

Participation & Registration in the Patient Registry





Participation

All persons with retinitis pigmentosa caused by an *RP2* gene variant are eligible.

Minors need the consent of both parents.

Your participation is voluntary.

You can withdraw your consent at any time without giving any reasons.

Secure communication

E-mail communication is encrypted using the SecuMails software.

Data collection

We collect genetic, clinical and imaging data of affected individuals.

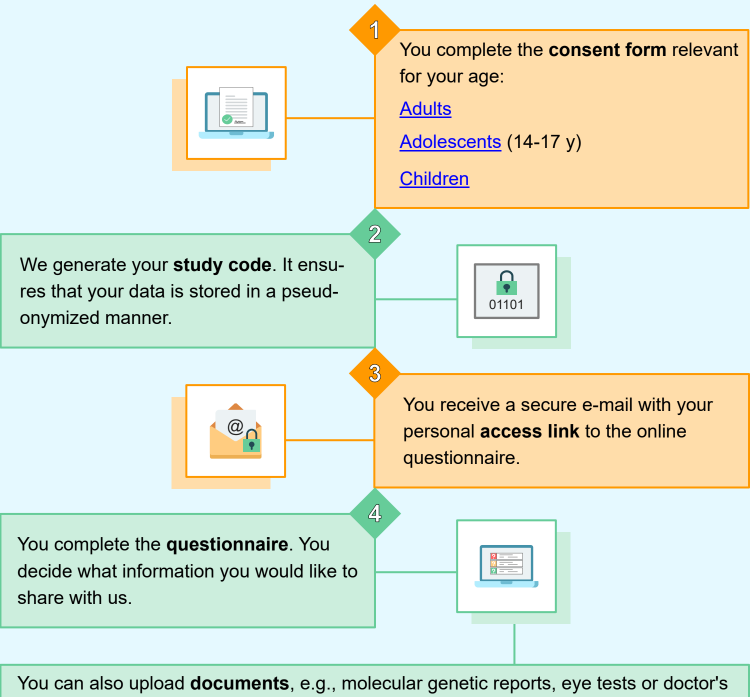
Data protection

Data is analyzed and stored in a pseudonymized manner in accordance with the European General Data Protection Regulation (GDPR). Personal data is replaced to make it more difficult to identify participants.

Data will be used exclusively for the purposes of the study and will not be passed on to third parties.

Registration

Participants will register online in two steps. First, you complete a consent form and then a questionnaire. This keeps your personal and medical data separate from each other.



You can also upload **documents**, e.g., molecular genetic reports, eye tests or doctor's letters.

Please remove/black out all personal data (e.g. name, date of birth, address) before uploading your documents.

Questions & contact

If you have any questions about the registration process or the study, please send an e-mail to the study manager, Dr. med. Nina Bögershausen: