

CLINICAL TRIAL

NCT00422721



Retinal RED

Genetic Study of Patients Suffering From Congenital Amaurosis of Leber or From an Early Severe Retinal Dystrophy

Rare Eye Disease concerned by the trial :
Amaurosis | Retinal Diseases

Status of the trial : completed
Orphan drug recognition : N/A

Inclusion criteria : [Click here to see detailed criteria](#)

Exclusion criteria : [Click here to see detailed criteria](#)



Inclusion
opening date : 01/04/2007



Inclusion
closing date : 24/11/2011

Children



Adults



Within ERN-EYE members



Principal location of the trial :

CHU de Nantes

1 Place Alexis-Ricordeau
44093 Nantes
France



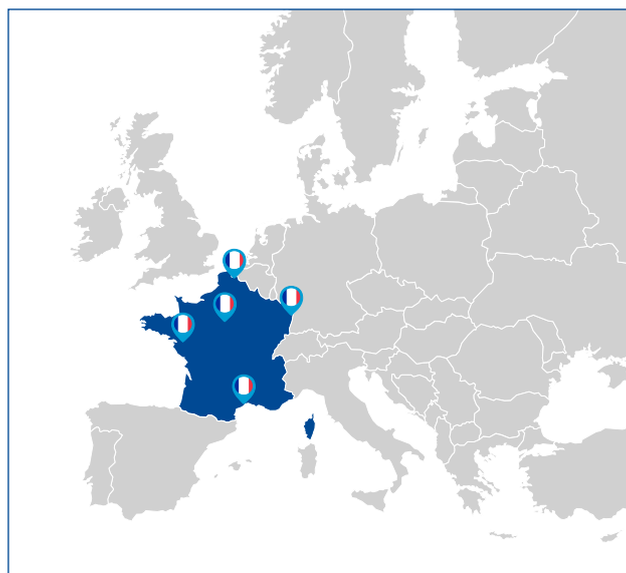
Principal investigator name :

Pr Michel Weber



Other investigators :

Dr Sabine Defoort, CHU de Lille
Dr Bernard Puech, CHU de Lille
Dr Isabelle Drumaré, CHU de Lille
Pr Christian Hamel, CHU de Montpellier
Pr Carl Arndt, CHU de Montpellier
Dr Olivier Roche, Hôpital Necker, Paris
Dr Christophe Orssaud, Hôpital Necker, Paris
Dr Emmanuel Bui Quoc, Hôpital Necker, Paris
Dr Saddek Mohand Saïd, CNO XV-XX, Paris
Pr José-Alain Sael, CNO XV-XX, Paris
Pr Hélène Dollfus, CHU de Strasbourg



Funder type : Public

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