



Worldwide Cystinosis Community Advisory Board

What are the key principles of the CAB?

- 1. Neutral setting
- 2. Confidentiality
- 3. Transparency
- 4. Sharing
- 5. Openness
- 6. Optimisation of research and development
- 7. A joint action plan as an outcome of the meeting

Who are the Worldwide Cystinosis Community Advisory Board?

The CAB includes representatives from:

AIRG-España AIRG France

Asociación Cistinosis España

Associazione Cistinosi, Italy

Cystinose France

Cystinosis Group Belgium

Cystinosis Ireland (incl Northern

Ireland)

Cystinosis Foundation South Africa

Cystinosis Foundation UK

Cystinosis Research Network, USA

Cystinose-Selbsthilfe e.V., Germany Cystinosis Support Group Australia

Dutch Cyctinocic Group

Dutch Cystinosis Group

What is a Community Advisory Board (CAB)?

A group trained of patients and patient representatives who offer their expertise to public or private sponsors of clinical research and early stage researchers.

The same group of patients advises several sponsors in their field.

Agenda and secretariat driven by the patients.

The CAB remains an autonomous and independent board.



What can the CAB offer?

Expertise on:

- patient outreach
- education on research
- clinical studies and their design
- criteria for participation
- informed consent forms and processes
- compassionate use programmes
- better retention of participants
- reporting on results
- autonomy and independence

Worldwide Cystinosis CAB work to date

July 2018: Briefing from Eurordis at Berlin Cystinosis Conference.

January 2019: First internal meeting and commitment to proceed. Chair elected.

March 2019: Training begins.

April 2019: Initial engagement with sponsors begin.

November 2019: First CAB and Sponsors meeting to be held in Dublin, Ireland.

