

GENOTYPING PROTOCOL

MUTANT MOUSE RESOURCE & RESEARCH CENTER: UC DAVIS

mmrrc@ucdavis.edu

530-754-MMRRC

Protocol Name: CR11674 Nudt8 EXDEL

Stock # 75942

Protocol: GoTaq® G2 Colorless Master Mix(Promega)

Reagent/Constituent	Volume (µL)
Water	4.5
GoTaq® G2 Colorless Master Mix,2X	7.5
Primer 1. (stock concentration is 20µM) comF	0.5
Primer 2. (stock concentration is 20µM) wtR	0.5
Primer 3. (stock concentration is 20µM) mutR	0.5
DNA (example) extracted w/ "Qiagen DNeasy columns or other similar silica based kits"	1.5
TOTAL VOLUME OF REACTION:	15.00 µL

Comments on protocol:

- Protocol may work with other DNA extraction methods.

Strategy:

Steps	Temp (°C)	Time (m:ss)	# of Cycles
1. Initiation/Melting HOT START? <input type="checkbox"/>	94	2:00	1x
2. Denaturation	94	0:10	
3. Annealing steps 2-3-4 cycle in sequence	65 (↓1°C/cycle)	0:30	10x
4. Elongation	68	2:00	
5. Denaturation	94	0:15	
6. Annealing steps 5-6-7 cycle in sequence	55	0:30	25x
7. Elongation	68	2:00 (↑20sec/cycle)	
8. Finish	4	∞	n/a

Primers:

Name	Nucleotide Sequence (5' - 3')	Electrophoresis Protocol:		
1. CR_Nudt8_comF	CCTGGTTGGGAGACACAAAGGG	Agarose: 1.5%	V: 90	
2. CR_Nudt8_wtR*	TACCGCCTGGGAAACTAAATGG	Estimated Running Time: 90 min.		
3. CR_Nudt8_mutR	AGGGTGAAGAGTCAGAAGGAGC	Primer Combination	Band (bp)	Genotype
		1 & 2, 1 & 3	392,798	wildtype
		1 & 3	328	mutant

Allele Description: Exon 2 ENSMUSE00001398765 and flanking splicing regions were constitutively deleted from the Nudt8 Gene ENSMUSG00000110949 using CRISPR Cas9 gene editing technology in mouse zygotes. The subsequent 470bp deletion from Chr 19: 4051007 - 4051476 GRCm39 was screened by PCR analysis. The selected founder animal was backcrossed to C57BL6/N to produce sequence confirmed heterozygous animals for establishing and archiving the line.

*wtR primer untested (ePCR verified)

