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This protocol has been prepared by Claudia Caradec, Engineer

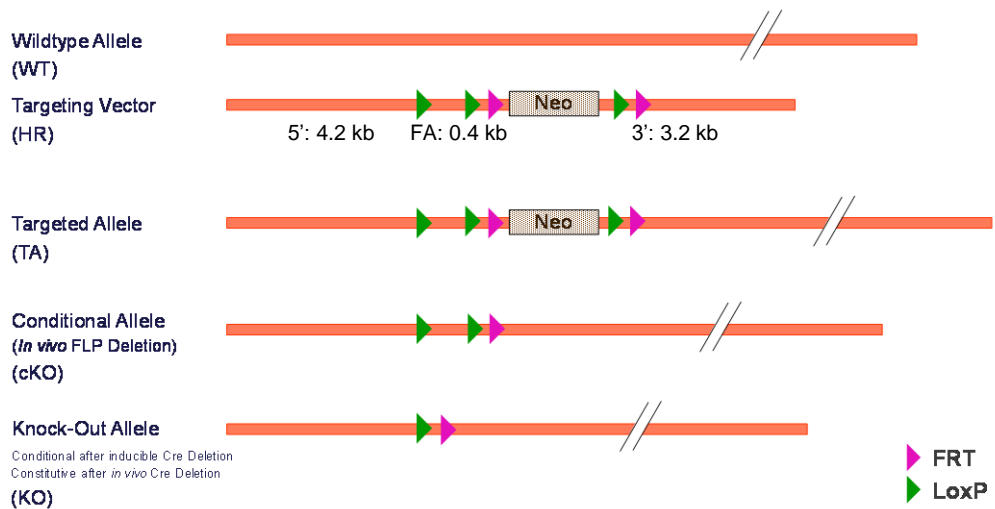
This protocol has been validated by Sylvie Jacquot, Ph.D., Project Manager

1. Schematic representation of the locus

1.1. Overview



Overview Targeting Strategy



Legend:

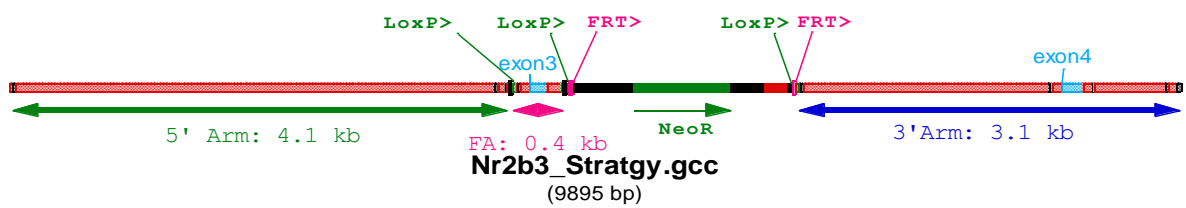
5': 5' homology arm; FA: floxed fragment; 3': 3' homology arm

This schematic representation is not on scale

1.2. Strategy chosen: flox of exon 3

Nr2b3 gene (also named RXRgamma) is a member of the nuclear receptor family. Additional information on this gene can be accessed at <http://www.informatics.jax.org/javawi2/servlet/WIFetch?page=markerDetail&key=13296>

Strategy used to generate the conditional knock out model

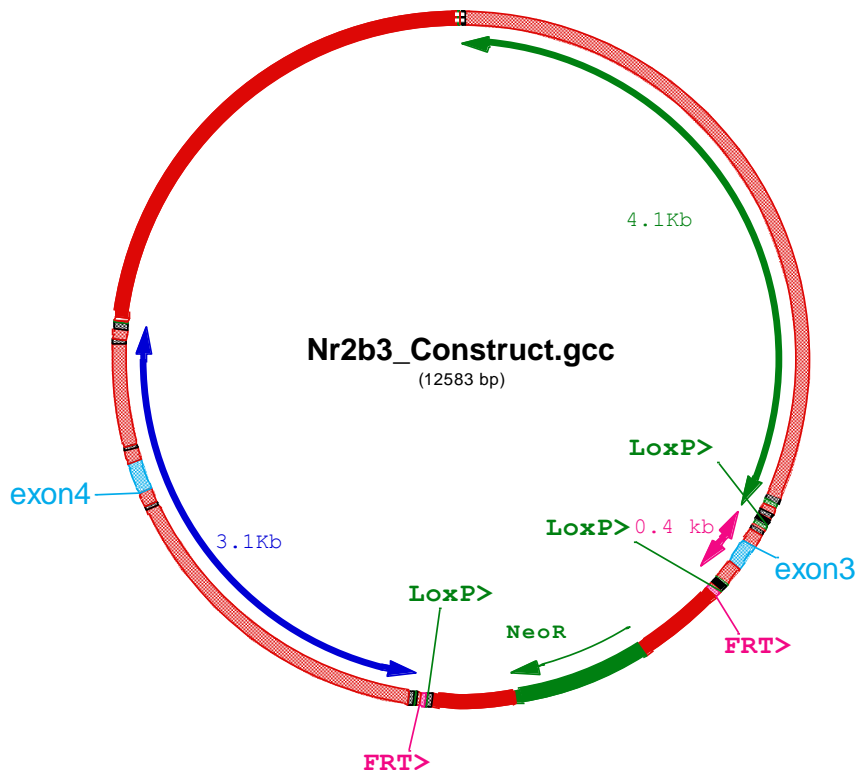


2. Construct used for homologous recombination in ES cells: Nr2b3 project

2.1. Legend

loxP sites are indicated in green ; FRT sites are indicated in purple; *Mus musculus* sequences are indicated in uppercase ; exogenous sequences are marked in lowercase.
 The targeting vector was generated in 129Sv/Pas and was not fully sequenced.

2.2. Map of targeting vector plasmid



2.3. 5' homology arm (4.2 kb)

GCCACTCGTCCTGAGCACTTTGTCCCACCACCGTTGGAATTCATGGAGAAGTGGGAGGAGCTGCTGACTCAGAAC
 GGAAGAGAATGTGTAACACTGCTCTGCTCCTTCATGGCTTCCGAGTCAAGGGGGTGGGGTAGGAAGCTTCAAGG
 ACAGGTGAGGGCTCAGTGTCTCCTGCCAGCCCCAGTACCTTGGATCCAGGAGCCCCGGGCACTTGGCACCAGGCAA
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 GTTTGAAAGCTATAATTCCTCTGCTCAAAATTTCCCTAATTGCCATCCTCCTAGCCAGGCAGCGGAGCTTCAAG
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2.4. Floxed fragment (0.4 kb)

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2.5. PGK-Neo region

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2.6. 3' homology arm (3.2 kb)

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TGGGAAATCTTGGTGGTTGTGCAGGAAGGGTGGCCTCCCCTTGCAGGTTGGGATGGCTCACGGAGAGAGTTTCGCT
CCATGCCTCT

2.7. Vector backbone sequence

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ggaaacagctatgacctgattacgccaagcgcgcaattaaccctcactaaagggaacaaaagctggagctcgcg
gccgcggcgcgc

3. ES cell lines targeted and validation data

3.1. ES cell lines targeted

The targeting vector was electroporated in P1 ES cells [MCI-129Sv/Pas background]

Number of clones screened: 372

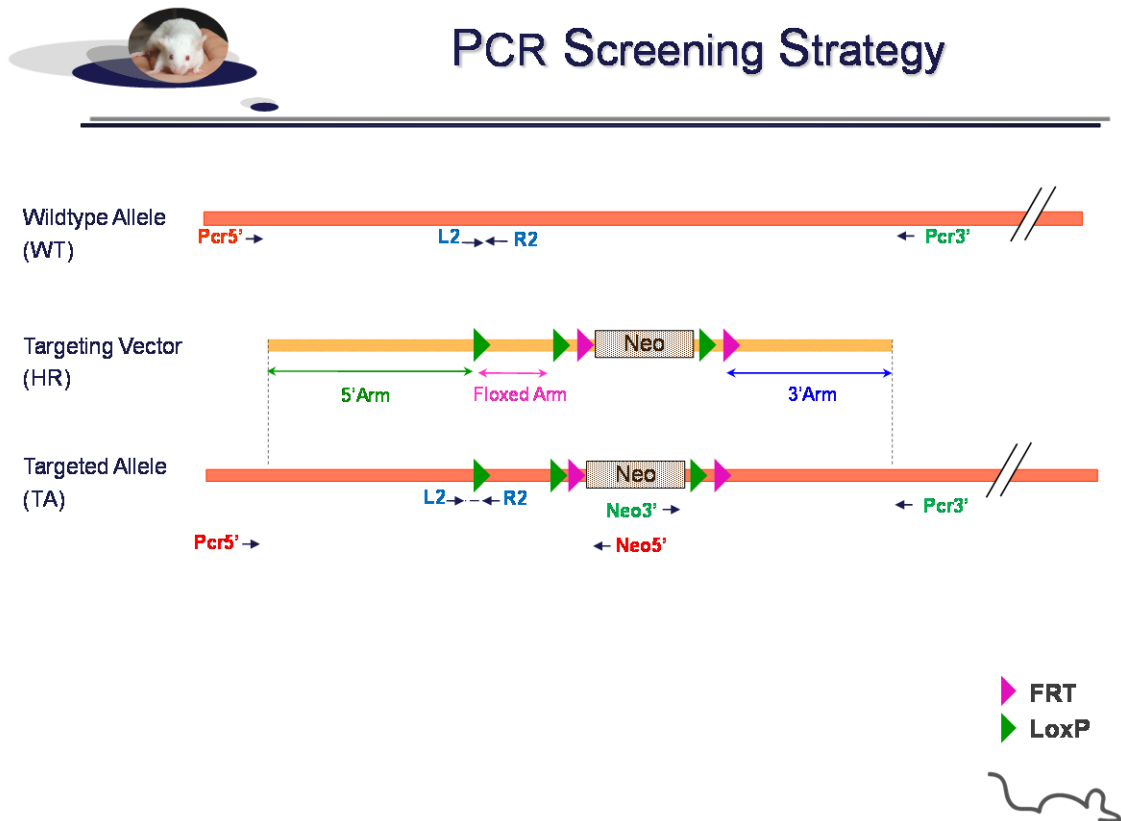
Number of positives: 2

Reference of clone used to generate the mouse line:

- clone **K210P1-22**

3.2. PCR data on positive clone

3.2.1. PCR screening strategy



Primers used for PCR validation of ES clone

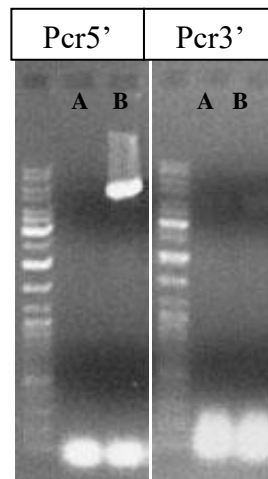
PCR	Primer Name	Primer sequences	PCR product size
5' external	Pcr5'	GGAGTCTATACCCTAGAACACCGAC	5.7kb in TA
	Neo5'	GCGGCCGGAGAACCTGCGTGCAATC	
loxP	L2	ATGCTTGGGTTTATGTTTGAGCTTC	0.20 kb WT 0.25 kb TA
	R2	AAGACACAGATAGGCAGGGTAAGCC	
3' external	Pcr3'	TGTGAACTGGCTCCAGTTGGTTGGG	4.2kb in TA
	Neo3'	AGGGGCTCGCGCCAGCCGAAGTGT	

3.2.2. Picture of PCR

PCR: 5' and 3' arm validation

ladder

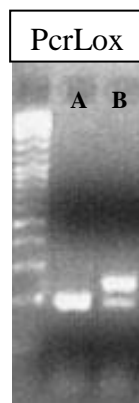
A: negative clone
 B: clone **K210P1-22**



PCR: LoxP validation

ladder

A: negative clone
 B: clone **K210P1-22**

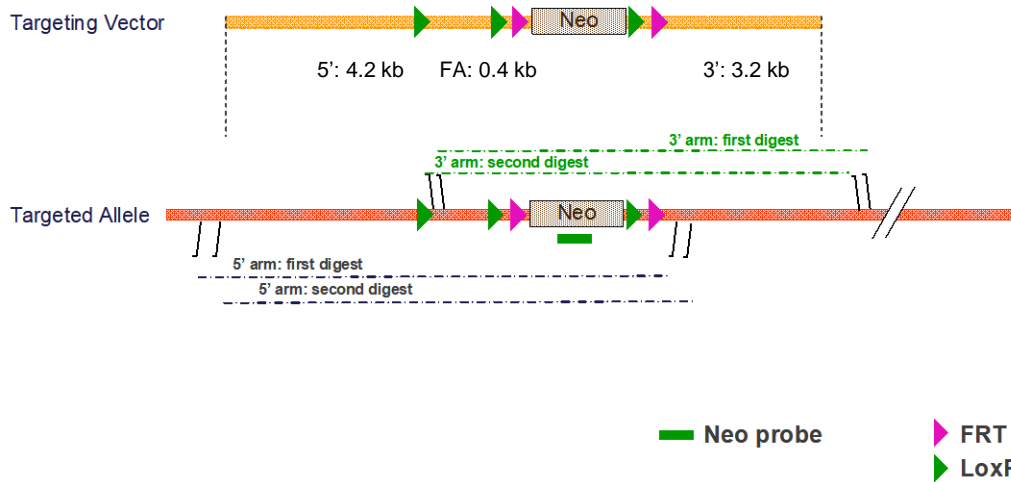


3.3. Southern data on positive clone

3.3.1. Neo Southern strategy



Southern Screening Strategy



Digestions used to validate the 5' and 3' insertion

Probe	Name	Genomic DNA digest	WT allele (kb)	Targeted Allele (kb)
Neo	5' arm first digest	PshAI	/	9.2
	3' arm first digest	Hind III	/	11.0

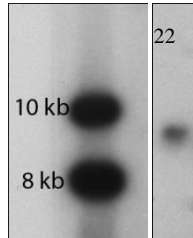
Two different digests are used to validate correct HR event: one digest validate the 5' insertion, one other digest validate the 3' insertion

3.3.2. Picture of Neo Southern

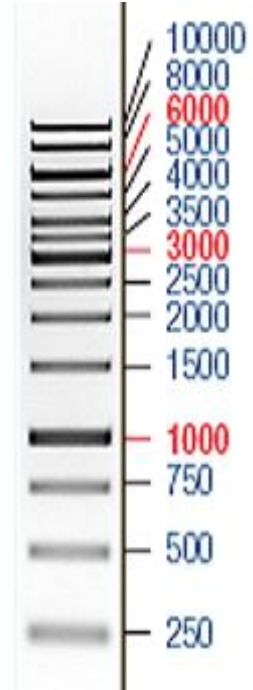
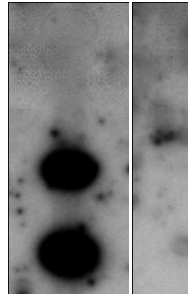
Neo southern blot: 5' and 3'arm validation

ladder

Digestion PshAI



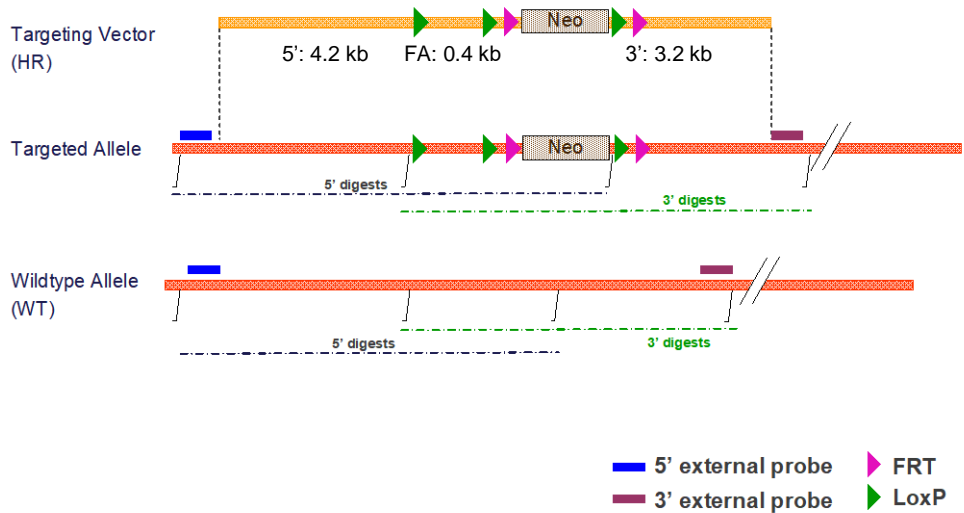
Digestion HindIII



3.3.3. External Southern strategy



Southern Screening Strategy



Digestions used to validate with 5' and 3' probes

Probe	Name	Genomic DNA digest	WT allele (kb)	Targeted Allele (kb)
5' external	first digest	Spe I	11.0	9.7
	second digest	PshAI	7.2	9.2
3' external	first digest	Nde I	9.2	11.2
	second digest	Hind III	9.0	11.0

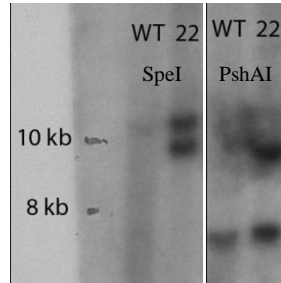
Two different digests are used with each probe.

Sequence of primers used for probes synthesis

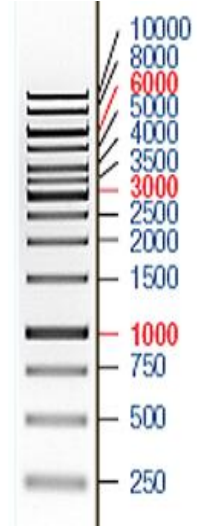
Primers	Sequence
5' probe	CTGTATACTTGTTACTATAATTCTG
	CTGCTGTGTCCAGCTAGCCAAGACT
3' probe	ATCTCTACATGCTACAGAGTATTGG
	TCAAGAAGCGACACAGACCTCATAC

3.3.4. Picture of External Southern

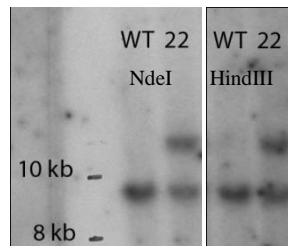
External southern blot: 5'arm validation



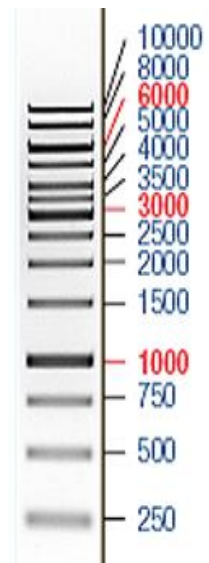
Ladder



External southern blot: 3'arm validation



Ladder



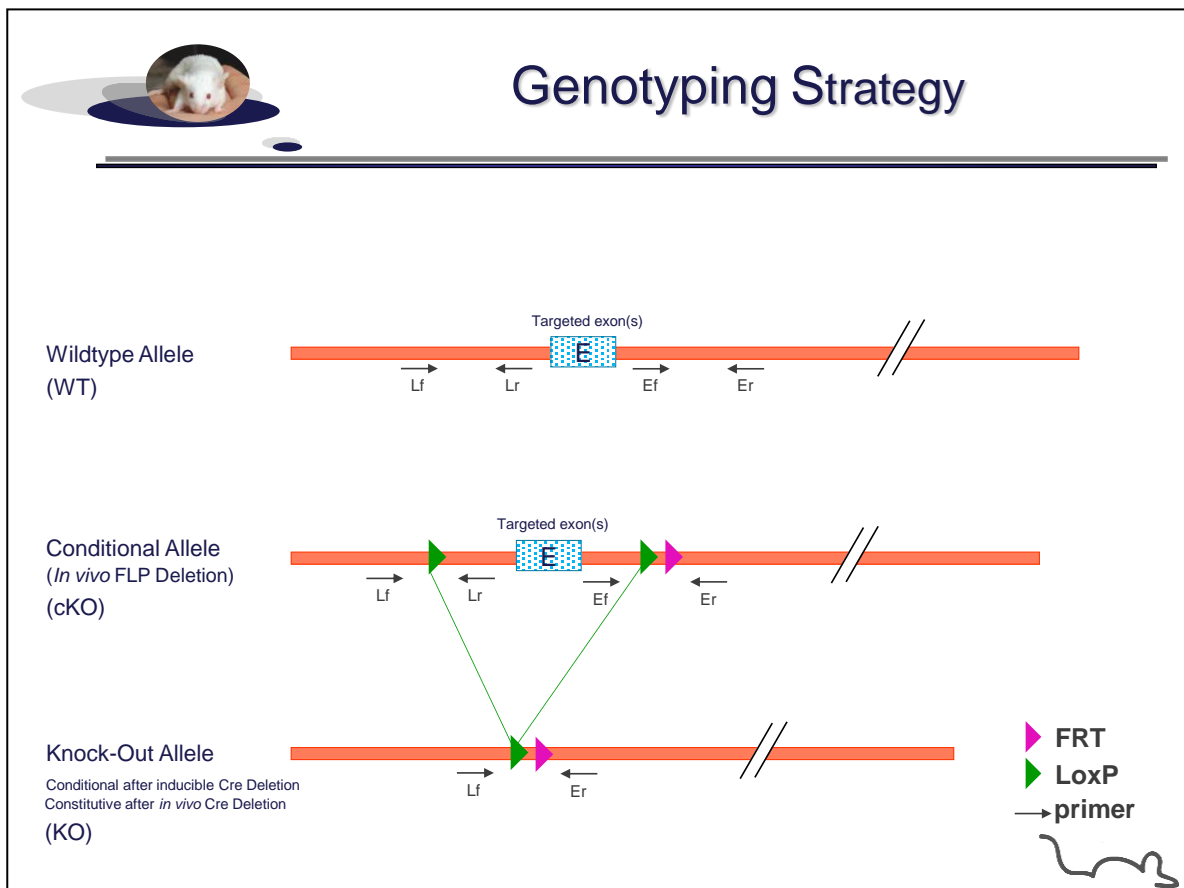
4. Data on conditional and knock-out animals

Both conditional and knock-out mouse models were backcrossed in C57BL/6J background.

4.1. Genotyping protocol and data

4.1.1. Genotyping strategy

The map below describes the position of the primers used for genotyping for each possible allele.



Sequence of primers used for genotyping

Primers	Sequence
Lf	ACAGCTGACATCGTCCTAAGTGTGC
Lr	AAGGCGCGTGCATGTGAGATGAG
Ef	ACATCAAGCCCTTACCAGGTCTGC
Er	TCATCAAGACTGCAGGCTCTGAACC



PCR fragments expected size (bp):

Region analyzed	Primers used	Position on the primer (see the map above)	Conditional allele (cKO)	Knock-Out allele (KO)	WT allele (WT)
Presence of the distal 5'loxP	1220-1221	Lf / Lr	302	---	252
Excision of the selection marker	1223-1225	Ef / Er	377	---	268
Total Excision (excision of the floxed exon(s), i.e. knock out)	1220-1225	Lf / Er	790*	324	630

* This PCR product will not be observed using our PCR genotyping conditions (see description below)
 --- No Amplicon should be obtained

4.1.2.PCR protocol

This section describes the composition of the mix and cycling conditions used for genotyping.

Reagents:	Volume:
-10x Buffer (Roche)	2.5µl
-dNTPs 10mM (Amersham Biosciences)	0.5µl
-Taq DNA Polymerase (Roche)	0.2µl
-DNA (50ng/µl)	3µl
-5' primer (100 µM)	0.125µl
-3' primer (100 µM)	0.125µl
-Sterile H2O	up to 25 µl

Cycling conditions:

Temp	Time	#Cycles
94°C	3min	1
94°C	1min	2
62°C	1min	
72°C	1min	
94°C	30s	30
62°C	30s	
72°C	30s	
72°C	3min	1
4°C	∞	

NB: These PCR conditions have been optimized for high-throughput genotyping. Adaptation to small-scale may be required.

4.1.3. Picture of genotyping with various alleles

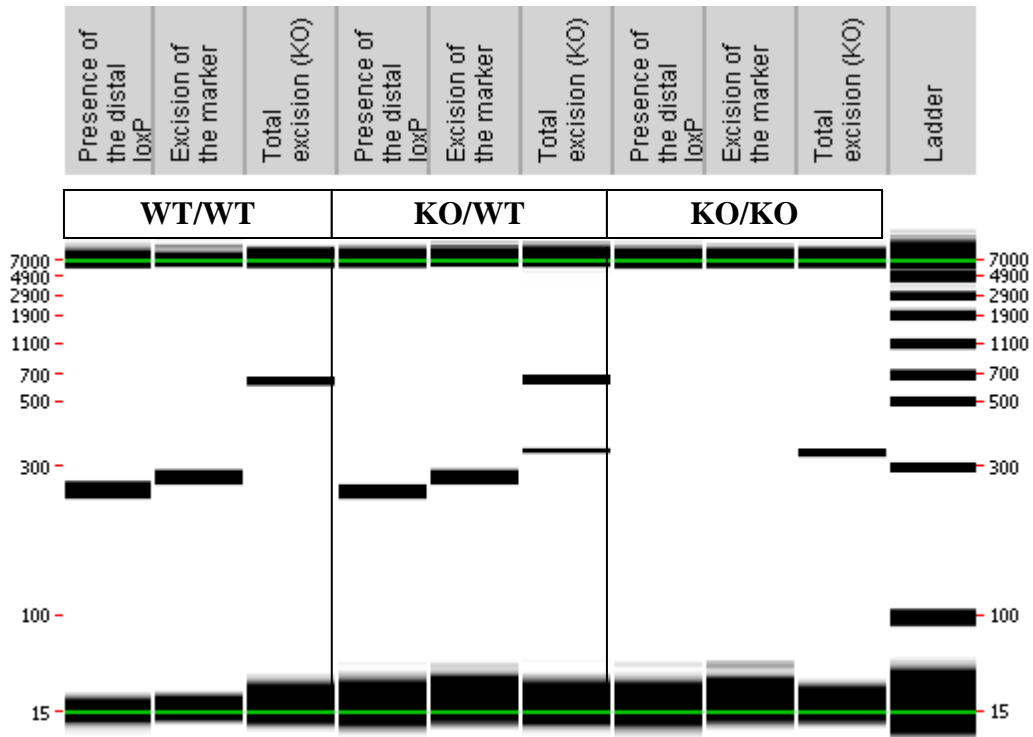
- Picture of genotyping with conditional knock-out (cKO) allele

Data not shown.

- Picture of genotyping with knock-out (KO) allele

Analysis of PCR products pattern was not done by gel electrophoresis but using LabChip® 90 microfluidic apparatus. PCR products were run on the HT DNA 5K LabChip® 90 Assay Kit.

Representative genotyping picture



Note that as this technology is more sensitive than gel analysis, non specific signals and/or primer dimers may be visible on the picture.

4.2. Evaluation of lethality of homozygote KO (KO/KO)

Males knock-out heterozygotes (KO/WT) were crossed with females knock-out heterozygotes (KO/WT). Offspring was genotyped to evaluate the ratio of the different genotypes. Results are provided in the table below.

Genotype	WT/WT	KO/WT	KO/KO	Total
Number of pups obtained	15	66	22	103
Experimental Ratio	14,6%	64,1%	21,4%	100%
Theoretical Ratio	25%	50%	25%	100%
Theoretical Ratio if KO/KO are not viable	33%	66%	0%	100%

The Nr2b3 knock-out homozygotes are viable.

Legend:

- >13% Homozygous = Viable
- >0% and ≤13% = Subviable
- 0% = Lethal