

Pard3 Cas9-KO Strategy To hall alto color color

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Project Overview



Project Name Pard3

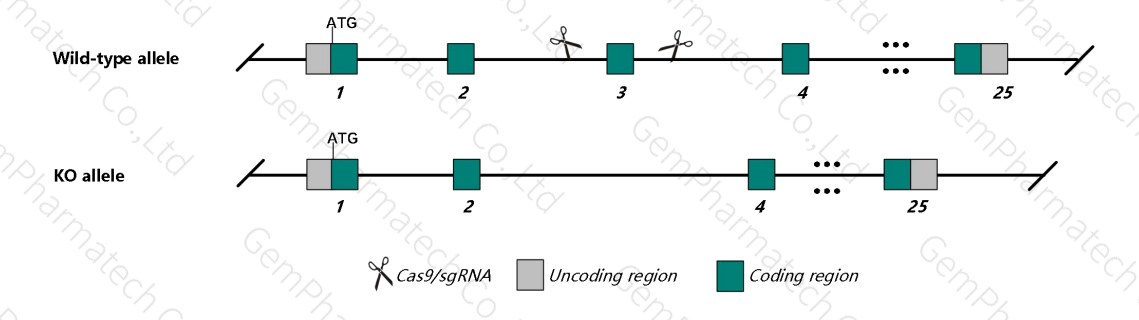
Project type Cas9-KO

Strain background C57BL/6J

Knockout strategy



This model will use CRISPR/Cas9 technology to edit the *Pard3* gene. The schematic diagram is as follows:



Technical routes



- ➤ The *Pard3* gene has 28 transcripts. According to the structure of *Pard3* gene, exon3 of *Pard3-201* (
 ENSMUST00000026921.12) transcript is recommended as the knockout region. The region contains 181bp coding sequence.

 Knock out the region will result in disruption of protein function.
- ➤ In this project we use CRISPR/Cas9 technology to modify *Pard3* gene. The brief process is as follows: sgRNA was transcribed in vitro.Cas9 and sgRNA were microinjected into the fertilized eggs of C57BL/6J 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/6J mice.

Notice



- ➤ According to the existing MGI data, Mice homozygous for a null allele exhibit embryonic lethality at E12.5 associated with growth retardation, abnormal heart development, and abnormal epicardial cell development.
- ➤ The *Pard3* gene is located on the Chr8. 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 gene transcription and translation processes, all risks cannot be predicted under existing information.

Gene information (NCBI)



Pard3 par-3 family cell polarity regulator [Mus musculus (house mouse)]

Gene ID: 93742, updated on 7-Apr-2019

Summary

☆ ?

Official Symbol Pard3 provided by MGI

Official Full Name par-3 family cell polarity regulator provided by MGI

Primary source MGI:MGI:2135608

See related Ensembl: ENSMUSG00000025812

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 AA960621, Al256638, Asip, D8Ertd580e, Par-3, Par3, Pard-3, Pard3a, Phip

Expression Ubiquitous expression in ovary adult (RPKM 7.1), bladder adult (RPKM 6.3) and 28 other tissuesSee more

Orthologs <u>human</u> all

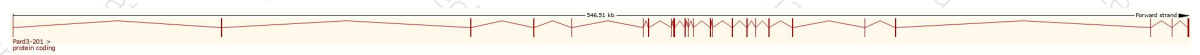
Transcript information (Ensembl)



The gene has 28 transcripts, all transcripts are shown below:

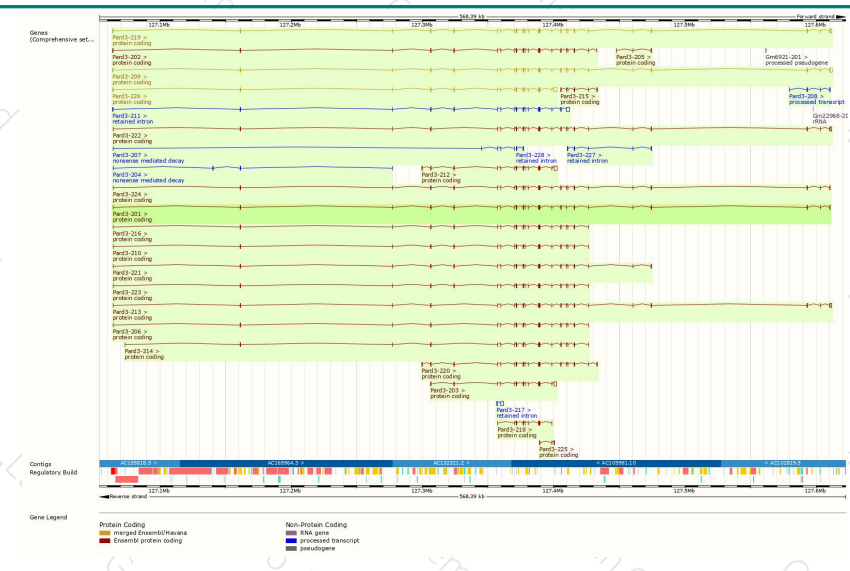
Name	Transcript ID	bp	Protein	Biotype	CCDS	UniProt	Flags
ard3-209	ENSMUST00000160272.7	5815	1334aa	Protein coding	CCDS80945@	A5D6P2Ø	TSL:1 GENCODE basic APPRIS ALT2
Pard3-219	ENSMUST00000162309.7	5659	<u>1333aa</u>	Protein coding	CCDS22788₽	G3XA13₽	TSL:1 GENCODE basic APPRIS P3
ard3-226	ENSMUST00000162907.7	4682	741aa	Protein coding	CCDS52712@	E9PYJ2₽	TSL:1 GENCODE basic APPRIS ALT2
Pard3-212	ENSMUST00000160717.7	4214	606aa	Protein coding	CCDS40523₽	Q99NH2₽	TSL:1 GENCODE basic
ard3-201	ENSMUST00000026921.12	4050	<u>1319aa</u>	Protein coding	CCDS80946₽	B7ZNY3Ø	TSL:1 GENCODE basic APPRIS ALT2
Pard3-203	ENSMUST00000108752.9	3497	606aa	Protein coding	CCDS40523₽	Q99NH2@	TSL:5 GENCODE basic
Pard3-202	ENSMUST00000079777.11	3394	896aa	Protein coding	CCDS22789₽	A0A0R4J1Y4®	TSL:5 GENCODE basic
ard3-220	ENSMUST00000162456.7	3204	896aa	Protein coding	CCDS22789@	A0A0R4J1Y4@	TSL:1 GENCODE basic
ard3-222	ENSMUST00000162536.7	5175	1289aa	Protein coding	6-9	E0CXL4⊌	TSL:5 GENCODE basic APPRIS ALT2
Pard3-213	ENSMUST00000160766.7	4949	1247aa	Protein coding	323	E0CY24®	TSL:5 GENCODE basic APPRIS ALT2
Pard3-224	ENSMUST00000162665.7	4005	1324aa	Protein coding	6-9	<u>F6S7Z1</u> ₽	CDS 5' incomplete TSL:1
ard3-221	ENSMUST00000162531.7	3293	<u>1094aa</u>	Protein coding	-	E0CZ83₽	CDS 3' incomplete TSL:5
ard3-223	ENSMUST00000162602.7	3083	1024aa	Protein coding	17-1	E0CZE2₽	CDS 3' incomplete TSL:5
ard3-216	ENSMUST00000161355.7	2981	990aa	Protein coding	323	E0CXR8₽	CDS 3' incomplete TSL:5
ard3-210	ENSMUST00000160581.7	2954	984aa	Protein coding	6-9	F6TSJ9₽	CDS 5' and 3' incomplete TSL:5 APPRIS ALT
ard3-206	ENSMUST00000159537.7	2838	943aa	Protein coding	72	E0CX45₽	CDS 3' incomplete TSL:5
ard3-214	ENSMUST00000161277.7	2758	919aa	Protein coding	6-8	F6UGU7₽	CDS 5' and 3' incomplete TSL:5
ard3-218	ENSMUST00000162176.1	1266	422aa	Protein coding		Q4JJC0@	CDS 5' and 3' incomplete TSL:1
Pard3-215	ENSMUST00000161348.2	1124	269aa	Protein coding	6-9	F6URI7₽	CDS 5' incomplete TSL:5
ard3-225	ENSMUST00000162727.1	622	116aa	Protein coding	828	F6TB42₽	CDS 5' incomplete TSL:2
ard3-205	ENSMUST00000159511.1	307	102aa	Protein coding	6-9	<u>F6T4Y3</u> ₽	CDS 5' and 3' incomplete TSL:3
Pard3-207	ENSMUST00000159818.1	568	44aa	Nonsense mediated decay	323	E0CYB6₽	TSL:5
Pard3-204	ENSMUST00000159141.7	520	<u>59aa</u>	Nonsense mediated decay	17-1	EDCXD3@	TSL:2
ard3-208	ENSMUST00000159940.1	746	No protein	Processed transcript	120	120	TSL:2
Pard3-211	ENSMUST00000160593.7	5310	No protein	Retained intron	6-3	(1-0)	TSL:1
Pard3-217	ENSMUST00000162035.1	3017	No protein	Retained intron	25	727	TSL:1
Pard3-227	ENSMUST00000163002.1	815	No protein	Retained intron	0-9	(1-0)	TSL:3
Pard3-228	ENSMUST00000163021.1	614	No protein	Retained intron	25	120	TSL:2

The strategy is based on the design of Pard3-201 transcript, The transcription is shown below



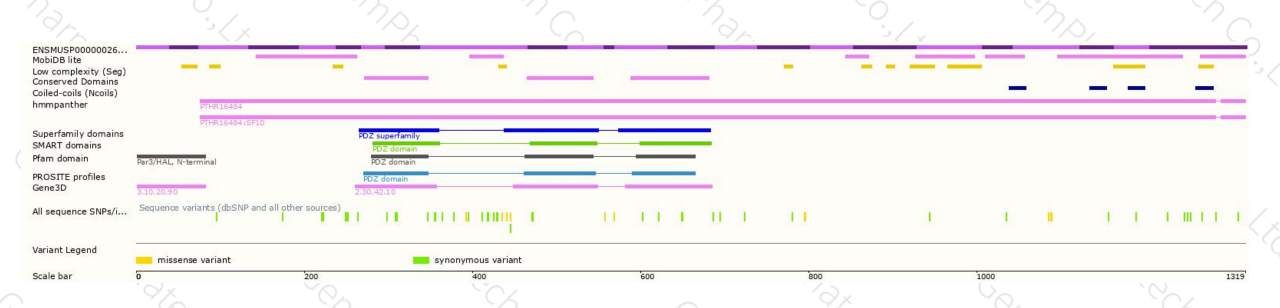
Genomic location distribution





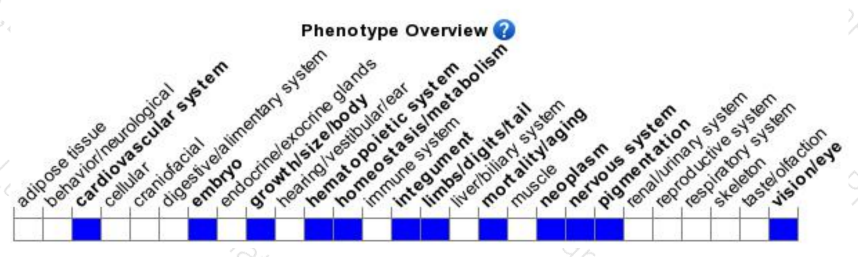
Protein domain





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, Mice homozygous for a null allele exhibit embryonic lethality at E12.5 associated with growth retardation, abnormal heart development, and abnormal epicardial cell development.



If you have any questions, you are welcome to inquire.

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