

# CLINICAL TRIAL

## CURETINA

Retinal RED



### CURETINA personalized medicine in hereditary retinal degenerations

Rare Eye Disease concerned by the trial :  
hereditary retinal degenerations caused by  
mutations in genes CRB1, CRX, PROM1

Status of the trial : recruiting

Orphan drug recognition : N/A

**Inclusion criteria :** Retinal degeneration caused by mutations in either of the genes CRB1, CRX, PROM1

**Exclusion criteria :** Additional mutations in genes known for causing hereditary retinal degenerations; additional eye diseases influencing the retinal morphology and function (glaucoma, retinal detachment, etc.)



Inclusion  
opening date : 01/02/2016



Inclusion  
closing date : 30/06/2018

Children



Adults



### Within ERN-EYE members



Principal location of the trial :

**Center for Ophthalmology, University of Tübingen**

Elfriede-Aulhorn-Str. 7  
72076 Tübingen  
Germany

[Contact details](#)



Principal investigator name :

Katarina Stingl

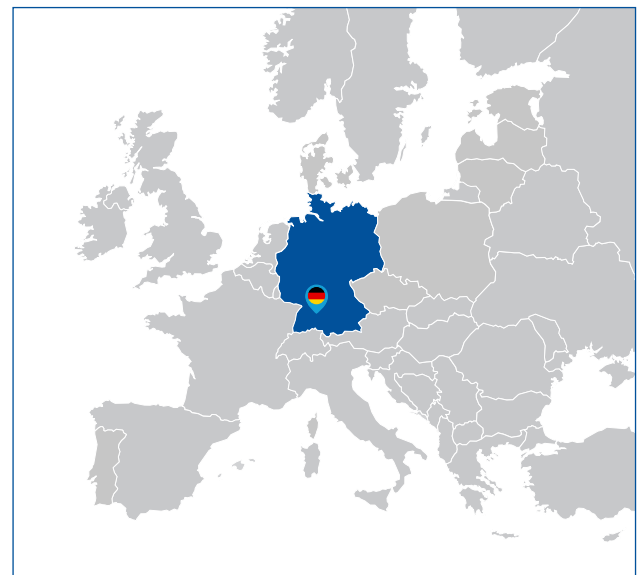


Other investigators :

Laura Kühlewein

**Center for Ophthalmology, University of Tübingen**

Tübingen, Germany



Funder type : National institution

[More informations](#)