ADVANTAGES TO CLINICAL SITE MEMBERS

- **Participation in subspecialty Scientific Sessions**
- Support for multinational Investigator-Initiated Research
- Receive for free EVICR.net organisational SOPs and become a certified Clinical Site
- Opportunity to participate in new multinational clinical research studies
- Support for EU Clinical Research applications
- Support in grant applications to industry
- Access to educational programme and training
- Promotion of your Clinical Research Centre as member of EVICR.net
- Networking with other European and USA networks

BECOME A MEMBER



SCIENTIFIC

REVIEW

STEERING COMMITTEE

Hendrik Scholl (Chairman) Francesco Bandello Marie-José Tassignon Cecília Martinho



Your partner in **INVESTIGATOR-INITIATED RESEARCH** (IIR)

CLINICAL RESEARCH **EXPERTISE**

SUPPORT IN

GRANT

APPLICATIONS

95 CLINICAL

SITES

IF YOU ARE AN INVESTIGATOR AND YOU HAVE AN IDEA FOR AN **OPHTHALMOLOGY MULTINATIONAL CLINICAL STUDY**, EVICR.net CAN HELP YOU

VISIT US



CONTACT US

EVICR.net Coordinating Centre AIBILI, Edif. Prof. Doutor José Cunha-Vaz Azinhaga de Santa Comba 3000-548 Coimbra, Portugal



JANUARY 2024

EDUCATIONAL PROGRAMME



TAKE ADVANTAGE OF WEBINARS ON OPHTHALMOLOGY CLINICAL RESEARCH

José Cunha-Vaz Francesca Cordeiro Pasquale Aragona **Birgit Lorenz** Tunde Peto



A NETWORK OF 95 CLINICAL RESEARCH SITES FROM 16 COUNTRIES

WE SUPPORT:

- Design and implementation of multinational clinical research activities in ophthalmology
- Grant applications to Pharmaceutical industry or to European Union

SCIENTIFIC REVIEW AND EXPERTISE



EXPERT COMMITTEES AND COORDINATORS

- AGE-RELATED MACULAR DEGENERATION Francesco Bandello
- **RETINAL DYSTROPHIES** Hendrik Scholl
- DIABETIC RETINOPATHY & VASCULAR DISEASES José Cunha-Vaz
- **GLAUCOMA** Francesca Cordeiro

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- ANTERIOR SEGMENT Marie-José Tassignon
- Ocular Surface, Inflammation, Dry Eye & Allergies Pasquale Aragona

TRANSVERSAL SECTIONS AND COORDINATORS

- RARE DISEASES Birgit Lorenz
- **READING CENTRES** Tunde Peto

WE PROVIDE SUPPORT TO

PRE-STUDY SERVICES

- Study design
- Study Protocol development
- Informed Consent Form development
- Case Report Form design
- Database validation and implementation
- eCRF management and support

- Clinical Sites feasibility
- SOP development
- Study Documents development
- Regulatory affairs
- (submission and reports)
- Contracts

POST-STUDY SERVICES

Biostatistics

- Final Study report
- Publication
- Medical Writing

How to proceed?

IN-STUDY SERVICES

Study management

IMP management

Data management

Pharmacovigilance

Monitoring

If you are a member:

 Submit your idea by filling the abstract available at <u>www.evicr.net</u> (MEMBERS RESTRICTED AREA)



If you are not an EVICR.net member yet:

Please contact the Coordinating Centre: evicrnet@aibili.pt