2	0 Kd subunit			
Gene3D		3,40,	50,12280				
All sequence SNPs/i	Sequence variants	(dbSNP and all other s	ources)	ii -	10.1		6
Variant Legend	missense varia						
Scale bar	0 20	40 60	80 100	120	140 10	0 180	200 224
							~?~
10		$\sim O$	×>.		$\sim 2 \sim$		

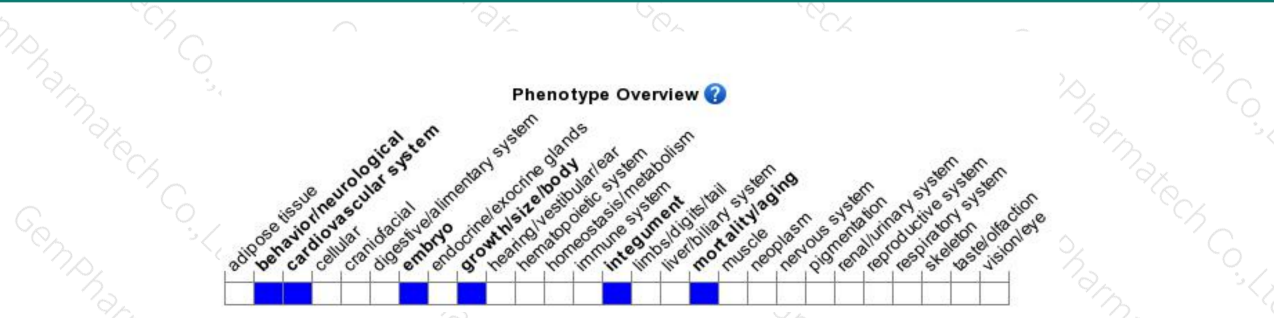
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Mouse phenotype description(MGI)





Phenotypes affected by the gene are marked in blue.Data quoted from MGI database(http://www.informatics.jax.org/).



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



