



Genotyping protocol

5330417C22Rik

IR00003910 / E259

(ICS internal reference)

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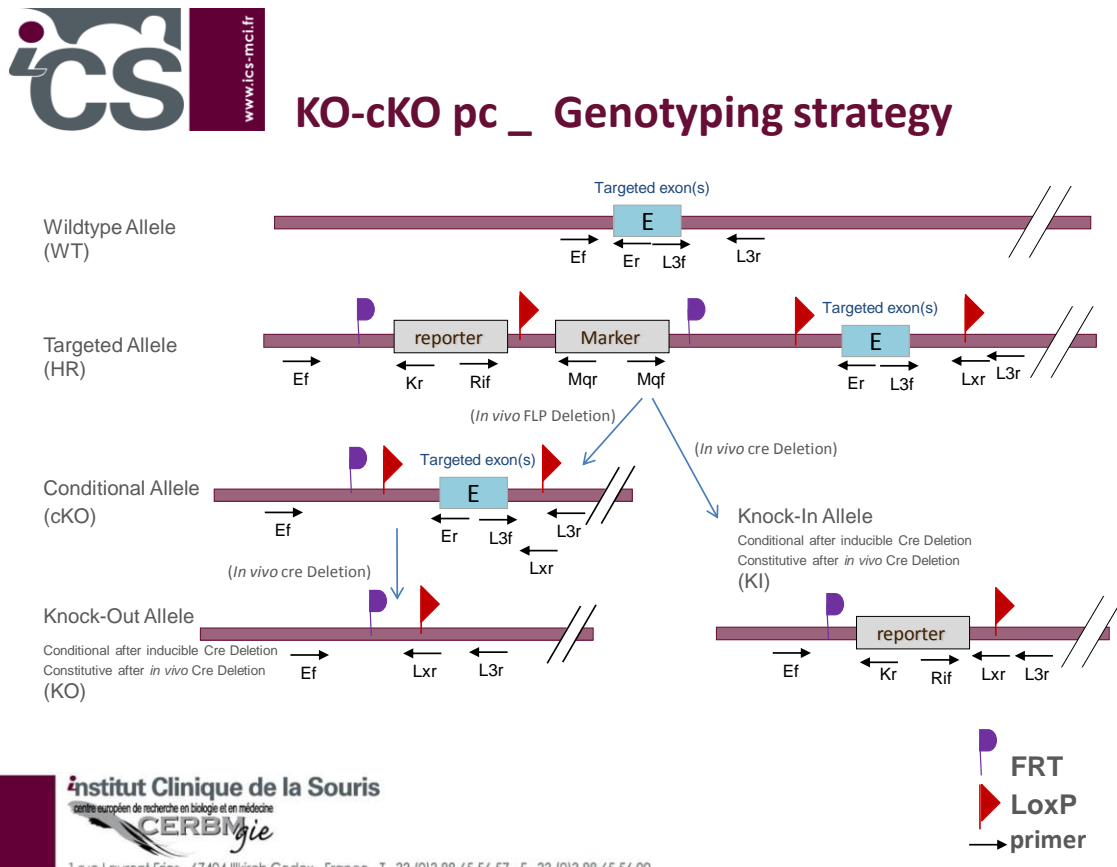
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1. Genotyping protocol and data

This section describes the condition used at the Mouse Clinical Institute (ICS) to genotype your **5330417C22Rik** Constitutive Knockout / Conditional Knockout (KO-cKO x Cre) project.

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:

| Position | Primers | Sequence |
|-----------------|---------|-----------------------------|
| Ef | 6777 | GATCTTAAATGCACAACAATCCTCCCC |
| Ef ² | 6778 | TTCACCCCTTACTGAGCAAAGTTGGAG |
| Er | 6782 | TCTTAATGCTCCCAGACTTGAGTGGTT |
| Kr | 3278 | GGGCAAGAACATAAAGTGACCCTCC |
| L3f | 6779 | TCGATGACAGCATCTGGAGTCCAC |
| L3r | 6780 | CCCTGCTAAGCCATTAGATGCCTG |
| Lxr | 3255 | ACTGATGGCGAGCTCAGACCATAAC |
| Ri1f | 5966 | GCACATGGCTGAATATCGACGGT |

²: for a selected position, a second primer was designed

PCR fragments expected size (bp):

| Region analyzed | Primers used | Position on the primer <i>(see the map above)</i> | Targeted allele (HR) | conditional allele (cKO) | KI allele | WildType allele |
|----------------------------------|--------------|--|----------------------|--------------------------|-----------|-----------------|
| 5' part of the selection marker | 6777-3278 | Ef / Kr | 337 | --- | 337 | --- |
| Presence of the distal loxP | 6779-6780 | L3f / L3r | 341 | 341 | --- | 305 |
| Distal loxP specific PCR | 6779-3255 | L3f / Lxr | 300 | 300 | --- | --- |
| Excision of the selection marker | 6778-6782 | Ef ² / Er | 7274* | 370 | --- | 235 |
| Cre total excision | 5966-3255 | Ri1f / Lxr | 3259* | --- | 471 | --- |

*: this PCR product will not be observed using our PCR genotyping conditions (see description below)

**: this PCR is only verified if mice are generated

---: no Amplicon should be obtained

1.2. PCR protocol

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

Reagents:

- FastStart PCR Master (Roche)
- DNA (50ng/ μ l)
- 5' primer (100 μ M)
- 3' primer (100 μ M)
- Sterile H₂O

Volume:

- 7.5 μ l
- 1.5 μ l
- 0.06 μ l
- 0.06 μ l
- up to 15 μ l

Cycling conditions:

| Temp | Time | #Cycles |
|------|------|---------|
| 95°C | 4min | 1 |
| 94°C | 30s | |
| 62°C | 30s | 34 |
| 72°C | 1min | |
| 72°C | 7min | 1 |
| 20°C | 5min | 1 |

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

2. Cre and Flp genotyping method

You will find the genotyping protocol in the publication:

[Highly-efficient, fluorescent, locus directed cre and FlpO deleter mice on a pure C57BL/6N genetic background.](#)

Birling MC, Dierich A, Jacquot S, Héault Y, Pavlovic G.
Genesis. 2012 Jun;50(6):482-9. doi: 10.1002/dvg.20826. Epub 2012 Mar 20.