

Gene: Tgm3

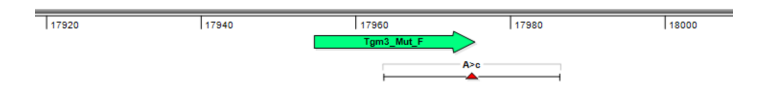
Colony prefix: DAGG

Allele: Tgm3<sup>em2(IMPC)Wtsi</sup>

Allele type: Crispr/Cas9 mediated Point mutation

Allele information: 1163L

Further information about the allele can be found on the 'International Mouse Phenotyping Consortium' (IMPC) web site at <u>http://www.mousephenotype.org/</u>



## Mouse QC information

Loss of WT Allele (LOA) qPCR	na	Mutation Sequence confirmed	pass
Mutant Specific SR- PCR	na	Off-target analysis complete	na

# Guide RNAs and mutant oligos used in initial experiment

Sequence	Chr	Chr Start	Chr End
GATTCTGGCATCATCTATGTGGG	2	130025311	130025333
TCCATTTACCTGTCCAAAGTTCCAGCCAACCATGCCAATTCGAT		130025266	130025385
TTGTGCTGCCCACATAGAGGATGCCAGAATCTTCTTCAACATAC			
TCTTGTCTTTCGGCGTGGTTACTCATAAAGAC			

This technical data sheet and information ("Datasheet") is supplied by Genome Research Limited ("GRL").

Although reasonable care is taken in the preparation of this Datasheet, GRL gives no warranties express or implied for any use of the Datasheet or for the accuracy of the Datasheet. GRL assumes no responsibility or liability for any decisions based upon the Datasheet. Without limiting the foregoing the Datasheet was prepared for mice supplied directly from GRL and where copies of this Datasheet are available from third party repositories or distribution centres ("Third Parties") GRL shall not be liable for any inconsistency between the mouse strain supplied by the Third Party and the Datasheet howsoever arising.

Report Generated on: 12-August-2020

Page 1 of 4



## Mutant allele sequence:

TCTCTCTCCCCCCCCCCCCCCCCTCTTGTCAAACAGCGGATGATGTCTTTATGAGTAACCACGCCGAAAGACAAGAGTATGTTGAAGAAGATTCTGGCATC[A/C]TCTATGTGGGCAGCACAAATCGAATTGGCATGGTTGGCTGGAACTTTGGACAGGTAAATGGACCATAAGACTTGGGTTGTCTCCCTAATACACAAGATGCTTGGGGCTTATAGTA

## Genotyping by end-point PCR

#### PCRs primer pairs and expected size bands

Assay Type	Assay	Forward Primer	Reverse Primer	Expected Size Band (bp)
Standard PCR	Screening*	Tgm3_WT_F	Tgm3_R	336
Standard PCR	Wildtype	Tgm3_WT_F2	Tgm3_R	183
Standard PCR	Mutant	Tgm3_Mut_F	Tgm3_R	183

<sup>\*</sup>The screening PCR flanks the SNP region and can be used for sequence verification of the allele. The PCR will not distinguish wild type from mutant mice, however, as a product will be amplified in all cases.

We recommend that mice are sequence-verified with the screening primers to confirm the genotyping qPCR results when establishing the colony, in case of any cross-talk between the assays.

#### **Primer sequences**

Primer Name	Primer Sequence (5' > 3')	
Tgm3_WT_F	TTCATGATGGCATCTGTTCTC	
Tgm3_R	TGATGTTCCAAATGAATGAACA	
Tgm3_WT_F2	TTGAAGAAGATTCTGGCATCA	
Tgm3_Mut_F	TTGAAGAAGATTCTGGCATCc	

#### **Reaction setup**

Reagent	μl
DNA (~50-100 ng)	1
10x Buffer	2
MgCl2 (50 mM)	0.6
Platinum Taq (Invitrogen)	0.2
dNTPs (100 mM)	0.2
Primer 1 (10 μM)	0.4
Primer 2 (10 μM)	0.4
ddH20	15.2
Total	20

This technical data sheet and information ("Datasheet") is supplied by Genome Research Limited ("GRL").

Although reasonable care is taken in the preparation of this Datasheet, GRL gives no warranties express or implied for any use of the Datasheet or for the accuracy of the Datasheet. GRL assumes no responsibility or liability for any decisions based upon the Datasheet. Without limiting the foregoing the Datasheet was prepared for mice supplied directly from GRL and where copies of this Datasheet are available from third party repositories or distribution centres ("Third Parties") GRL shall not be liable for any inconsistency between the mouse strain supplied by the Third Party and the Datasheet howsoever arising.

Report Generated on: 12-August-2020



#### **Amplification conditions**

Step	Conditions	Time
1	94°C	5 min
2	94°C	30 sec
3	58°C	30 sec
4	72°C	1:30 sec
5	Go to '2' + 34 cycles	-
6	72°C	5 min
7	12°C	Forever

## **Genotyping by SNP qPCR**

#### Primers for LoA qPCR assay

Gene	Source	Forward Primer Seq.	Reverse Primer Seq.	Probe Primer Seq.
Tgm3	Life	GCCGAAAGACAAGAGTA	CCAACCATGCCAATTCGATTTGTG	[VIC]TCTGGCATCATCTATG[FAM]TTC
	Technologies	TGTTGAAG		TGGCATCCTCTATG

Reactions are performed in a  $10\mu$ l volume using an Applied Biosystems 7900HT Fast Real-Time PCR System or Applied Biosystems Viia7 with DNA prepared using the Sample-to-SNP<sup>TM</sup> kit (Applied Biosystems) from mouse ear biopsies. GTXpress<sup>TM</sup> buffer is also used (Applied Biosystems).

Reagent	μl
2x GTXpressTM buffer	5
40x target assay	0.25
ddH2O	3.75
DNA	1

#### **Amplification conditions**

Step	Conditions	Time
Pre-read	60°C	30 sec
1	95°C	20 sec
2	95°C	10 sec
3	60°C	30 sec
4	Go to '2' + 34	-
Post-read	60°C	30 sec

This technical data sheet and information ("Datasheet") is supplied by Genome Research Limited ("GRL").

Although reasonable care is taken in the preparation of this Datasheet, GRL gives no warranties express or implied for any use of the Datasheet or for the accuracy of the Datasheet. GRL assumes no responsibility or liability for any decisions based upon the Datasheet. Without limiting the foregoing the Datasheet was prepared for mice supplied directly from GRL and where copies of this Datasheet are available from third party repositories or distribution centres ("Third Parties") GRL shall not be liable for any inconsistency between the mouse strain supplied by the Third Party and the Datasheet howsoever arising.

Report Generated on: 12-August-2020



## Links to information and frequently asked questions

MGP mouse phenotype data: http://www.mousephenotype.org

How the "critical" exon is decided: <a href="http://www.i-dcc.org/kb/entry/102/">http://www.i-dcc.org/kb/entry/102/</a>

## **Relevant publications**

White, J.K., Gerdin, A.-K., Karp, N.A., Ryder, E., Buljan, M., Bussell, J.N., Salisbury, J., Clare, S., Ingham, N.J., Podrini, C., et al. (2013). Genome-wide Generation and Systematic Phenotyping of Knockout Mice Reveals New Roles for Many Genes. Cell 154, 452–464.

Mali P, Yang L, Esvelt KM, et al (2013) RNA-guided human genome engineering via Cas9. Science 339:823–6. doi: 10.1126/science.1232033

Jinek M, Chylinski K, Fonfara I, et al (2012) A programmable dual-RNA-guided DNA endonuclease in adaptive bacterial immunity. Science 337:816–21. doi: 10.1126/science.1225829

Cong L, Ran FA, Cox D, et al (2013) Multiplex genome engineering using CRISPR/Cas systems. Science 339:819–23. doi: 10.1126/science.1231143

Singh P, Schimenti JC, Bolcun-Filas E (2014) A Mouse Geneticist's Practical Guide to CRISPR Applications. Genetics genetics.114.169771—. doi: 10.1534/genetics.114.169771

Brandl C, Ortiz O, Röttig B, et al (2015) Creation of targeted genomic deletions using TALEN or CRISPR/Cas nuclease pairs in one-cell mouse embryos. FEBS Open Bio 5:26–35. doi: 10.1016/j.fob.2014.11.009

Zhou J, Wang J, Shen B, et al (2014) Dual sgRNAs facilitate CRISPR/Cas9 mediated mouse genome targeting. FEBS J. doi: 10.1111/febs.12735

Kraft K, Geuer S, Will AJ, et al (2015) Deletions, Inversions, Duplications: Engineering of Structural Variants using CRISPR/Cas in Mice. Cell Rep. doi: 10.1016/j.celrep.2015.01.016

Shen B, Zhang J, Wu H, et al (2013) Generation of gene-modified mice via Cas9/RNA-mediated gene targeting. Cell Res 23:720–3. doi: 10.1038/cr.2013.46

Wang H, Yang H, Shivalila CS, et al (2013) One-step generation of mice carrying mutations in multiple genes by CRISPR/Cas-mediated genome engineering. Cell 153:910–8. doi: 10.1016/j.cell.2013.04.025

Yang H, Wang H, Shivalila CS, et al (2013) One-Step Generation of Mice Carrying Reporter and Conditional Alleles by CRISPR/Cas-Mediated Genome Engineering. Cell 154:1370–1379. doi: 10.1016/j.cell.2013.08.022 This technical data sheet and information ("Datasheet") is supplied by Genome Research Limited ("GRL").

Although reasonable care is taken in the preparation of this Datasheet, GRL gives no warranties express or implied for any use of the Datasheet or for the accuracy of the Datasheet. GRL assumes no responsibility or liability for any decisions based upon the Datasheet. Without limiting the foregoing the Datasheet was prepared for mice supplied directly from GRL and where copies of this Datasheet are available from third party repositories or distribution centres ("Third Parties") GRL shall not be liable for any inconsistency between the mouse strain supplied by the Third Party and the Datasheet howsoever arising.

Report Generated on: 12-August-2020