

# GENOTYPING PROTOCOL

## MUTANT MOUSE RESOURCE & RESEARCH CENTER: UC DAVIS

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530-754-MMRRC

Protocol Name: CR10545 Mme N689K HDR Stock #: 66744

Reagent/Constituent	Volume (μL)
QuantiTect Multiplex PCR Master Mix Cat No./ID 204541	5.0
Water	3.4
<b>Target Probe mix</b>	0.3
-21 μM Mutant Forward Primer	
-21 μM Mutant Reverse Primer	
-7 μM Mutant probe	
<b>TCRD (endogenous control) mix</b>	0.3
-21 μM WT Forward primer	
-21 μM WT Reverse Primer	
-7 μM WT probe	
Sample	1.0
<b>TOTAL VOLUME OF REACTION:</b>	<b>10.00 μL</b>

### Comments on protocol:

Protocol may work with other DNA extraction methods. WT Vic probe may be substituted for WT Orange 540 probe. Reference: ABI User Bulletin #2 and #5 (updated 10/2001) for multiplex in same tube and validation of each assay to match relative efficiencies of reference and target primer/probe combinations. Also reference: Rapid and accurate determination of zygosity in transgenic animals by real-time quantitative PCR. TransRes (2002).

### Strategy:

Steps	Temp (°C)	Time (m:ss)	# of Cycles
1. Initiation/Melting	95	15:00	1
2. Denaturation	95	0:30	40x
3. Annealing/Elongation	60	1:00	40x
4. To step 2 for 40 cycles			

### Primers:

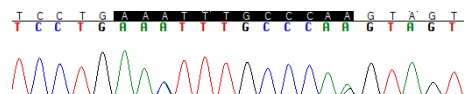
Name	Nucleotide Sequence (5' - 3')
1. TM_Mme_WT-F	GACTTGACCTCAATCACAAACAAC
2. TM_Mme_WT-R	CATATTTATCAATCATGTAAGACATACTACC
3. Mme-WT Orange 560 BHQ-1 Probe	Orange 560-pdC-pdC-pdU-GAA-pdC-pdU-pdU-pdU-G-pdC-pdC-pdC-AGG-pdU-BHQ-1
4. TM_Mme_KI-F	GACTTGACCTCAATCACAAACAAC
5. TM_Mme_KI-R	TTCATATTTATCAATCATGTAAGACATACTACT
6. Mme-KI Fam BHQ-1 Probe	Fam-pdU-pdU-pdU-pdU-pdC-pdC-pdU-GAAA-pdU-pdU-pdU-G-pdC-pdC-pdC-AAG-pdU-BHQ-1

**Allele Description:** The mouse N689K model was created using optimized CRISPR Cas9 KI technology utilizing Ribonucleoprotein (RNP) in the presence of a synthetic single strand DNA repair template harboring the desired KI. Zygotes were electroporated and subsequent progeny were screened for the presence of the correctly targeted allele via homology directed repair (HDR). The KI nucleotide is 4 bp from the cleavage site, and one silent PAM mutations was engineered into ssODN to prevent cleavage of the KI allele by Cas9. Key progeny were sequence confirmed.

**aagaatggtgaagaaaaattactccctggacttgacctaatacacaacaactattttctgaaAttgcccAgtagtagtctctacatgattgataaatatgaaaagttgtaatttc**

WT AAC > AAA KI

Sample	ΔCt	Genotype
Mme-ntc		No Rxn
Mme-WT	0.89	WT
CR10545-56	-0.38	Het
CR10545-57	-6.15	Hom



Het

aaCttgcccA
WT
aaAttgcccA
KI

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### Alternative Genotyping Protocol

### Standard PCR and Sequencing

**Protocol:** *GoTaq® G2 Colorless Master Mix(Promega)*

Reagent/Constituent	Volume (μL)
Water	5.0
GoTaq® G2 Colorless Master Mix,2X	7.5
Primer 1. (stock concentration is 20μM) IVF	0.5
Primer 2. (stock concentration is 20μM) IVR	0.5
DNA (example) extracted w/ "Qiagen DNeasy columns or other similar silica based kits"	1.5
<b>TOTAL VOLUME OF REACTION:</b>	<b>15.0 μL</b>

**Comments on protocol:**

- Protocol may work with other DNA extraction methods.
- When crossing Alg13 and Glt28d2 the Taqman picks up the "other" mutation. Sequencing will be needed if mixed.

**Strategy:**

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

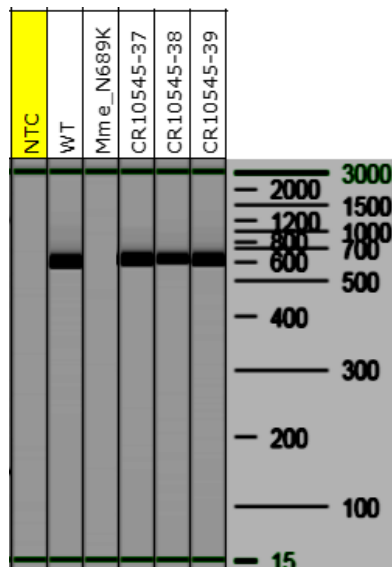
**Primers:**

Name	Nucleotide Sequence (5' - 3')	Agarose: 1.5% : 90
1. CR-Mme_N689K-IVF	GACATTCAGTGTGATCATCTGACATGC	Estimated 90 min.
2. CR-Mme_N689K-IVR	GGAAGTAACAGAGTCAACACTGGTGTAGC	<b>Primer</b> <b>Band (bp)</b> <b>Seq Primer</b>
		1 & 2 774 CR-Mme_N689K-IVF

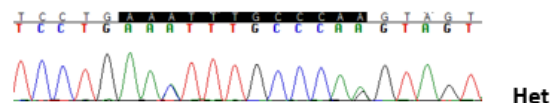
**Electrophoresis Protocol:**

Sequencing across Exon 21 to verify the HDR using standard PCR and PCR purification is used to determine the presence of HDR.

Note: Pup #38 was not sequenced as previous TM data indicated that this pup did not have HDR. It is necessary to sequence all WT-sized bands to verify HDR when using only standard PCR.



Sample	Genotype
CR10545-37	Het
CR10545-39	Het



aaCtttgcccaG	WT
aaAttgcccA	KI