

Slc22a6 Cas9-CKO Strategy

Designer:

Reviewer:

Design Date:

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2019-11-29

Project Overview



Project Name

Slc22a6

Project type

Cas9-CKO

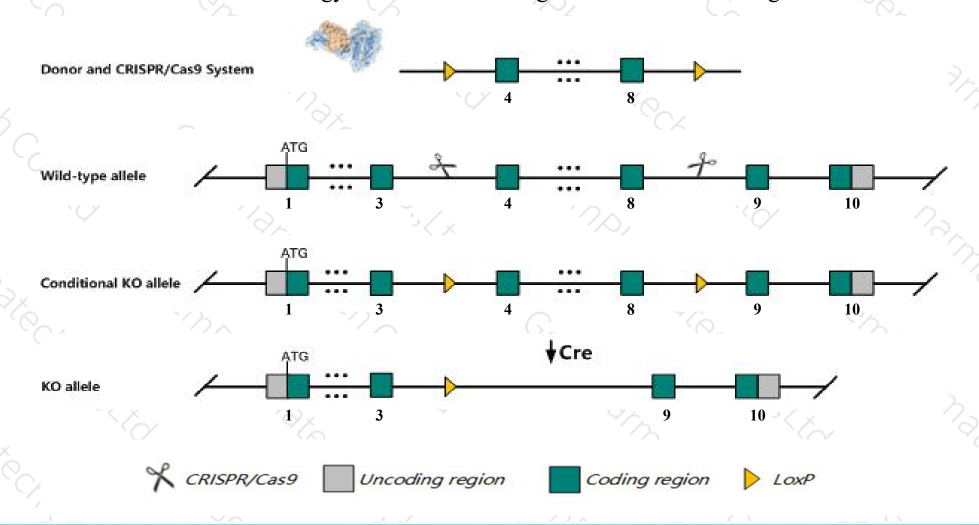
Strain background

C57BL/6JGpt

Conditional Knockout strategy



This model will use CRISPR/Cas9 technology to edit the Slc22a6 gene. The schematic diagram is as follows:



Technical routes



- ➤ The Slc22a6 gene has 1 transcript. According to the structure of Slc22a6 gene, exon4-exon8 of Slc22a6-201 (ENSMUST00000010250.3) transcript is recommended as the knockout region. The region contains 733bp coding sequence. Knock out the region will result in disruption of protein function.
- In this project we use CRISPR/Cas9 technology to modify *Slc22a6* 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.

Notice



- > According to the existing MGI data, Homozygous mutation of this gene may result in increased thymus weight or impaired renal organic anion excretion for a subset of organic anions.
- > The Slc22a6 gene is located on the Chr19. 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)



Slc22a6 solute carrier family 22 (organic anion transporter), member 6 [Mus musculus (house mouse)]

Gene ID: 18399, updated on 10-Oct-2019

Summary



Official Symbol Slc22a6 provided by MGI

Official Full Name solute carrier family 22 (organic anion transporter), member 6 provided by MGI

Primary source MGI:MGI:892001

See related Ensembl: ENSMUSG00000024650

Gene type protein coding
RefSeq status VALIDATED

Organism Mus musculus

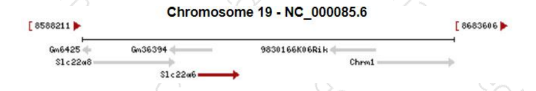
Lineage Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Euarchontoglires; Glires; Rodentia; Myomorpha;

Muroidea; Muridae; Murinae; Mus; Mus

Also known as NKT; Oat1; mOat1; Orctl1

Expression Restricted expression toward kidney adult (RPKM 141.9) See more

Orthologs human all



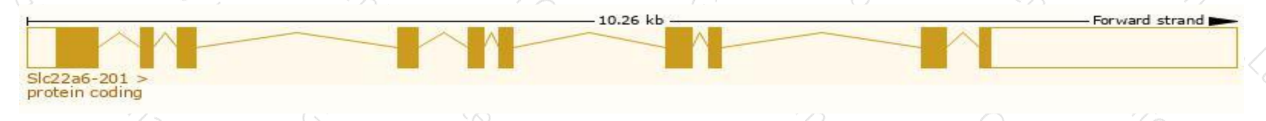
Transcript information (Ensembl)



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

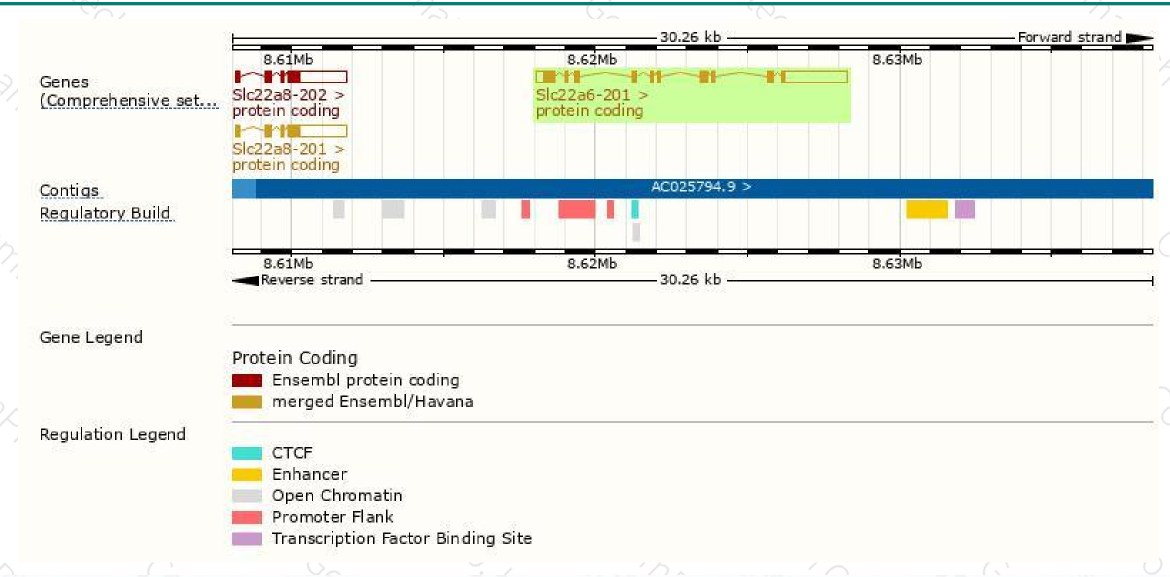
Name 🍦	Transcript ID 👙	bp 🌲	Protein 🍦	Translation ID 🝦	Biotype 🍦	CCDS 🍦	UniProt 🌲	Flags		
Slc22a6-201	ENSMUST00000010250.3	3983	<u>545aa</u>	ENSMUSP00000010250.2	Protein coding	CCDS29538₽	Q8VC69₽	TSL:1	GENCODE basic	APPRIS P1

The strategy is based on the design of Slc22a6-201 transcript, The transcription is shown below



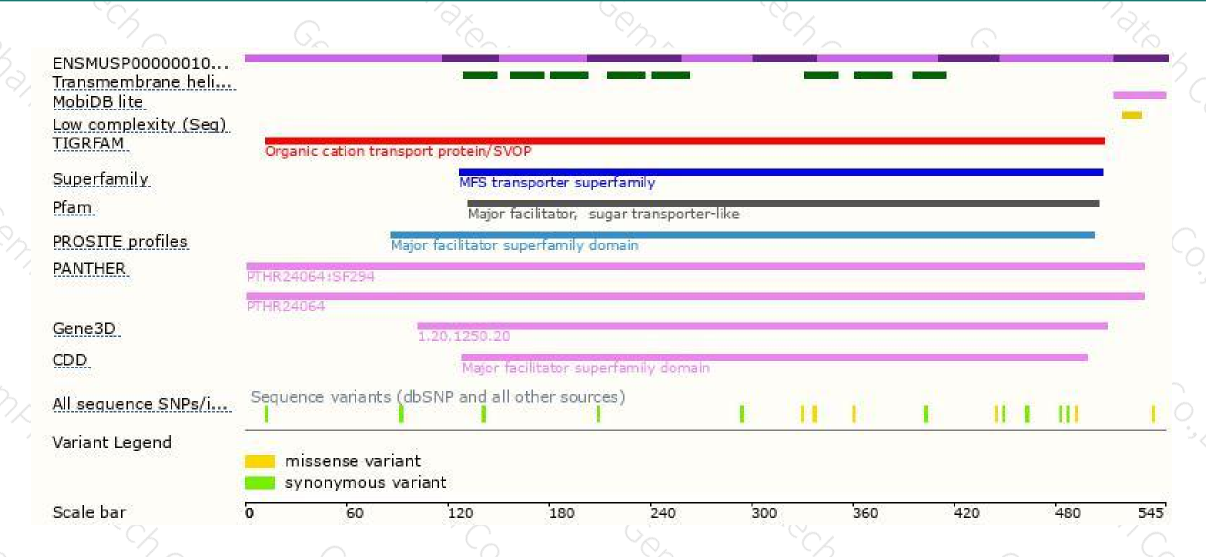
Genomic location distribution





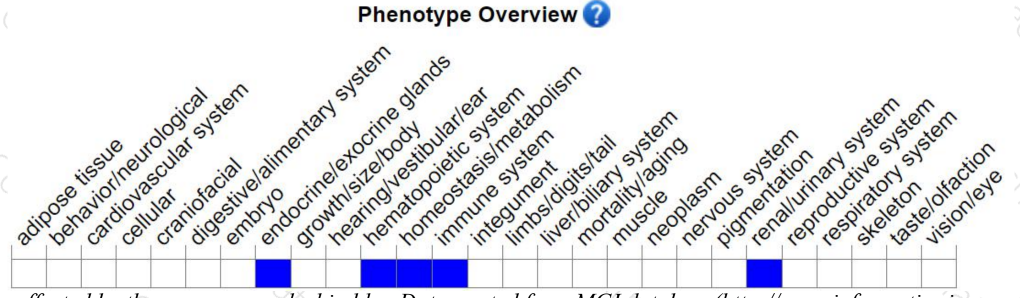
Protein domain





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, Homozygous mutation of this gene may result in increased thymus weight or impaired renal organic anion excretion for a subset of organic anions.



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





