

# KPC

系統名	C57BL/6Smoc- <i>Trp53<sup>em4(R172H)</sup>Kras<sup>em4(LSL-G12D)</sup></i> Tg(Pdx1-cre)Smoc
SMOC番号	NM-KI-210096
維持形態	Repository Live

## 遺伝子の概要

Gene Symbol Trp53	Synonyms	bbl; bfy; bhy; p44; p53; Tp53
	NCBI ID	<u>22059</u>
	MGI ID	<u>98834</u>
	Ensembl ID	ENSMUSG0000059552
	Human Ortholog	TRP53
Gene Symbol Kras	Synonyms	Ki-ras; K-ras; Kras2; Kras-2
	NCBI ID	<u>16653</u>
	MGI ID	<u>96680</u>
	Ensembl ID	ENSMUSG0000030265
	Human Ortholog	KRAS
Gene Symbol pdx1	Synonyms	lpf1, IDX-1, IPF-1, Mody4, STF-1, pdx-1
	NCBI ID	<u>18609</u>
	MGI ID	<u>102851</u>
	Ensembl ID	ENSMUSG0000029644

#### 説明

The KPC mouse is an established and clinically relevant model of pancreatic ductal adenocarcinoma (PDAC) which develops many key features observed in human PDAC including pancreatic intraepithelial neoplasia alongside a robust inflammatory reaction including exclusion of effector T cells. Metastases are observed in around 80% of KPC animals located primarily in the liver and lungs. Mutations in both KRAS and TP53 genes are found in around 80% and 70% of all human PDACs respectively. Trp53-R172H (NM-KI-18028)、Kras-LSL-G12D (NM-KI-190003)were crossed with Pdx1-Cre-Tg to generate KPC mice. The KPC mouse contains a dominant negative point mutation in the transformation related protein 53 gene (TP53R172H),

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and a conditional activation point mutation in the KRAS gene (KRASG12D). A lox-stop-lox termination sequence is encoded upstream of KRAS mutated genes to prevent expression in the absence of Cre recombinase. The pancreas-specific Pdx-1 promoter enables expression of Cre recombinase in acini, islet and duct cells of the pancreas. Cre-mediated recombination excises the lox-stop-lox termination sequences and enables expression of KRASG12D in pancreatic tissue.

#### 応用分野: Spontaneous pancreatic tumor

\*Literature published using this strain should indicate: KPC mice (Cat. NO. NM-KI-210096) were purchased from Shanghai Model Organisms Center, Inc..

### 病気の予測

	表現型	
Pancreatic Carcinoma	参考文献	Pancreatology . 2020 Jan;20(1):79-88. doi: 10.1016/j.pan.2019.11.006. Epub 2019 Nov 18.

## 表現型デロタ



Fig 1 The spontaneous pancreatic tumor of KPC mouse model with large volume, uneven surface and multiple nodular projections.



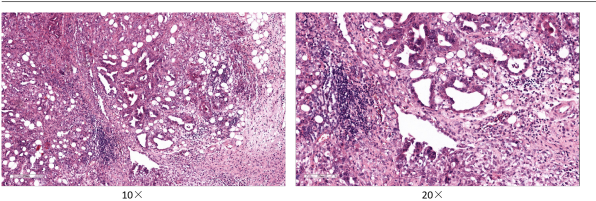


Fig 2 HE staining of pancreatic tumor from KPC mouse model.

The tumor cells from KPC mouse model demonstrated disorderly arranged pancreatic cells, irregular tissue structure, dilated pancreatic ducts, inflammatory cells infiltration and stromal fibrosis as was seen in pancreas adenocarcinoma.