

Name of Mouse model or mutation:**GRM1-Y792C-EM1-B6N****Description:**

Point mutation model made using CRISPR/Cas9.

Type of mutation:

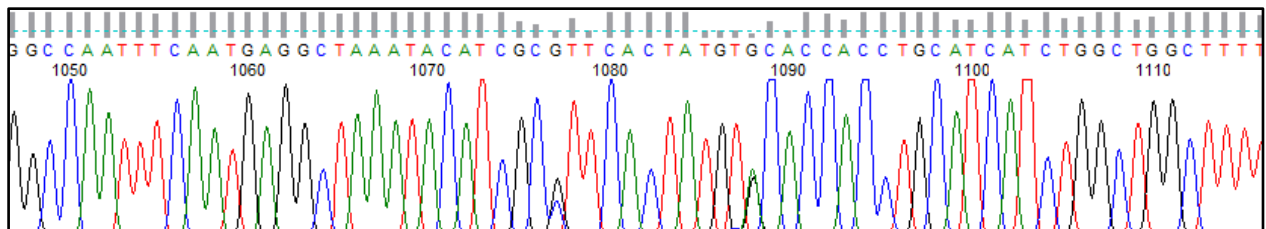
Point mutation: Y792C

Sequence details**WT**

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TGGGCCCAAGTGATCATAGCCTCCATTCTGATTAGTGTACAGCTGACACTAGTGGTGACCTTGATCA
TCATGGAGCCTCCCATGCCATTTTGTCTACCCGAGCATCAAGGAAGTCTATCTTATCTGCAATACC
AGCAACCTGGGTGTAGTGGCACCTGTGGGTTACAACGGACTTCTCATCATGAGCTGTACCTACTATG
CCTTCAAGACCCGCAACGTGCCGGCCAATTTCAATGAGGCTAAATACATCGCCTTCACTATGTACACC
ACCTGCATCATCTGGCTGGCTTTTGTCCATTTACTTTGGGAGCAACTACAAGATTATCACTACCTGC
TTCGCAGTGAGCCTCAGTGTGACGGTGGCCCTGGGCTGCATGTTCACTCCCAAGATGTACATCATT
TTGCCAAACCCGAGAGGAATGTCCGCAGTGCCTTACCACCTCTGATGTAGTGCGCATGCAC
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GRM1-Y792C-EM1-B6N

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TGGGCCCAAGTGATCATAGCCTCCATTCTGATTAGTGTACAGCTGACACTAGTGGTGACCTTGATCA
TCATGGAGCCTCCCATGCCATTTTGTCTACCCGAGCATCAAGGAAGTCTATCTTATCTGCAATACC
AGCAACCTGGGTGTAGTGGCACCTGTGGGTTACAACGGACTTCTCATCATGAGCTGTACCTACTATG
CCTTCAAGACCCGCAACGTGCCGGCCAATTTCAATGAGGCTAAATACATCGCgTTCACTATGTgCACC
ACCTGCATCATCTGGCTGGCTTTTGTCCATTTACTTTGGGAGCAACTACAAGATTATCACTACCTGC
TTCGCAGTGAGCCTCAGTGTGACGGTGGCCCTGGGCTGCATGTTCACTCCCAAGATGTACATCATT
TTGCCAAACCCGAGAGGAATGTCCGCAGTGCCTTACCACCTCTGATGTAGTGCGCATGCAC
```

GRM1-Y792C-EM1-B6N Heterozygous F1 animal sequence trace:

Nucleotide Alignment:

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*      20      *      40      *      60      *      80      *      100     *      120     *      140
Grml_WT   : TGGGCCCAAGTGATCATAGCCTCCATTCTGATTAGTGTACAGCTGACACTAGTGGTGACCTTGATCATCATGGAGCCTCCCATGCCATTTTGTCTACCCGAGCATCAAGGAAGTCTATCTTATCTGCAATACCAGCAA
Grml_Y792C : TGGGCCCAAGTGATCATAGCCTCCATTCTGATTAGTGTACAGCTGACACTAGTGGTGACCTTGATCATCATGGAGCCTCCCATGCCATTTTGTCTACCCGAGCATCAAGGAAGTCTATCTTATCTGCAATACCAGCAA
TGGGCCCAAGTGATCATAGCCTCCATTCTGATTAGTGTACAGCTGACACTAGTGGTGACCTTGATCATCATGGAGCCTCCCATGCCATTTTGTCTACCCGAGCATCAAGGAAGTCTATCTTATCTGCAATACCAGCAA

*      160     *      180     *      200     *      220     *      240     *      260     *      280
Grml_WT   : CCTGGGTGTAGTGGCACCTGTGGGTTACAACGGACTTCTCATCATGAGCTGTACCTACTATGCCTTCAAGACCCGCAACGTGCCGGCCAATTTCAATGAGGCTAAATACATCGC TTCACTATGT CACCACCTGCATCA
Grml_Y792C : CCTGGGTGTAGTGGCACCTGTGGGTTACAACGGACTTCTCATCATGAGCTGTACCTACTATGCCTTCAAGACCCGCAACGTGCCGGCCAATTTCAATGAGGCTAAATACATCGC TTCACTATGT CACCACCTGCATCA
CCTGGGTGTAGTGGCACCTGTGGGTTACAACGGACTTCTCATCATGAGCTGTACCTACTATGCCTTCAAGACCCGCAACGTGCCGGCCAATTTCAATGAGGCTAAATACATCGC TTCACTATGT CACCACCTGCATCA

*      300     *      320     *      340     *      360     *      380     *      400     *      420
Grml_WT   : TCTGGCTGGCTTTTGTTCCCATTTACTTTGGGAGCAACTACAAGATTATCACTACCTGCCTTCGCAGTGAGCCTCAGTGTGACGGTGGCCCTGGGCTGCATGTTCACTCCCAAGATGTACATCATTATTGCCAAACCCGAG
Grml_Y792C : TCTGGCTGGCTTTTGTTCCCATTTACTTTGGGAGCAACTACAAGATTATCACTACCTGCCTTCGCAGTGAGCCTCAGTGTGACGGTGGCCCTGGGCTGCATGTTCACTCCCAAGATGTACATCATTATTGCCAAACCCGAG
TCTGGCTGGCTTTTGTTCCCATTTACTTTGGGAGCAACTACAAGATTATCACTACCTGCCTTCGCAGTGAGCCTCAGTGTGACGGTGGCCCTGGGCTGCATGTTCACTCCCAAGATGTACATCATTATTGCCAAACCCGAG

*      440     *      460
Grml_WT   : AGGAATGTCCGCAGTGCCTTCACCACCTCTGATGTAGTGGCGCATGCAC : 468
Grml_Y792C : AGGAATGTCCGCAGTGCCTTCACCACCTCTGATGTAGTGGCGCATGCAC : 468
AGGAATGTCCGCAGTGCCTTCACCACCTCTGATGTAGTGGCGCATGCAC

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Predicted Protein Alignment:

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*      20      *      40      *      60      *      80      *      100     *      120     *      140
Grml_WT   : WAQVIIASILISVQLTLVVTLIIMEPPMPILSYPSIKEVYLI CN TSNLGVVAPVGYNGLLIMSCTYYAFKTRNVPANFNEAKYIAFTM T TCI IWLAFVPIYFGSNYKIITTCFAVSLSVTVALGCMFPPKMYIIIAKPE
Grml_Y792C : WAQVIIASILISVQLTLVVTLIIMEPPMPILSYPSIKEVYLI CN TSNLGVVAPVGYNGLLIMSCTYYAFKTRNVPANFNEAKYIAFTM T TCI IWLAFVPIYFGSNYKIITTCFAVSLSVTVALGCMFPPKMYIIIAKPE
WAQVIIASILISVQLTLVVTLIIMEPPMPILSYPSIKEVYLI CN TSNLGVVAPVGYNGLLIMSCTYYAFKTRNVPANFNEAKYIAFTM T TCI IWLAFVPIYFGSNYKIITTCFAVSLSVTVALGCMFPPKMYIIIAKPE

*
Grml_WT   : RNVRSAF TTS DVVRMH : 156
Grml_Y792C : RNVRSAF TTS DVVRMH : 156
RNVRSAF TTS DVVRMH

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QC strategy employed at Harwell to check the edited allele:

Genomic DNA was extracted from ear clip biopsies and amplified in a PCR reaction using the following conditions/primer sequences:

Geno_Grm1_F1	TGGGCCCAAGTGATCATAGC
Geno_Grm1_R1	GTGCATGCGCACTACATCAG
Taq Polymerase used	Roche Expand Long Range DNTPack
Annealing Temperature (°C)	63
Elongation time (min)	1
WT product size (bp)	468
Mutant product size (bp)	468
Notes	Sequenced using internal primer Geno_Grm1_F2 (5'-3' CAATACCAGCAACCTGGGTG)

All amplicons were sent for Sanger sequencing to check for integration of the donor oligo sequence at the target site. F1 sequences should be heterozygous unless on Y chromosome.

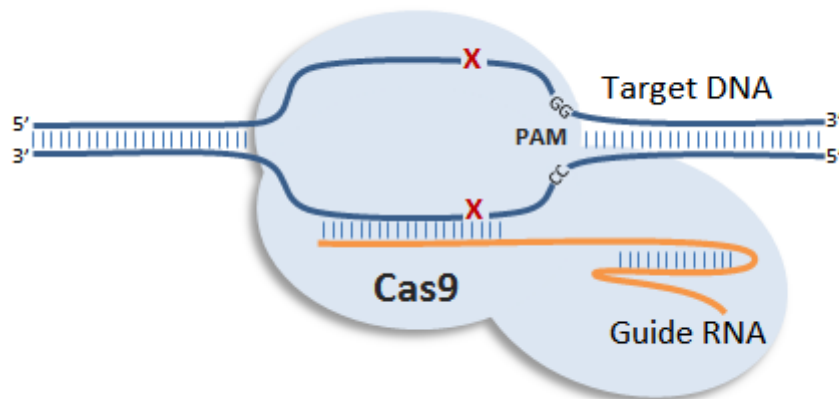
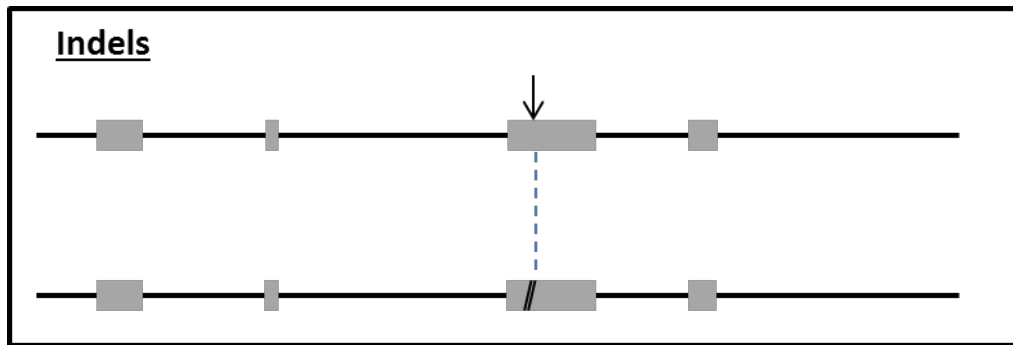
Copy counting of the donor sequence was carried out by ddPCR at the F1 stage to confirm donor oligos were inserted once on target into the genome. The following Taqman assay was used to copy count the donor sequence compared against a VIC-labelled reference assay for Dot1l:

Assay name	Grm1-Y792C-DONOR-WT1
Forward Primer	GCCAATTTCAATGAGGCTAAATACATCG
Reverse Primer	TGCTCCCAAAGTAAATGGGA
Probe	CACCACCTGCATCATCTGGCTGG
Label	FAM-BHQ1

The ddPCR assay is universal and will bind to both WT and mutant Grm1 Y792C sequences. Therefore, WT animals are expected to call at 2 copies and a copy of the desired Point mutation is also expected to call at 2 copies for F1 (HET) animals.

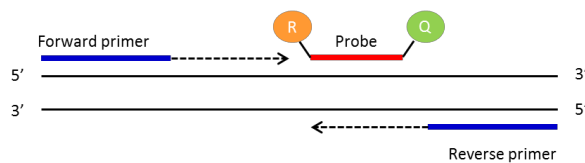
Grm1-Y792C Genotyping Strategy

Animals have been engineered using the CRISPR/Cas9 technology. Most of the knockout alleles generated through this method will be obtained by deletion of a critical exon or by introduction of an indel (insertion/deletion) within the coding sequence of a critical exon (see picture below).

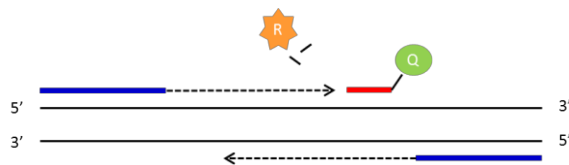


qPCR genotyping strategy

Standard PCR is the amplification of DNA between a pair of primers. Quantitative PCR employs the same principal as standard PCR, although it actually monitors the progress of the DNA synthesis as it occurs. The progress of the reaction is measured by using a Taqman probe. This is a short DNA oligo that is complimentary to part of the DNA sequence between the forward and reverse primers. At the 5' end of the probe there is a fluorescent reporter (R) and at the 3' end a quencher (Q). Whilst they are in close contact with each other there is no fluorescent signal.



As the forward primer is extended the reporter is cleaved from the probe resulting in a fluorescent signal being detected. Once the primer extends enough to release the quencher this signal is blocked. By using probes with different fluorescent signals multiple PCR assays can be multiplexed and run together.



Allele specific primer and probe amplification assay (ASPPAA) PCR

This is a new real-time PCR method (Billard *et al.*, 2012) in which an allele specific primer and an allele specific probe designed specific to the SNPs. The primer is designed such a way that its 3' end ends with a specific SNP. The probe is also designed specific to the SNPs at its 3' end giving a primer probe overlap. A maximum of 3nt overlap between a primer and probe is allowed.

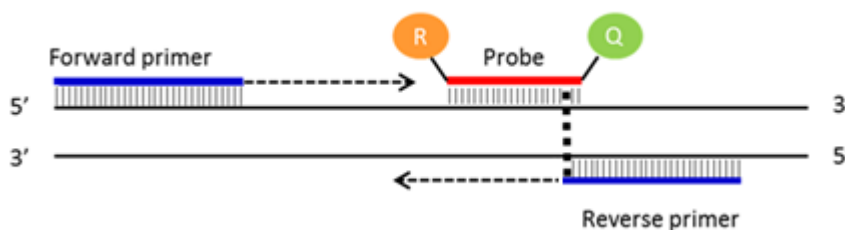
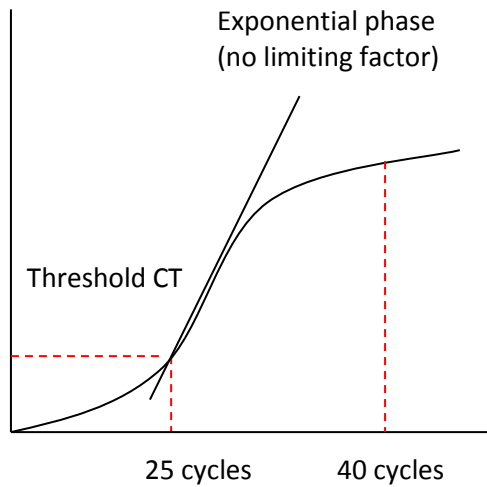


Figure1: Figure showing principle of ASPPAA PCR. The dotted line indicates the position of the SNP.

PCR reaction plot



The number of cycles the PCR takes to reach a set threshold is known as the CT value. This is inversely correlated to the amount of template DNA in the sample.

e.g. CT 25 = 2 x template DNA
CT 26 = 1 x template DNA
CT above 30 = no template represented in the sample

CT value can be used to determine how many copies of a particular allele samples have.

All our qPCR are run in duplicate. A FAM labelled genotyping assay is run in multiplex with a VIC labelled internal control Dot1l.

References:

Billard A., Laval V., Fillinger S., Leroux P., Lachaise H., Beffa R., et al. (2012).
The allele-specific probe and primer amplification assay, a new real-time PCR method for fine quantification of single-nucleotide polymorphisms in pooled DNA.
Appl. Environ. Microbiol. 78 1063–1068. 10.1128/AEM.06957-11



Grm1-Y792C Genotyping Strategy

Samples are genotyped with a WT assay. This is a FAM labelled assay that is designed to detect the critical exon that has been modified. If the animal contains the modified allele the copy number of the WT assay should drop by 1. For autosomal genes that have been targeted this means the following

- WT= 2 copies of the WT assay
- HET = 1 copy of the WT assay
- HOM = 0 copies of the WT assay

Grm1-Y792C CRISPR/Cas9 mutant in which SNPs are as highlighted

WT GCTAAATACATCGCCTTCACTATGTACACCACCTGCATCATCT
 Mutant GCTAAATACATCGCGTTCACTATGTGCACCACCTGCATCATCT

Grm1-Y792C-WT1 assay (FAM labelled probe)

TGGGCCCAAGTGATCATAGCCTCCATTCTGATTAGTGACAGCTGACACTAGTGGTGACCTTGATCATCATGGAG
 CCTCCCATGCCCATTTTGTCTACCCGAGCATCAAGGAAGTCTATCTTATCTGCAATACCAGCAACCTGGGTGTA
 GTGGCACCTGTGGGTTACAACGGACTTCTCATCATGAGCTGTACCTACTATGCCTTCAAGACCCGCAACGTGCCG
GCCAATTTCAATGAGGCTAAATACATCGCCTTCACTATGTACACCACCTGCATCATCTGGCTGGCTTTTGTGCC
 ATTTACTTTGGGAGCAACTACAAGATTATCACTACCTGCTTCGCAGTGAGCCTCAGTGTGACGGTGGCCCTGGGC
 TGCATGTTCACTCCCAAGATGTACATCATTATTGCCAAACCCGAGAGGAATGTCCGCAGTGCCTTACCACCTCT
 GATGTAGTGCGCATGCAC

Grm1-Y792C-WT1 primers and probe

Primer 1 = ACGTGCCGGCCAATTTCAATG
 Primer 2 = GCCAGATGATGCAGGTGGTGT
 Probe = CATCGCCTTCACTATGTACA

Allele specific primer and probes

Dot1l internal control (VIC labelled)

CCCCCTAGTCGTTTTCTGTTAGTAGTTGGCATCCTTATGCTTCATCTTACAGTCGACTTGAGAGCTGCCCCTGA
ATGGTCGTGCTGGGGCAAGGCTTTATTTTCAGGCGTAGCACACATGGTGCCAATGGGACTCTGTAGGATCTGCC

Primer 1 = GCCCCAGCAGACCATT
 Primer 2 = TAGTTGGCATCCTTATGCTTCATC
 Probe = CCAGCTCTCAAGTCG

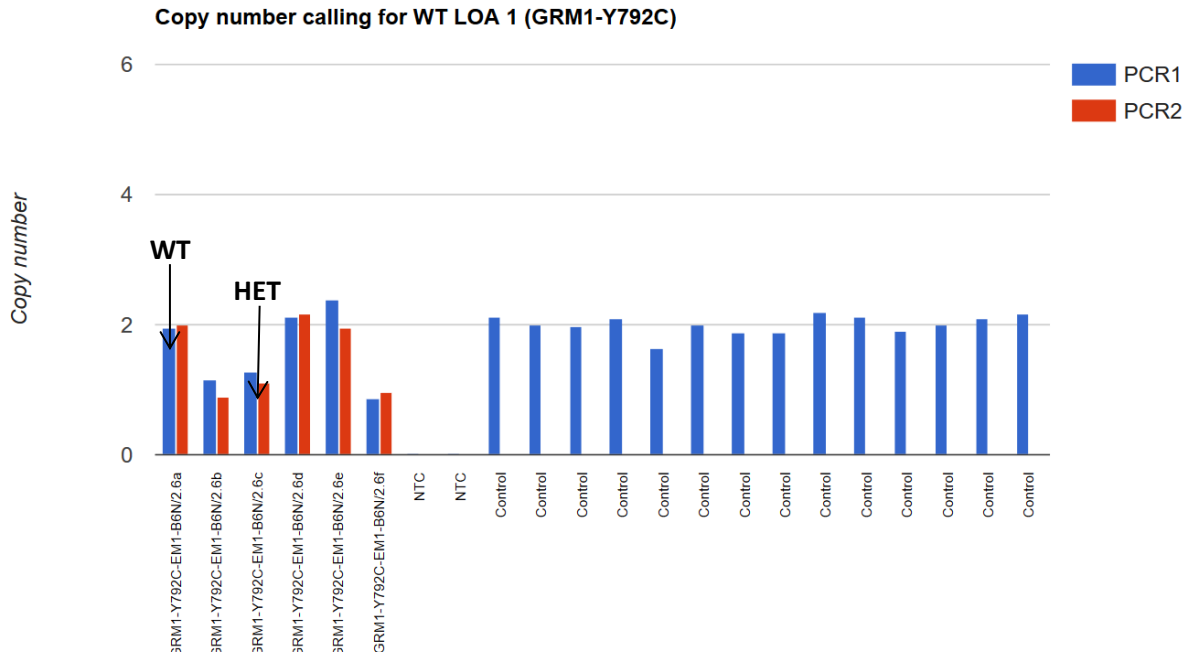
qPCR master mix

ABI GTX Taqman master mix	5µl
Primers Dot1L_2F (20µM)	0.225µl
Primers Dot1L_R (20µM)	0.225µl
Probe DotL_2M (5µM)	0.2µl
FAM Assay (probe 5µM & primers 15µM each)	0.3µl
ddH2O	1.55µl
DNA (1/10 dilution of ABI Sample-to-SNP prep)	2.5µl



Grm1-Y792C copy called result, image showing both replicates and controls

Task 199866 Results



Version No. 1
Date: 10.05.2018
Created/Updated by: Ramakrishna Kurapati
Approved by: Daniel Ford