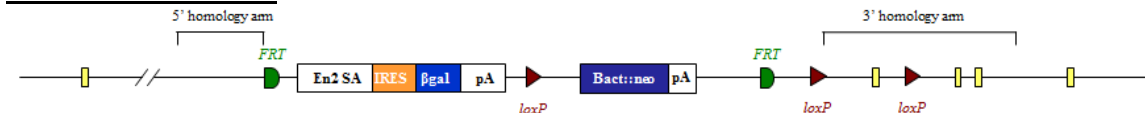


EUCOMM/KOMP-CSD ‘Knockout-First’ Genotyping

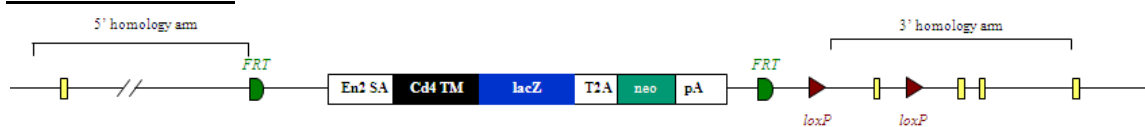
Introduction

The majority of animals produced from the EUCOMM/KOMP-CSD ES cell resource contain the Knockout-First-Reporter Tagged Insertion allele. As well as gene specific elements, the targeting cassette contains many common elements e.g. encoding beta-galactosidase (*lacZ* gene), neomycin-resistance (*neo* gene), FRT and loxP sites. Further details can be found on <http://www.knockoutmouse.org/about/eucomm>. The majority of cassettes will be either promoter driven where the *neo* gene contains its own promotor and is separated from the *lacZ* gene by a 3rd loxP site, or promotorless (PL) where the *neo* and *lacZ* genes are adjacent and there are only 2 loxP sites. These have further implications in our genotyping strategies detailed later.

Promotor driven line



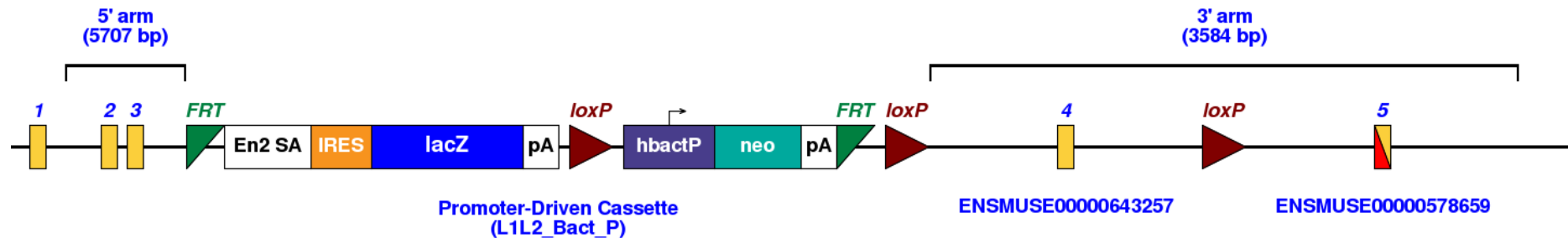
Promotorless line



Animals containing the full Knockout-First mutant allele (Tm1a) above can be crossed to mice expressing Flp or Cre recombinase. The progeny from these mating may contain converted forms of the Tm1a allele known as Tm1b, Tm1c and Tm1d depending on the breeding strategy performed. In order to genotype animals generated from the EUCOMM/KOMP-CSD ‘Knockout-First’ ES cell resource, a combination of mutant assays looking for sequences occurring in the targeting cassette, and WT specific assays will need to be performed. We typically use real time qPCR and copy count the various alleles. This assay design sheet will give an outline of how to genotype these lines by gel based and qPCR based technologies, as well as including details of assays we have tried and tested.



Ahnak-Tm1a

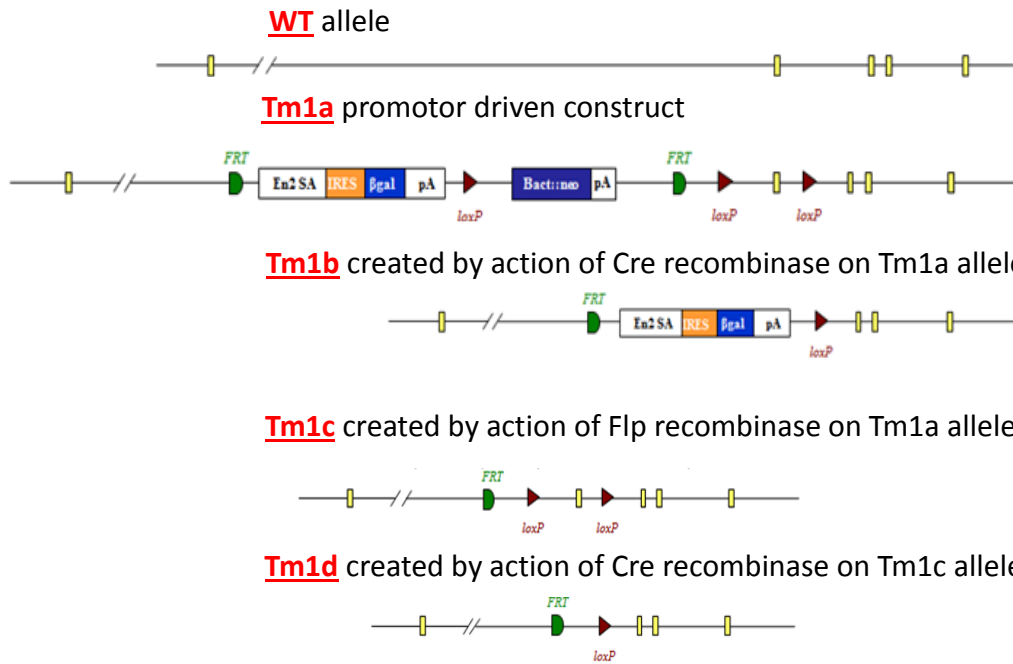


Details for the tm1a allele can be found below

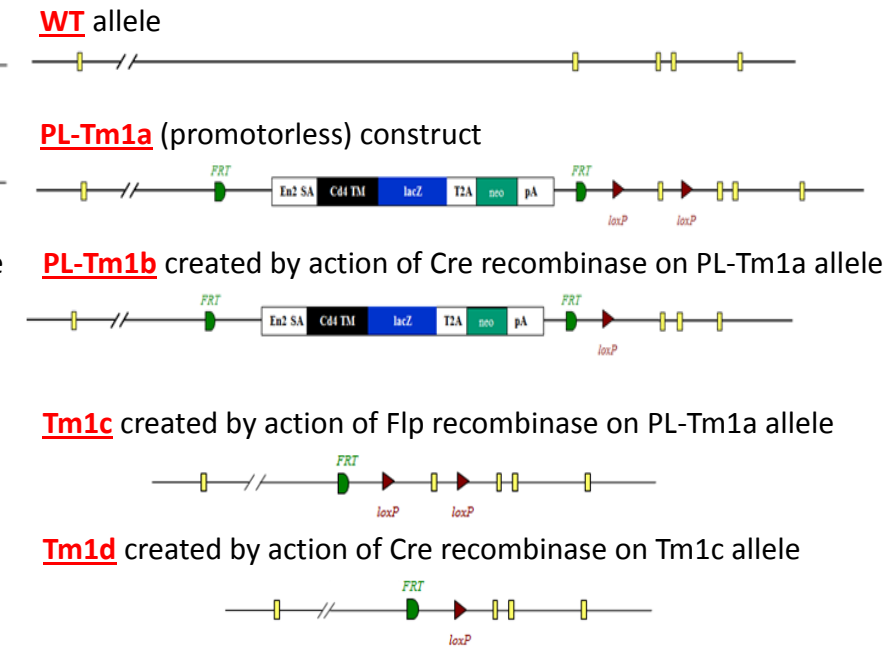
http://www.mousephenotype.org/martsearch_ikmc_project/martsearch/ikmc_project/69475

Types of alleles

Promotor driven lines



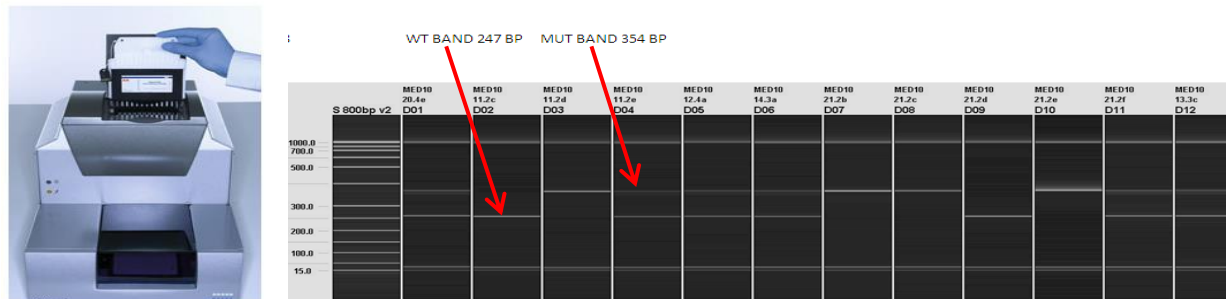
Promotorless lines





Gel based genotyping

The gel based assays are normally run on the Qiagen QIAxcel. This is a capillary based system that provides clearer resolution and is quicker than running standard agarose gels. Different size ladders maybe loaded onto runs depending on the fragment sizes being analysed. Typically samples are run with a 50-800bp size ladder.



PCR is performed using KAPA fast Taq polymerase, although alternatives may be used.

Generic 4 primer PCR mix

KAPA Taq PCR master mix	5µl
Primer 1F	0.5µl
Primer 1R	0.5µl
Primer 2F	0.5µl
Primer 2R	0.5µl
H ₂ O	2.0µl
DNA	1µl

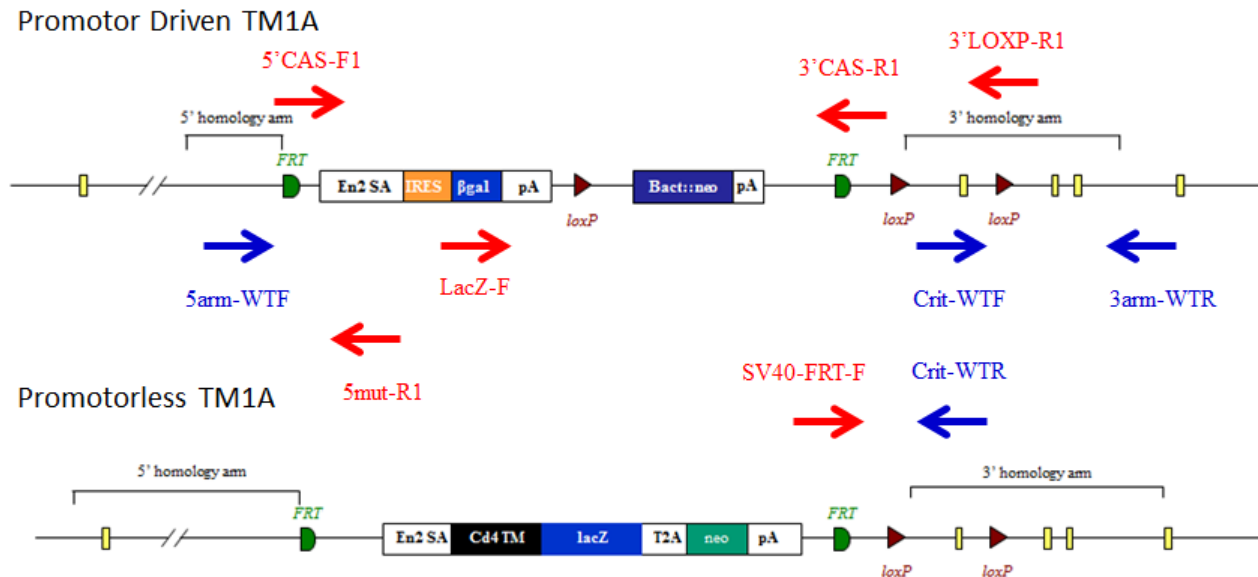
60°C Cycling conditions

60TM30FA

1. 95°C 1min
2. 95°C 10sec
3. **60°C** 10sec
4. 72°C 1sec
5. Go to 2 for 29 cycles
6. 72°C 30sec
7. 16 °C forever

Types of gel based assay

The diagram below shows the location of the various gel based primers.



Mutant primer sequences

5mut-R1	GAACCTCGGAATAGGAACCTCG
LacZ-F	CCAGTTGGTCTGGTGTCA
SV40-FRT-F	CGCGTCGAGAACGTTCTATT
5'CAS-F1	AAGGCGCATAACGATACCAC
3'CAS-R1	CCGCCTACTGCGACTATAGAGA
3'LOXP-R1	ACTGATGGCGAGCTCAGACC

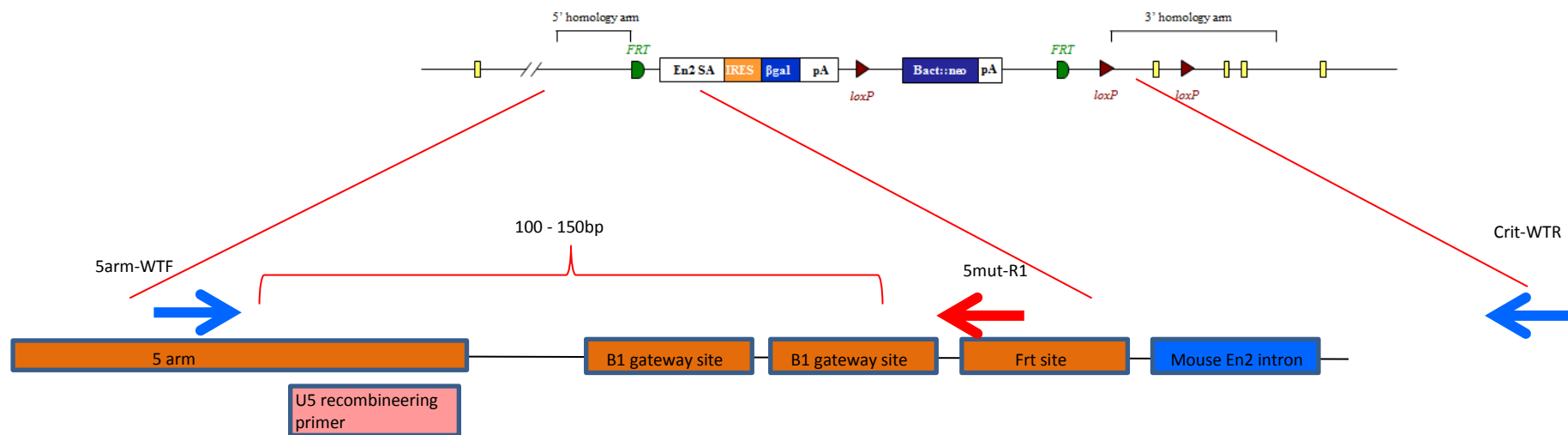
Tm1a gel based genotyping

Tm1a gel based assays are designed using a universal mutant reverse primer that sits in the sequence just after the 5' homology arm

5mut-R1 GAACTTCGGAATAGGAACCTCG

A forward primer is designed to the 5' homology arm and designed to give a mutant specific band, typically between 100-150bp that will only be present if the cassette is present.

A WT reverse primer is designed to the critical region and should give a product of >200bp. If the mutant cassette is present the product between the two primers is too large to PCR under standard conditions.





Ahnak Tm1a gel based genotyping

Ahnak 'Knockout'-First and WT sequences

5'homology arm (last 300bp)

AGAAGGAGGAAGAGACAACCCGGGAGCTGCTGCCAACTGGCAGGGCAGTGGTCCCACGGGCTGACCATTGCCAGAGGGATGATGGAGTCAGGAGGTGATGCAG
AACTCCCCTCGGGCCCGACTGGGTGGTCAAGGAGGGTGAGTCCTCAGGATAATGACTGGGCCTGCAGGGGGGGGGCAGGAGGCAGGGGACAGGCAGAAATGACCAAAT
GACTGTGGAAGGCCTGGGTCTAACAACATAAGGAAGGTCACTACCAAGGCCCTGTCCCTTGCTACTCT

Mutant 5'sequence location of 5mut-R1 (mutant reverse primer)

AAGGCGCATAACGATACCACGATATCAACAAGTTGACAAAAAAGCAGGCTGGCGCCGAAC **CGAAGTTCCTATTCCGAAGTTCTATTCTCTAGAAAGTATAGGAACCTCGAAC**
CCTTCCCCACACCACCTCCACACTGCCAACACTGCCAAC

Critical region

TCAGGGGAGGTAGCCTGTGGAGTACCGACTGTATCTCAGATCTGATTCTGCCTTCACCAATTCTGCAGCTATGTGTAAGTAGTTGGCCTTCCAGCTAGTTTC
CTCCTCTGTGCTTGAATGGCAAACTATGTCTGTGGTGGTCTGAAAGCTAAAGGAAAGTCAGTTTAGTGGCTGTTATGTGGTCTTCTGAGACCTGTGGTCTTACCTG
ATTGTTGTAACCCCTACTTAGGGGACCAGATTGTGGTGCACCATCTACTTGACAAACCTGCAGTCTGGTGGAGGTGACCCAGTTGCTGAATACCATGGGCATCACACTGTTG
CTTGAAGTTGCACCGTAAAGGGGACCGTCCCTGAGCCTGGACAGACCTGGACCCATGAAGTCTCAGTCCCGTAGCTCTGAAGTGGTCTGGTGAGTACCTGGTACCCACCT
TTGTTCAAGAACTCCCCACGATCCTGATGGCTTCTGGCCTACTCCTGAGCCTCCTTATCTCTCTGCATTCTTGACTTCTCTGCTGGTTCTTCTGTTA
TATGTGGCATGTGGCTGGTCCATTCTACAAGTATAAGTGGTCCAGTCAGTCTCTTAAGA

3'homology arm (1st 300bp)

CTGCTTTACTCTGAAATGACCTCACTGGTGTCAACCTGCCTCCCTACATTCCATGAGTGAGAAGATGAACAGCACAATGCTTGGAAAGCGCTTCTTACTCCTAATGGCTTGC
TTACTGAATAAAGCATGTGCCCTAAAGTAAAGTATTACCCAATCACAGTCAGCCATCAGCTGTCACAAATGCTTCCAGTGTGCAAAGGCTCTGGGTGACACCTTAAACAAT
CAAGCTAGTGAGCTGAGAATTGAAAGATGAAGGGCCTGATTGGGATGGGTCTCTGGAACAGGTTCTT



Ahnak Tm1a gel based genotyping

Ahnak 'Knockout'-First and WT sequences

Genomic sequence

Black sequence is missing in the construct due to the insertion of loxP sites and can be used to design break point (BP) qPCR assays

AGAAGGAGGAAGAGACAACCCGGGAGCTGCTGCCA**ACTGGCAGGGCAGTGGTCCCACGGGCTGACCATTGCC**CAGGGATGATGGAGTCAGGAGGTGATGCAG
AACTCCC**CTCGCGCCCGCACTGGGTGGTCAAGGAGGGT**GAGTCCTCAGGATAATGACTGGGCC**CTGCGGGGGGGGGCAGGAGGCAGGGG**GACAGGCAGAAATGACCAAT
GACTGTGG**AAGGCCTGGGTCTAACACATAAGGAAGGT**CAGTACCCAA**AGCCCTG**CCCTGCTACTCT**GTGCTTGCTTG**GAATCTCAGGGAGGTAGTCC**TGTGGAGT**CTTGAA
TACCG**ACTGTATCTTCAGATCTGATTCTGCGCTCTCACCAATTCTGCAGCTATGTG**TA**AAATAGTTGGC**TTCCAGCTAGTTCC**CCTCTGTG**CTGAATGGCAAA**TATATG**
TCTGTGTGG**TGTG**GAAG**CTAAAGGAAAGTCAGTTAGTGGCTGTTATGTGGT**G**CCTTCTTGAGACCTG**GG**GTCTTAC**CTGATTGTT**GACTCCCTACTTAG**GGGACCAG
ATTGTGGGTGCCACCAT**CTACTTTGACAAACCTGCAGTCTGGTGAGGTGACCCAGTTGCTGA**AT**ACCATGGGCATCACACTGTTGGCTGAAGTTGCACCGTAAAGGGGACCGT**C
CCCTGAGCCTGGACAGACCTGGACCCATGAAGTCTTCAGTCCCGTAGCTCTGAA**ACTGTTCTGGTGAGTACCTGGT**AC**CCACCTTGTGAGA**ACTCCCCCACGATC**CTGATGG**
CTTCTGG**CCTACTTCTGAGCCTTCTTATCTCTCTGCATTCTTGACTTCTCTGCTG**TTCTGTT**ATATGTGGCATG**TGG**CTGGTGGT**CCATTCTAC
AAGTATAGTGTCCCTCAGTCTTAAGACTATGCC**ATGGTGAGGGAAACA**ACTCAAGCAGTGGGG**CTTAGGTGCCAAGAGGAAGGTTGGCTG**TTACTCTGAA**ATGACCTC**
TGGTGT**CAACCTGCCTCC**TACATTCCATGAGTGAGAAGATGACAGCACA**ATGCTTG**GGGAAGCG**CTTACTCTTA**ATGG**CTTGCTTA**CTGAATAAAGCATG**TGCCCT**
AAGTGAAAGTATTACCCAA**ATCACAGTCAAGCCATCAGCTGT**CACAA**ATGCTTCCAGTGTGCAAAGGCTCTGGT**GACAC**CTTAAACA**ATCAAGCTAGTGAGCTGAGAATTGAAA
GATGAAGGGC**CTGATTGGATGGGTCTCTGG**AA**CAGGTTCT**

5' homology arm and cassette sequence

AGAAGGAGGAAGAGACAACCCGGGAGCTGCTGCCA**ACTGGCAGGGCAGTGGTCCCACGGGCTGACCATTGCC**CAGGGATGATGGAGTCAGGAGGTGATGCAG
AACTCCC**CTCGCGCCCGCACTGGGTGGTCAAGGAGGGT**GAGTCCTCAGGATAATGACTGGGCC**CTGCGGGGGGGG**CAGGAGGCAGGGG**GACAGGCAGAAATGACCA**AT
GACTGTGG**AAGGCCTGGGTCTAACACATAAGGAAGGT**CAGTACCCAA**AGCCCTG**CCCTGCTACTCT**AAGGCGCATAACG**ATAC**ACGATATCACAA**AGTTGTA**AAAAAAAGCA**
GGCTGGCGCCGGAA**ACCGAAGTTCC**ATT**CCGAAGTTC**CTATTCT**CTAGAA**GTATAGGA**ACTTCGAAC**CC**TTCCCACACCAC**CTGCC**AAACACTGCCA**AC



Tm1a gel based genotyping

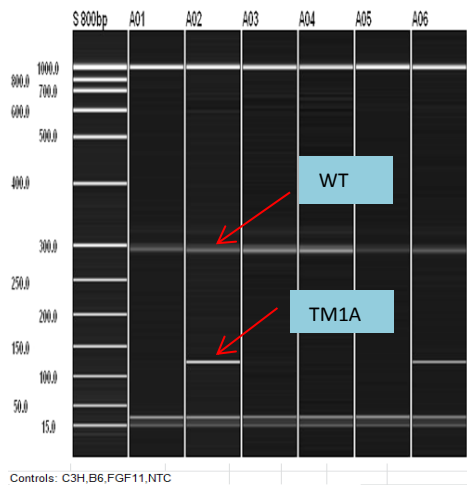
Ahnak tm1a gel based primers

Ahnak-5arm-WTF	AAGGCCTGGTCTAACACA
Ahnak-Crit-WTR	CCCACAGGTCTCAAGAAAGC
5mut-R1	GAACCTCGGAATAGGAACCTCG
WT band	300
Mutant band	145

PCR mix

KAPA Taq PCR master mix	5µl
5arm-WTF	0.5µl
Crit-WTR	0.5µl
5mut-R1	0.5µl
H ₂ O	2.5µl
DNA	1µl

Example of a generic TM1A assay



Cycling conditions

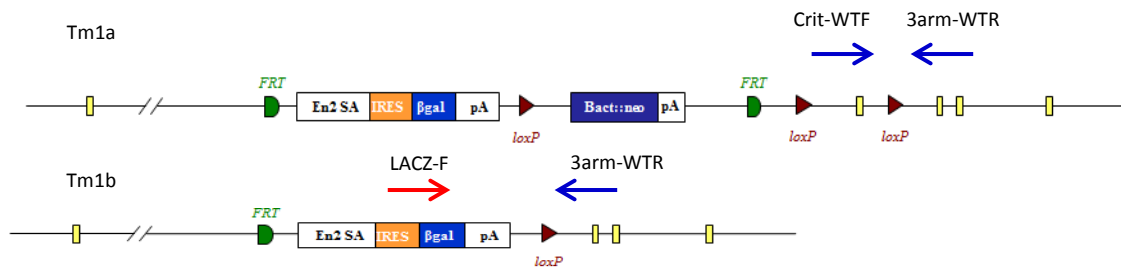
60TM30FA

1. 95°C 1min.
2. 95°C 10sec.
3. **60°C** 10sec.
4. 72°C 1sec.
5. Go to 2 for 29 cycles
6. 72°C 30sec.
7. 16 °C for ever
8. end

Tm1b gel based genotyping

Promotor driven genotyping

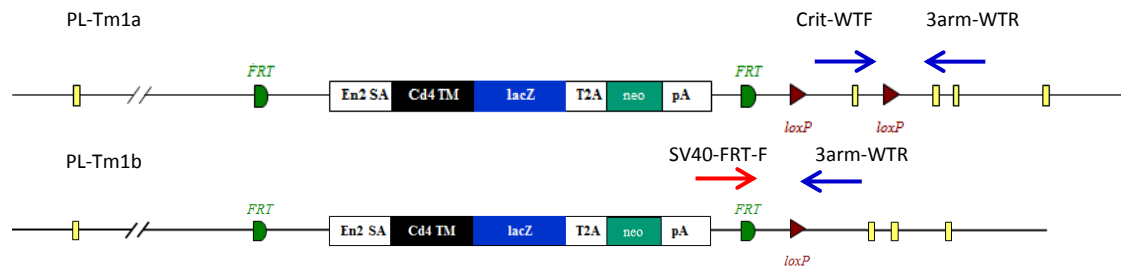
Tm1b alleles are produced by exposing tm1a alleles to cre recombinase which causes the deletion of Neo and the critical region.



If the deletion of the critical region has not occurred then a product is produced between the -Crit-WTF and the -3arm-WTR. Complete conversion by Cre will mean a product is produced from the LacZ-F and -3arm-WTR as the two primers now lie close enough to generate a PCR product under standard conditions. Run both the tm1a genotyping protocol to detect WT and tm1a alleles and the tm1b protocol to detect converted alleles.

Promotorless genotyping

PL-Tm1b (promotorless tm1b) mice are produced in a similar fashion but only the deletion of the critical region occurs and Neo is still present.

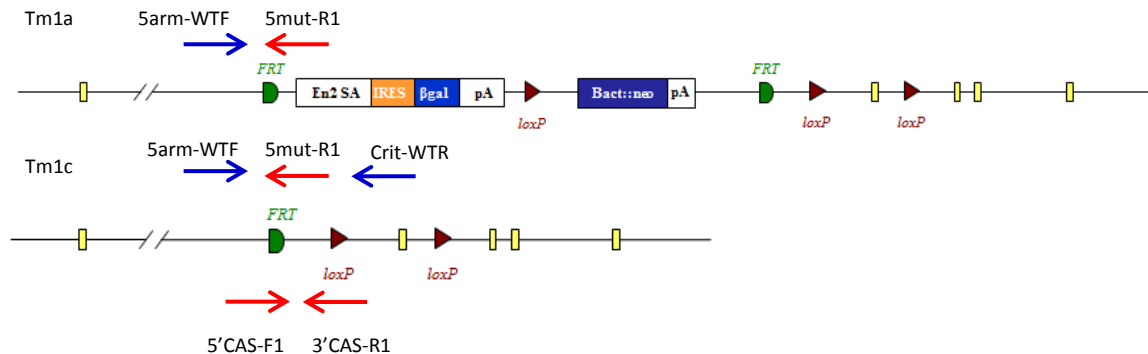


If the deletion of the critical region has not occurred then a product is produced between the Crit-WTF and the 3arm-WTR. Complete conversion by Cre will mean a product is produced from the SV40-FRT-F and 3arm-WTR as the two primers now lie close enough to generate a PCR product under standard conditions. This will not work for promotor driven lines as the presence of the 5' loxP site between the Neo and lacZ sequence causes this region to be removed by cre deletion. Run both the tm1a genotyping protocol to detect WT and tm1a alleles and the tm1b protocol to detect converted alleles.

Tm1c and Tm1d gel based genotyping

Tm1c genotyping

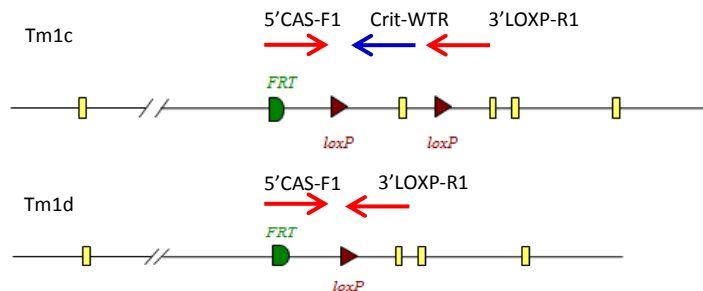
Tm1c alleles are produced by exposing tm1a alleles to flp recombinase which causes the deletion of Neo and LacZ sequences that are between the FRT sites.



The protocol for tm1a genotyping needs to be run. Both tm1a and tm1c alleles will produce a mutant specific product from -5arm-WTF and 5mut-R1. WT and tm1c alleles will produce PCR products from -5arm-WTF and Crit-WTR although the tm1c product will be around 200bp bigger due to the presence of the FRT-loxP part of the cassette. In addition to this a generic tm1c assay can be used. This uses the 5'CAS-F1 and 3'CAS-R1 primers run with an internal control. The tm1c allele produces a 218bp product.

Tm1d genotyping

Tm1d mice are produced by crossing tm1c animals with mice expressing cre recombinase. This causes the deletion of the critical region.



The tm1d allele produces a 174bp product from the 5'CAS-F1 and 3'LOXP-R1 primers. A tm1c allele will produce a larger product which includes the critical region sequence. If the critical region is very large then a gene specific primer lying in the critical region may be needed to detect it (Crit-WTR).

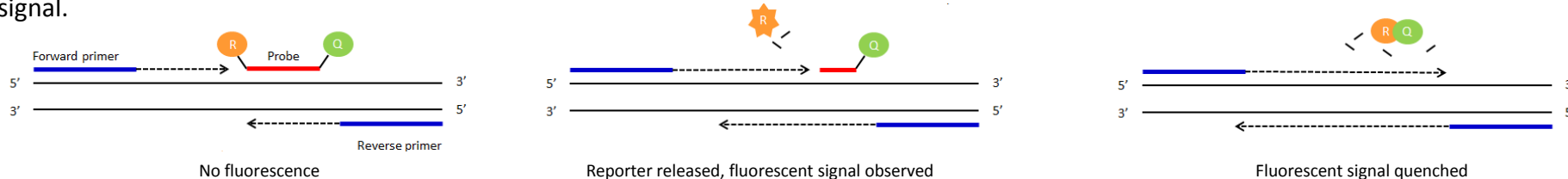
Gel based genotyping strategy summary

Primers to use	5'CAS-F1	5mut-R1	LacZ-F	SV40-FRT-F	3'CAS-R1	3'LOXP-R1	5arm-WTF	Crit-WTF	Crit-WTR	3arm-WTR	Comments
Tm1a + PL		✓					✓		✓		
Tm1b			✓					✓		✓	Run tm1a protocol to identify WT alleles
PL-Tm1b				✓				✓		✓	Run tm1a protocol to identify WT alleles
Tm1c	✓	✓			✓		✓		✓		Blue =separate reaction
Tm1d	✓					✓			✓		Run tm1a protocol to identify WT alleles

- WT Product between 5arm-WTF and Crit-WTR (product too big in tm1a).
- Tm1a Product between 5arm-WTF and 5mut-R1.
- Tm1b Product between LacZ-F and 3arm-WTR (product too big in tm1a but PCR product from Crit-WTF and 3arm-WTR is made).
- Tm1b-PL Product between SV40-FRT-F and 3arm-WTR (product too big in tm1a but PCR product from Crit-WTF and 3arm-WTR is made).
- Tm1c 218bp product between 5'CAS-F1 and 3'CAS-R1 made. Can also run tm1a protocol for a gene specific assay.
- Tm1d 174bp product produced from 5'CAS-F1 and 3'LOXP-R1. Larger product may be produced by tm1c.

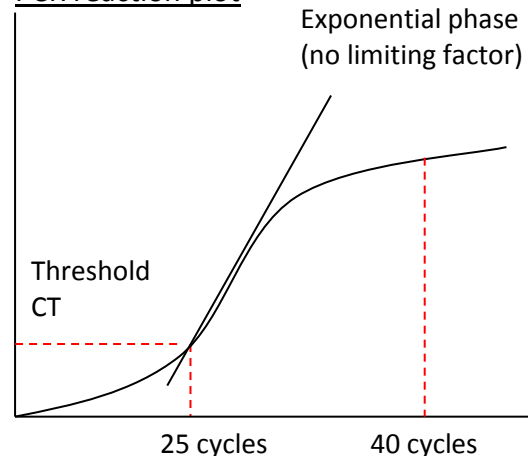
qPCR based genotyping

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.

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

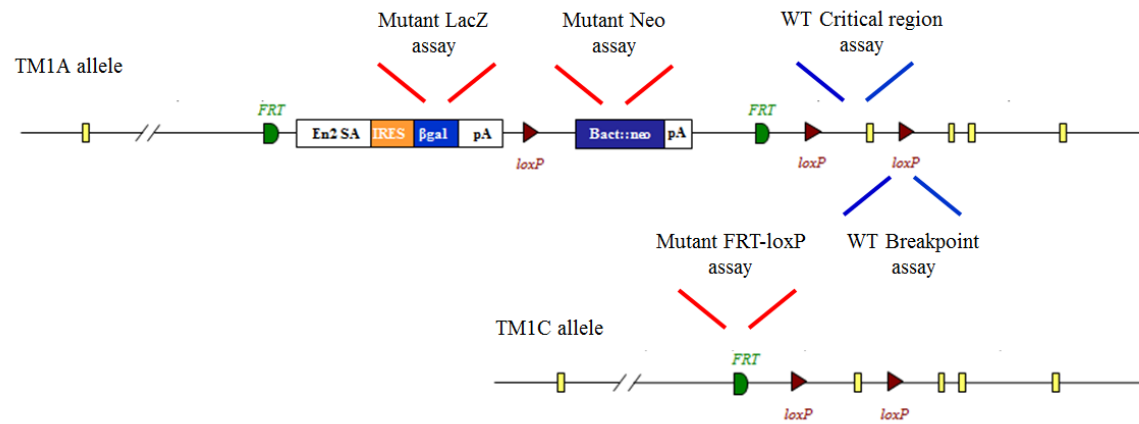
CT above 30 = no template represented in the sample

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

qPCR Assay Types

All qPCR assays are FAM labelled and run in duplex with a VIC labelled internal control, Dot1l. There are 3 mutant qPCR assays based around the targeting cassette. They are as follows:

- : Neomycin assay Designed around the sequence of the neomycin resistance cassette.
- : LacZ assay Designed around the sequence of the LacZ reporter.
- : FRT-loxP Designed to detect the recombined sequence of the FRT and loxP area created when the Tm1a allele is Flp converted to the tm1c. The forward primer is based on the sequence of the 5' region upstream of the 5' FRT site which means this assay will only work for Tm1c/d.



There are 2 WT qPCR assays based on WT sequence that may be lost in Tm1a/b/c/d alleles. They are as follows:

- : BP-LOA Break Point loss of allele assay. This is based on WT sequence that is lost around the loxP sites of the synthetic cassette.
- : CR-LOA Critical Region loss of allele assay. This is based on the critical region sequence that is flanked by the two loxP sites.



qPCR Mutant assay sequences

NEO assay (FAM labelled probe)

ATTGAACAAGATGGATTGCACGCAGGTTCTCCGGCCGTTGGTGGAGAGGCATT CGGC TATGAC TGGGCACAACAGACAATCGGCTGCTCTGATGCCGCCGTGTC CCGCTGTC
AGCGCAGGGCGCCGTTCTTTGTCAAGACCGACCTGTCCGGTGCCTGAATGAAC TG CAGGACGAGGCAGCGCGGTATCGTGGCTGCCACGACGGCGTCCCTGCGCAG
CTGTGCTGACGTTGTCACTGAAGCGGAAGGGACTGGCTGCTATTGGCGAAGTGCCGGGCAGGATCTCCTGTCATCTCACCTGCTCCTGCCGAGAAAGTATCCATCATGGCT

Primer 1 = GGTGGAGAGGCTATT CGGC

Primer 2 = GAACACGGCGGCATCAG

Probe = TGGGCACAACAGACAATCGGCTG

LacZ assay (FAM labelled probe)

CGATAAAAACA ACTGCTGACGCCGCTGCGCGATCAGTTCACCGTGCACCGCTGGATAACGACATTGGCGTAAGTGAAGCGACCCGATTGACCTAACGCCCTGGGTGCAACGCT
GGAAGGC GGCGGGCCATT ACCAGGCCGAAGCAGCAGCGTTGTCAGTGACGGCAGATACTTGCTGATGCGGTGCTGATTACGACCGCTCACCGTGGCAGCATCAGGGAAAACC
TTATT TATCAGCCGGAAAACCTACC GGATTGATGGTAGTGGTCAAATGGCGATTACCGTTGATGTGAAGTGGCGAGCGATAACCCGATCCGGCGGGATTGCCCTGA ACTGCCA
GCTGGCGCAGGTAGCAGAGCGGGTAAACTGGCTCGGATTAGGGCGCAAGAAA ACTATCCGACCGCCTACTGCCCTGTTGACCGCTGGGATCTGCCATTGTCAGACATGT

Primer 1 = CTCGCCACTTCAACATCAAC

Primer 2 = TTATCAGCCGGAAAACCTACC

Probe = TCGCCATTGACCACTACC ATCAATCC

FRT-LOXP-mut2 assay (FAM labelled probe)

AAGGCGCATAACGATACCA CGATA TCAACAAGTTGTACAAAAAAGCAGGCTGGCGCCGGAACCGAAGTT CCTAT TCCGAAGTT CCTATTCTCTAGA AAGTATAGGA ACTT C <GTC
GAGATAACTT>CGTATAGCA TACATTATACGAAG

Primer 1 = AGGCGCATAACGATACCA CGATA

Primer 2 = TGCTATACGAAGTTATCTGACGA

Probe = TCCGAAGTT CCTATTCTCTAGA



qPCR WT assay sequences

Example of a BP-LOA-wt1 assay (FAM labelled probe)

Blue = Sequence in homology arm Black = breakpoint Red = critical region

TACAACCTGTATGAGGCCATTCCCTCCTGGAAGAACACATATTGGATTGATTTAACTCGGATTTGCAGGCTGCTTGCACTTGAAACTGTTAAATATATTAC<CCAA
ACCAATCTTGGCAAGTAGCAAGCCTTTAGAGACACATCGGTGAGTCGGTGGTTTGAAACACTC>CGTCATCTCATTATTTGGGTTGTTAGATCAGAAACTTGCAA
TGAACTTTCGTCATGGCTGGTCAAGTGTCAAACCATGCTCTGAGATAAAGTTCTAACGCCTGAGGTTGCCCTCCAGACTTGTCAGCTGACGAGGGTAGAGTTCTGCTAC

Primer 1 = GGCTGCTTGCACTTGAAAC Primer 2 = CCAGCCATGCACGAAAAGTTC Probe = AGACACATCGGTGAGTCGGTGGTT

Ahnak CR-LOA-wt1 assay (FAM labelled probe)

GTTGACTCCCTACTTAGGGGACCAGATTGTGGGTGCCACCATCTACTTTGACAACCTGCAGTCTGGTGAGGTGACCCAGTTGCTGAATACCATGGGCATCACACTGTTGGCTT
GAAGTTGCACCGTAAAGGGGACCGTTCCCTGAGCCTGGACAGACCTGGACCCATGAAGTCTCAGTTCCCGTAGCTCTGAAGTGGTTCTGGTGAGTACCTGGTGACCCACCTTG
TTCAGAACTCCCCACGATCCTGATGGCTCTGGCTACTCCTGAGCCTCCTATCTCTCTTCT

Primer 1 = TGGGTGCCACCATCTACTTTG Primer 2 = GGTGCAACTTCAAGCCAACAG Probe = ACAACCTGCAGTCTGGTGAGGTGA

All qPCR assays are run in duplex with a VIC labelled internal control, Dot1l

Dot1l internal control (VIC labelled internal control)

AACCAGTGGGCAGTGGATGTGAAGGGCAGGAAGGCTCATAGGGTACTGGCAACCCAGGAAGCCGGAGTGCTCGTCTCTGTTCTTTCCCTCTAGCGTTCT
GTAGTAGTGGCATCCTATGCTTCATCTACAGTCAACTGAGAGCTGCCCTGAATGGCTGCTGGGCAAGGCTTATTCAGGCAGCACACATGGTGGCCAATGGGAC
TCTGTAGGATCTGCCACACCCATCAGGTGTGCAGGGAGACAGAGCTGAGTCAGGCTCAGCTGGGAATATGTTGAGTCACCACCTGTAGGGTGGTGTGCATAGAAC

Primer 1 = GCCCCAGCACGACCATT Primer 2 = TAGTTGGCATCCTTATGCTTCATC Probe = CCAGCTCTCAAGTCG



qPCR genotyping set up

qPCR master mix

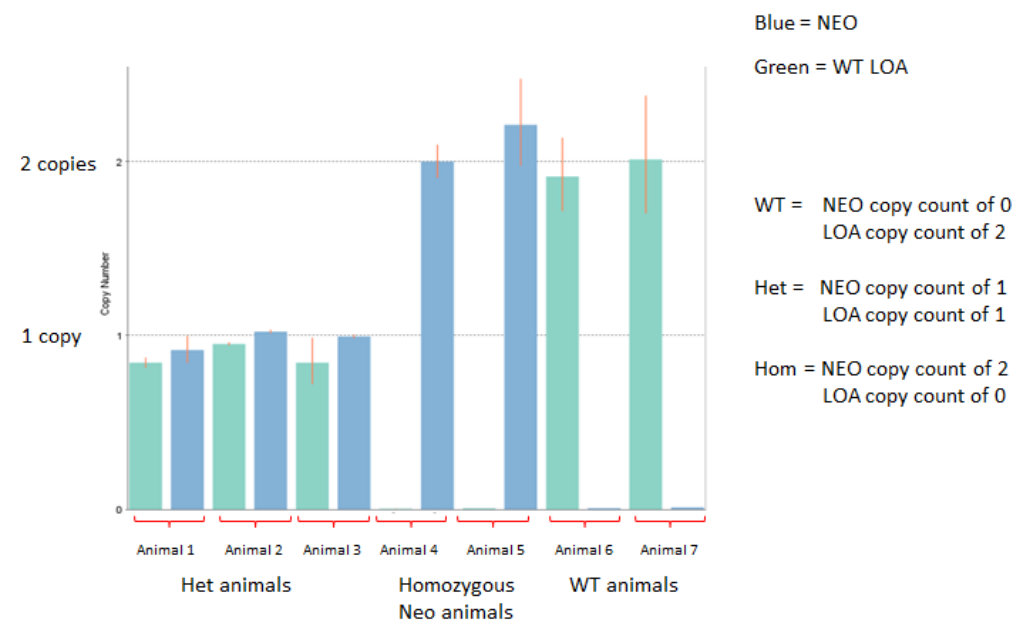
ABI GTX Taqman master mix	5µl
Primers Dot1L_2F (20µM)	0.225µl
Primers Dot1L_R (20µM)	0.225µl
Probe DotL_2M (5µM)	0.2µl
FAM Assay (probe 5µM & primers 15µM each)	0.3µl
Water	1.55µl

Generic example of a NEO + BP-LOA copy called result

ALIQUOT 7.5µL

DNA (1/10 dilution of ABI Sample-to-SNP prep)	2.5µl
--	-------

The results to the right have been run on an ABI 7500 and analysed using the ABI software CopyCaller. Each sample was run with a technical duplicate and for each assay, Neo and the BP-LOA, 7 controls of known copy number and a no template control, all in duplicate were run so the software can accurately copy count.



qPCR based genotyping strategy summary

Assay to use and copy number expected per allele					
	NEO	LACZ	FRT-loxP	BP-LOA	CR-LOA
WT	v 0			v 1	
Tm1a + PL-Tm1a	v 1			v 0	
Tm1b	v 0	v 1			v 0
PL-Tm1b	v 1	v 1			v 0
Tm1c		v 0	v 1	v 0	
Tm1d		v 0	v 1	v 0	v 0

Expected copy number per animal					
	NEO	LACZ	FRT-loxP	BP-LOA	CR-LOA
WT	0	0	0	2	2
Tm1a-HET	1	1	0	1	2
Tm1a-HOM	2	2	0	0	2
Tm1b-HET	0	1	0	1	1
Tm1b-HOM	0	2	0	0	0
PL-Tm1b-HET	1	1	0	1	1
PL-Tm1b-HOM	2	2	0	0	0
Tm1c-HET	0	0	1	1	2
Tm1c-HOM	0	0	2	0	2
Tm1d-HET	0	0	1	1	1
Tm1d-HOM	0	0	2	0	0

Assays that are normally run to detect allele

Assays not normally run

Please note copy numbers are based on what is expected for autosomal genes



Allele Conversions

The action of either Flp or Cre recombinase can be used to convert the KO 1st alleles described in this document. Both gel based and qPCR genotyping methods may be employed to detect animals carrying the DNA sequences coding for these proteins.

Cre Genotyping

Cre gel based assays are run using primers specific to DNA sequences encoding Cre recombinase as well as a set of primers detecting a house keeping gene Slc40a1. These are run using the same cycling conditions as described in page 4.

ActinB-Cre gel based primers

Cre_F1	GCGGTCTGGCAGTAAAAACTATC
Cre_R1	GTGAAACAGCATTGCTGTCACTT
Slc40a1_sh_F (control)	CCTTGTAACCTCCTCTGTGTC
Slc40a1_sh_R (control)	CTGAAGTCTTCATGATAACTGCATT

Fragment sequences

Slc40a1 147bp

CCTTGTAACCTCCTCTGTGCTTATTTAGCCTAACAGCAAAGACTAAAGATGGATCTTATCTGGAGAATAGTTCTATTAGTCCAGCAAACTTCTAGTAAATATGTTAG
ATGAAAATGCAGTTATCATGAAAGACTTCAG

Cre1 102bp

GCGGTCTGGCAGTAAAAACTATCCAGCAACATTGGGCCAGCTAACATGCTTCATCGTCGGCCGGCTGCCACGACCAAGTGACAGCAATGCTGTTCAC



Allele Conversions

Cre Genotyping

Cre qPCR assay (FAM labelled probe)

ATGTCCAATTTACTGACCGTACACCAAATTGCCTGCATTACCGGTCGATGCAACGAGTGATGAGGTT CGCAAGAACCTGATGGACATG TTCAGGGATGCCAGGCGTT TCTGA
GCATAACCTGGAAAATGC TTCTGTCGTTGCCGGT CGTGGGCGGCATGGT GCAAGTTGAATAACCGGAAATGGTTCCCGCAGAACCTGAAGATGTTCGCGATTATCTTCTATATC
TTCAGGCGCGGGTCTGGCAGTAAAAACTATCCAGCAACATTGGGCCAGCTAACATGCTTCATCGTCGGTCCGGCTGCCACGACCAAGTGACAGCAATGCTGTTCACTGGTT

Primer 1 = CGCAAGAACCTGATGGACATG

Primer 2 = ACCGGCAAACGGACAGAA

Probe = TTCAGGGATGCCAGGCGTT

qPCR set up for the Cre assay is exactly the same as previously, where the assay is run in duplex with a VIC labelled Dot1l internal control.



Allele Conversions

Flpe Genotyping

Flpe gel based assays are run using primers specific to DNA sequences encoding Flpe recombinase as well as a set of primers detecting a house keeping gene Slc40a1. These are run using the same cycling conditions as described in page 4.

Flpe gel based primers

ICS_Flpe_F	TCTTAGCGCAAGGGTAGGATCG
ICS_Flpe_R	GTCCTGCCACGGCAGAACG
Slc40a1_sh_F (control)	CCTTGTAACCCCTCTGTGTC
Slc40a1_sh_R (control)	CTGAAGTCTTCATGATAACTGCATT

Fragment sequences

Slc40a1 147bp

CCTTGTAACCCCTCTGTGCTTTATTTAGCCTAACAGCAAAGACTAAAGATGGATCTTATCTGGAGAAtagttctattnagtcCAGCAAACCTTCTAGTAAATATGTTAG
ATGAAAATGCAGTTATCATGAAAGACTTCAG

ICS-Flpe1 332bp

CCTTCTTAGCGCAAGGGTAGGATCGATCCACTTGTATATTGGATGAATTGGAGGAATTCTGAACCAGTCCTAAACGAGTAAATAGGACCGGAATTCTCAAGCAACAA
ACAGGAATACCAATTAAAAGATAACTTAGTCAGATCGTACAACAAGGCTTGAAGAAAAATGCGCCTTATCCAATCTTGCTATAAGAATGGCCAAAATCTCACATTGAA
GACATTGATGACCTCATTCTGTCAATGAAGGGCTAACGGAGTTGACTAATGTTGGAAATTGGAGCGATAAGCGTGCTTCTGCCGTGGCCAGGAC



Allele Conversions

F1pe Genotyping

F1pe qPCR assay (FAM labelled probe)

GGTAGGATCGATCCACTTGTATTTGGATGAATTGGAGGAATTCTGAACCAGTCCTAAAACGAGTAAATAGGACCGCAATTCTCAAGCAACAAACAGGAATACCAATTATT
AAAAGATAACTTAGTCAGATCGTACAACAAGGCTTGAGAAAAATGCCTTATCCAATCTTGTCTAAAGAATGGCCAAAATCTCACATGGAAGACATTGATGACCTCAT
TTCTGTCAATGAAGGGCTAACGGAGTTGACTATGTTGTGGAAATTGGAGCGATAAGCGTGCTTCTGCCGTGGCCAGGACAACGTATACTCATCAGATAACAGCAATACCTGAT
CACTACTTCGCACTAGTTCTCGGTACTATGCATATGATCCAATATCAAAGGAAATGATAGCATTGAAGGATGAGACTAATCCAATTGAGGAGTGGCAGCATAAGACAGCTAAA
GGTAGTGTGAAGGAAGCATACGATACCCGATGGAATGGATAATATCACAGGAGGTACTAGACTACCTTCATCCTACATAAATAGACGCATATAAT

Primer 1 = GTGGGAAATTGGAGCGATAAGC

Primer 2 = ACCGAGAAACTAGTGCAGTAG

Probe = CTTCTGCCGTGGCCAGGACAAC

qPCR set up for the F1pe assay is exactly the same as previously, where the assay is run in duplex with a VIC labelled Dot1l internal control.