

# Col4a5-R471X

**Nomenclature** C57BL/6Smoc-*Col4a5*<sup>em1(R471X)Smoc</sup>

**Cat. NO.** NM-KI-200183

Strain State Repository Live

### **Gene Summary**

Gene Symbol Col4a5	Synonyms	-
	NCBI ID	12830
	MGI ID	<u>88456</u>
	Ensembl ID	ENSMUSG00000031274
	Human Ortholog	COL4A5

## **Model Description**

These mice carry a R471X mutation of Col4a5 gene.

**Research Application**: Alport syndrome research

\*Literature published using this strain should indicate: Col4a5-R471X mice (Cat. NO. NM-KI-200183) were purchased from Shanghai Model Organisms Center, Inc..

#### **Disease Connection**

X-Linked Alport Syndrome	Phenotype(s)	MGI:6479076
	Reference(s)	Hashikami K, et al., Establishment of X- linked Alport syndrome model mice with a Col4a5 R471X mutation. Biochem Biophys Rep. 2019 Mar;17:81-86

### **Validation Data**



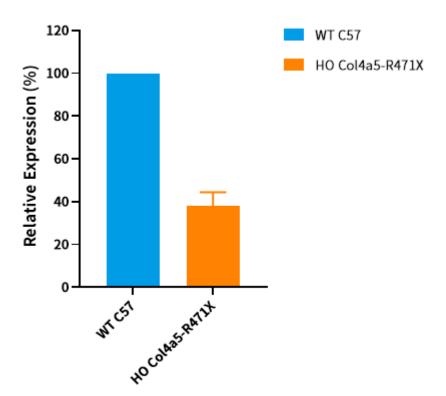


Fig1. *Col4a5* mRNA level was measured in Col4a5-R471X mice (n=3, male, 6 week-old) and the point mutation of *Col4a5* has been verified by sequencing.

Abbr. HO, homozygous; WT, wild type.

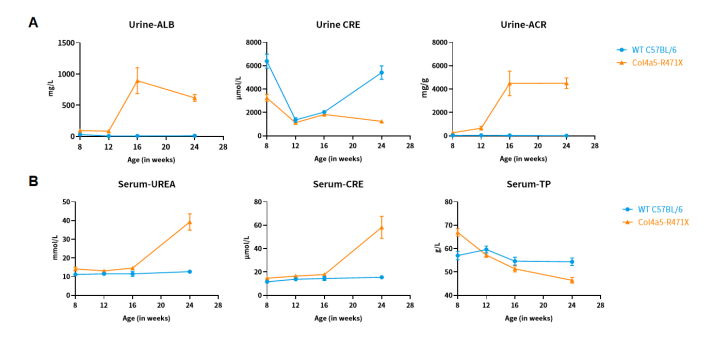


Fig2. The results of urine (A) and blood (B) biochemical indicators in Col4a5-R471X mice (n=2 male and 6 female).

Abbr. WT, wild type.



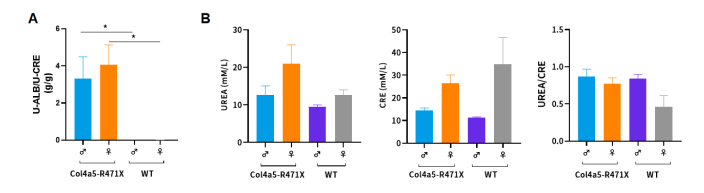


Fig3. The results of urine (A) and plasma (B) biochemical indicators in 21-weeks-old Col4a5-R471X mice (n=3/group).(Data from a cooperator)

Abbr. WT, wild type.