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Nf1 loxP site verification

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working

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Abstract

We have found that our Nf1 flox mouse colonies contained mice that had lost one of the two loxP sites inserted into the Nf1 gene (Parada laboratory, PMID:11297510, JAX# 017639), leading to loss of DNA recombination upon Crerecombinase activity and loss of previously identified skeletal phenotypes. This is likely to occur in other Nf1 flox mouse colonies.

We describe here a strategy to verify conservation of the loxP sites within the Nf1 allele and to sequence the loxP1 and loxP2 sites for detecting potential mutations or deletions. This may be done if the expression of Nf1 (qPCR) is not reduced in $Cre^{Tg/+}$; Nf1 flox/flox tissues of interest.

Attachments



Exon 31_32 and loxP ...

1.3MB

Materials

Heater (DNA denaturation/Lysis)

PCR machine

Agarose gel electrophoresis system (DNA separation)

Gel documentation system (DNA amplicon visualization and picture)

Protocol materials

Specific Primers Forward and Reverse Step 1

GoTaq® Green Master Mix Promega Corparation Catalog #M7123 Step 3.2



Nf1 loxP site verification

15m

15m

1 Specific Primers Forward and Reverse Contributed by users

Primers for the Nf1 loxP1 site:

P1: 5'-CTTCAGACTGATTGTTGTACCTGA-3' P4: 5'-TGATTCCCACTTTGTGGTTCTAAG-3' P3: 5'-ACCTCTCTAGCCTCAGGAATGA-3'

Primers for the Nf1 loxP2 site:

LoxP2For: 5'-GCTTTAGCTTCTGGAAATGTGAA-3' LoxP2Rev: 5'-GCGGGCTAAAATGGCAATGTCG-3'

2 gDNA preparation from NaOH lysate

1. Cut a 2-3mm piece of tail/ear

- 2. Add it to 300uL of lysis solution (NaOH 50mM)
- 3. Heat for 15 min at 95C
- 4. Room temperature, 00:15:00 , use max speed of your bench centrifuge

3 PCR reaction

3.1 From a working solution of 10uM, use 1 uL of each primer (200nM each in a total volume of 20uL)

Reaction for loxP1: P1 + P2 + P3

Reaction for lopP2: LoxP2For + LoxP2Rev

3.2 Per reaction, mix 0.5uL of gDNA from the NaOH lysate (from step 2), 1 uL of each primer (from step 3.1), 10 uL GoTaq® Green Master Mix Promega Corparation Catalog #M7123 mix, and add water for a total volume of 20 uL

10m

3.3 PCR cycle program

2h

	А	В	С
	Temp.	Duration	Cycles
	94C	5 min	
Г	94C	15 sec	3 to 5, 35x
Г	58C	30 sec	
Г	72C	1 min	
	72C	10 min	



PCR temperatures and cycles

4 Run PCR reaction products on a 2% ethidium bromide agarose gel

1h

5

Expected result

Expected PCR results:

LoxP1 site: a)

480bp: WT (+) allele (P1-P3) in $Nf1^{+/+}$ or $Nf1^{f/+}$ mice, or loss of the loxP1 site in Nf1350bp: Presence of loxP1 (P1-P4)

in Nf1^{f/f} mice

LoxP2 site: b)

699 bp: WT (+) allele in $Nf1^{+/+}$ or $Nf1^{f/+}$ mice, or loss of the loxP2 site in $Nf1^{f/f}$ mice 829 bp: Presence of loxP2 in Nf1 f/f mice

Protocol references

Ablation of NF1 function in neurons induces abnormal development of cerebral cortex and reactive gliosis in the brain, PMID: Y Zhu 1, M I Romero, P Ghosh, Z Ye, P Charnay, E J Rushing, J D Marth, L F Parada, PMID:11297510.