

Sep 19, 2024

Nf1 loxP site verification

DOI

dx.doi.org/10.17504/protocols.io.261ge3ryyl47/v1

Greig Couasnay¹, Florent Elefteriou¹

¹Baylor College of Medicine

Baylor College of Medicine



Florent Elefteriou

Baylor College of Medicine

OPEN  ACCESS



DOI: dx.doi.org/10.17504/protocols.io.261ge3ryyl47/v1

External link: <https://www.jax.org/strain/017639>

Protocol Citation: Greig Couasnay, Florent Elefteriou 2024. Nf1 loxP site verification. **protocols.io**
<https://dx.doi.org/10.17504/protocols.io.261ge3ryyl47/v1>

License: This is an open access protocol distributed under the terms of the [Creative Commons Attribution License](#), which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited

Protocol status: Working

We use this protocol and it's working

Created: December 20, 2022

Last Modified: September 19, 2024

Protocol Integer ID: 74278

Keywords: Neurofibromatosis type 1, NF1, Genotyping, PCR, loxP

Funders Acknowledgement:

NIH

Grant ID: R01AR077949



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 Cre-recombinase 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 Corporation Catalog #M7123** Step 3.2



Nf1 loxP site verification

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'-GCGGGCTAAAATGGCAATGTGCG-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. **max rpm, Room temperature**, 00:15:00 , use max speed of your bench centrifuge

15m

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 loxP2: 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 Corporation Catalog #M7123** mix, and add water for a total volume of 20 uL

10m

3.3

PCR cycle program

2h

A	B	C
Temp.	Duration	Cycles
94C	5 min	
94C	15 sec	3 to 5, 35x
58C	30 sec	
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:**a) LoxP1 site:**

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

350bp: Presence of loxP1 (P1-P4) in *Nf1*^{f/f} mice

b) LoxP2 site:

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.