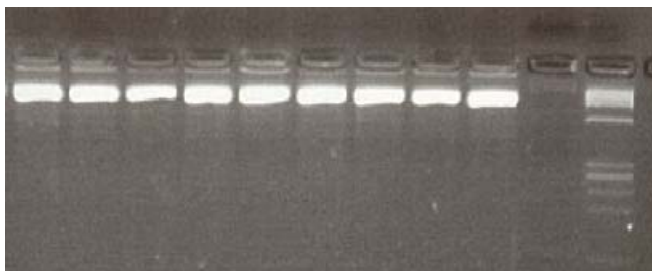


Transgene	Cre
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primer	Sequence(5'-3')	Size	target gene	accession
px1679/1701 px2802/2780	GCTGGTTGTTGTGCTGTCTCATC ACCATTGCCCTGTTTCACTATC		Cre	
comment				

PCR condition							
Taq polymerase	BIOTAQ™ HS DNA Polymerase(BIOLINE, London, UK)						
Thermal cycler	PC-808(ASTECC)						
PCR buffer	Ampdirect Plus						
	first denature	denature	anneal	extension	cycle	final extension	
PCR	°C	94	94	55	72	35	72
	min	3	0.5	1	0.45		3
comment	PCR method; Blood was applied to FTA®card (GE Healthcare UK Ltd.,UK) and dried. 1.5mm FTA disc was removed from the bloody-stained region on FTA®card. Untreated sample discs were placed directly in 15 μL PCR mixture containig 1 × Ampdirect®Plus, 0.2 μM each primer and 0.4 units of BIOTAQ™ HS DNA Polymerase.						

Electrocataphoresis ;4% agarose gel 300V, 30min	
	<p>1 2 3 4 5 6 7 8 <math>\lambda</math> M</p> 
Strain	W-Tg(CAG-cre)81Jmsk