

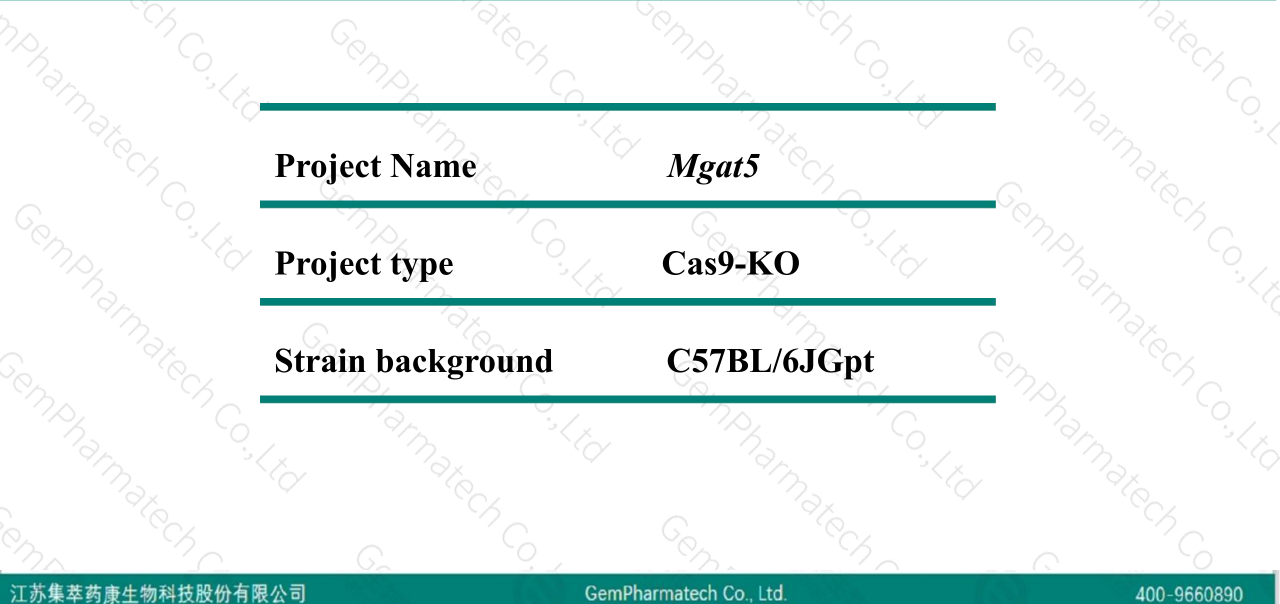
Mgat5 Cas9-KO Strategy

Designer: Reviewer: Design Date: Yang Zeng Ruirui Zhang

2019-11-22

Project Overview

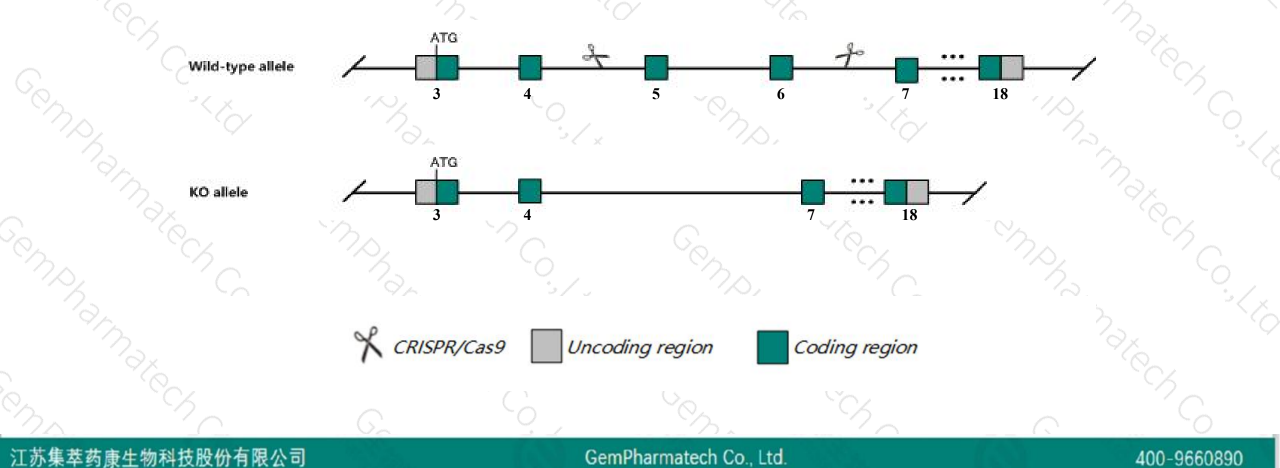




Knockout strategy



This model will use CRISPR/Cas9 technology to edit the Mgat5 gene. The schematic diagram is as follows:





- The Mgat5 gene has 4 transcripts. According to the structure of Mgat5 gene, exon5-exon6 of Mgat5-201 (ENSMUST00000038361.10) transcript is recommended as the knockout region. The region contains 167bp coding sequence. Knock out the region will result in disruption of protein function.
- > In this project we use CRISPR/Cas9 technology to modify Mgat5 gene. The brief process is as follows: CRISPR/Cas9 system

- According to the existing MGI data, Mice homozygous for deficiencies in this gene have immune system abnormalities and reduced cancer growth and metastasis.
 - The Mgat5 gene is located on the Chr1. 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 the gene knockout on gene transcription, RNA splicing and protein translation cannot be predicted at the existing technology level.

Notice

Gene information (NCBI)



☆ ?

Mgat5 mannoside acetylglucosaminyltransferase 5 [Mus musculus (house mouse)]

Gene ID: 107895, updated on 27-Aug-2019

Summary

Official SymbolMgat5 provided by MGIOfficial Full Nammannoside acetylglucosaminyltransferase 5 provided by MGIPrimary sourceMGI:MGI:894701See relatedEnsembi:ENSMUSG0000036155Gene typeprotein codingprotein codingVALIDATEDOrganianMus musculusLineageEukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Euarchontoglires; Glires; Rodentia; Myomorpha; Muroidea; Muridae;
Murinae; MusAlso known asAl480971; GlcNAc-TV; 4930471A21Rik; 5330407H02RikExpressionUbiquitous expression in colon adult (RPKM 9.9), frontal lobe adult (RPKM 9.4) and 27 other tissues See more
human all

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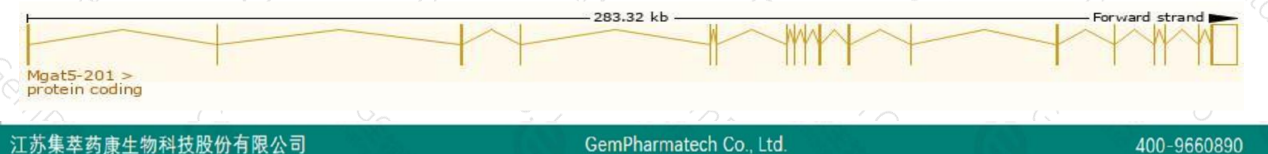
Transcript information (Ensembl)



The gene has 4 transcripts, all transcripts are shown below:

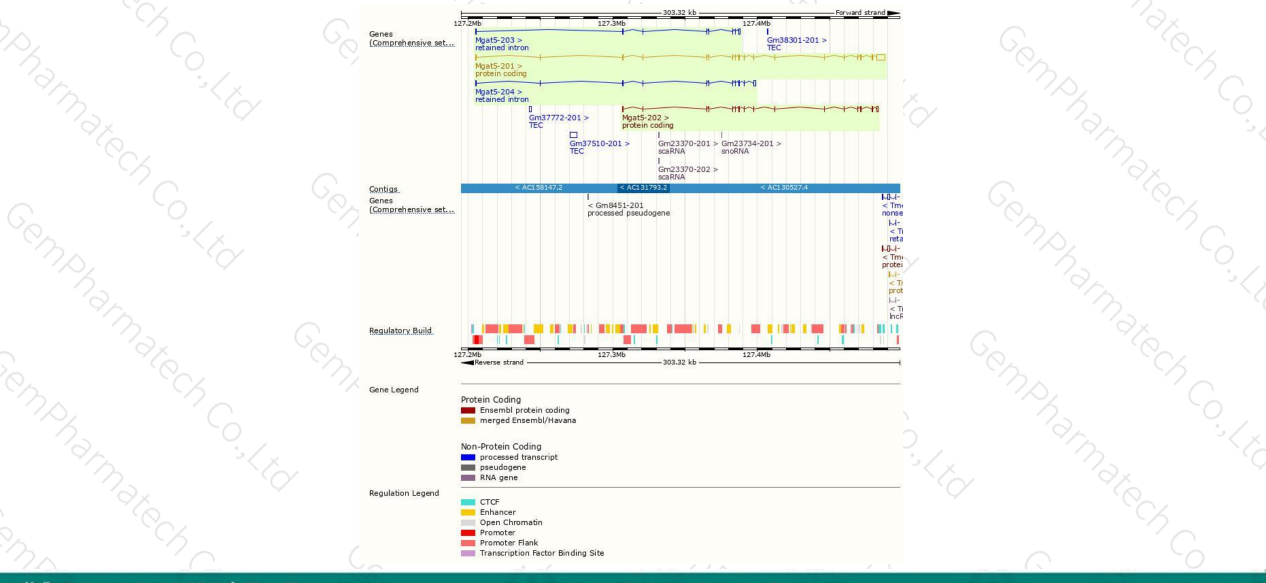
Name	Transcript ID	bp	Protein	Biotype	CCDS	UniProt	Flags
Mgat5-201	ENSMUST0000038361.10	8436	740aa	Protein coding	CCDS15245	Q059T5 Q8R4G6	TSL:1 GENCODE basic APPRIS P1
Mgat5-202	ENSMUST00000171405.1	3216	<u>740aa</u>	Protein coding	CCDS15245	Q059T5 Q8R4G6	TSL:1 GENCODE basic APPRIS P1
Mgat5-204	ENSMUST00000190921.1	2984	No protein	Retained intron	2	141	TSL:1
Mgat5-203	ENSMUST00000189427.6	2229	No protein	Retained intron	2	120	TSL:1
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The strategy is based on the design of Mgat5-201 transcript, The transcription is shown below



Genomic location distribution





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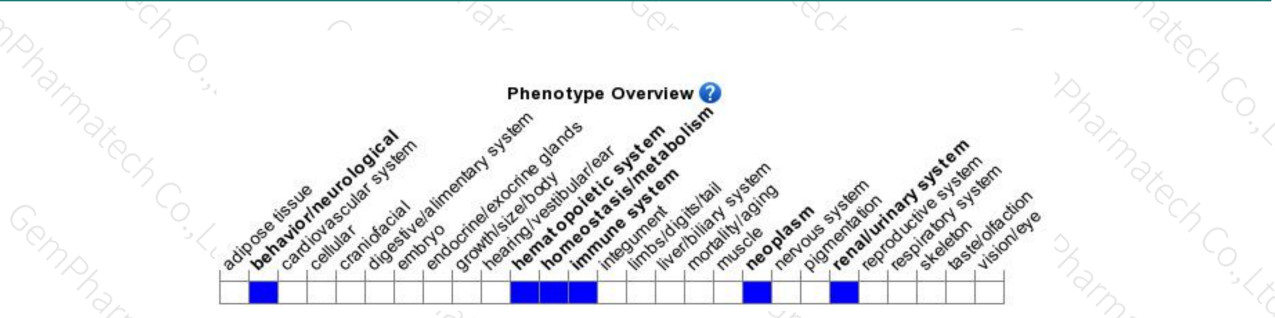
Protein domain



		C.C.M.S.	°° S			G. C.	Enjoy ~ (C)	3		
	ENSMUSP00000038 Transmembrane heli Low complexity (Seg) Pfam	Domain of unknown	function DUF4525					<		
Cen.	PANTHER	PTHR15075:SF5	Glycosyltrans	ferase family 18				6,		
	All sequence SNPs/i	PTHR15075 Sequence variants (dbSNP and all other sources)								
Sent	Variant Legend	missense variant splice region variant synonymous variant								
	Scale bar	0 80	160 :	240 320	400	480 560	640 74	r ×¢		
	Nate Ch	9 G	Con Con	Gen		``A	C AC			
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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/).

According to the existing MGI data, Mice homozygous for deficiencies in this gene have immune system abnormalities and reduced cancer growth and metastasis.



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



