



The role of patient engagement in clinical research

Kerry Leeson-Beevers National Development Manager, Alström Syndrome UK

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ZAANDAM





Alström Syndrome UK



https://youtu.be/XZx1LSrWZyo





Living with Alström Syndrome









Developing my Understanding

- ➤ World turned upside down
- ➤ Resigned from work due to Kion's ill health
- > Joined the team at ASUK
- ➤ Training & Development
- ➤ 2008 & 2010 attended the EURORDIS Summer School to develop my understanding of clinical trials and EU regulatory affairs
- ➤ Became a volunteer for EURORDIS and joined the Paediatric Committee at the EMA as a patient representative until December 2016





Continued....

- ➤ Joined the European Public Affairs Committee with EURORDIS
- ➤ Joined the Patient Engagement Group with Rare Disease UK
- ➤ Patient Think Tank member for the Asterix Project Advances in Small Trials dEsign for Regulatory Innovation and eXcellence http://www.asterix-fp7.eu/
- > Patient Representative on the UK Rare Disease Policy Board
- ➤ National Development Manager with a particular focus on research, NHS clinics and clinical trials
- ➤ Project Lead for Breaking Down Barriers





Patient Support

- ➤ Organise clinic lists
- Advocacy
- ➤ Clinic support
- ➤ Link between local professionals and specialist centres
- ➤ Annual audit patient feedback

Current Projects

- ➤ Transition T-Kash resources developed by young people
- ➤ Breaking Down Barriers
- > Research Consortium
- ➤ Clinical Trial



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Developing supportive & inclusive services for individuals and families affected by genetic disorders



Understanding genetics together

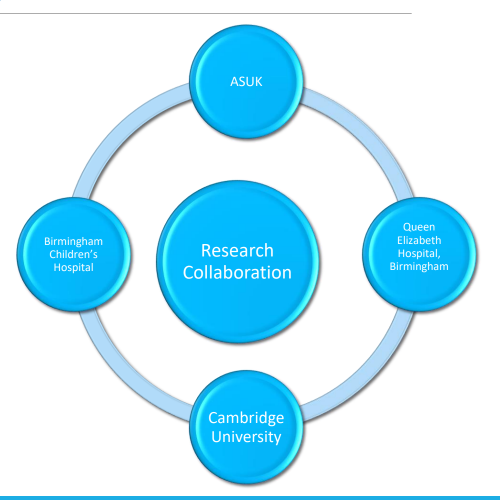




Research – Big Lottery Funding

"....research project aimed at deepening our understanding of Alström Syndrome, improving diagnosis and management while investigating the basic cellular disease mechanisms in AS patients"

- ➤ Database
- ➤ Tissue Bank
- **EuroWABB**







Clinical Trials

"PBI-4050 aims to halt the progression and possibly reverse some of the effects of fibrosis which is one of the main problems associated with Alström Syndrome. The trial drug may also have the potential to improve sugar and fat levels within the body, improving diabetes and obesity and

reducing insulin resistance."

"The primary aim of the trial is to evaluate the safety and tolerability of PBI-4050 and its effects on inflammatory, fibrosis, diabetes and obesity biomarkers in people affected by Alström Syndrome."







ASUK Conference

Health, Happiness and Well-being!

Medical and Scientific Symposium & Family and Professional Conference

6th and 7th October 2017



Aston University Conference Centre in Birmingham, UK





Patient Think Tank Members

- ➤ Elizabeth Vroom
- **≻**Oliver Timmis
- ➤ Jörg Richstein
- > Radoslaw Kaczmarek
- ➤ Marleen Kaatee
- ➤ Veronica van Nederveen
- ➤ Mieke Donk
- ➤ Noirin O'Neil
- ➤ Christine Lavery
- ➤ Kerry Leeson-Beevers

Supported by the VSOP (Dutch umbrella organisation for rare, genetic and congenital disorders)





Patient Leaflets

- ➤ Goal Attainment Scaling
- ➤ Multiple Endpoints
- ➤ Adaptive Designs
- ➤ Meta Analyses (with zero events)
- ➤ Clustering Framework of Diseases
- ➤ Patient Registries
- **Randomisation**
- ➤ Patient Involvement in Rare Disease Clinical Research
- ➤ Why trials may stop early
- ➤ Adaptive Pathways

Patient Involvement

in Rare Disease Clinical Research



Why should patients be involved?

Patients and parents of a child with a rare or ultra-rare disease can make an important contribution to the trial becoming a success. As patients manage their disease on a day to day basis, they have more knowledge on which aspects of their disease should have priority for further investigation by researchers. Patients are able to offer a unique perspective based on this experiential knowledge. Patients should be involved in the choice of outcome measures in a trial and the composition of patient information. They better understand the burden of trial participation for them and their families, and can help researchers design a more accessible trial. In this leaflet, we will show how patients can be involved in research, and which tools may be used.



Roles of Patients in clinical research

This diagram represents the different roles that patient representatives play in the clinical trial process: a research subject, an information provider, an advisor, a reviewer, a co-researcher and a driving force. This diagram was developed in the EU project patient partner, based on the Participation ladder of Arnstein, a vertical ladder. All roles are necessary and important and there is no hierarchy of one above the other, thus the ladder was turned. Patients can be involved in clinical trials in various ways: setting the research agenda, design of clinical trials, recruitment and dissemination of the results. The Asterix project studies the design of better clinical trials for small populations.

European collaborations on patient involvement in research

EURORDIS

Eurordis is the alliance of rare disease patient organizations in Europe. Over 700 patient organisations are members of Eurordis, and they represent over 4000 rare diseases. The organization is non-governmental and driven by patients. Eurordis represents the patients' voice on a European level. Eurordis strengthens the capacity of patient advocates and provides training on clinical trials, drug development and regulatory processes in their Summer

School. Trained patients are partner of researchers, and are members of various committees at the EMA, such as the Patients' and Consumers' Working Party (PCWP).

EUPAT

EUPATI, short for 'European Patients' Academy on Therapeutic Innovation' offers information and training to patients about healthcare research. EUPATI organises Patient Expert Training Courses for patients. The EUPATI toolbox on Medicines R&D is an information tool in several languages, which patients can use to contribute to Research & Development of new treatments with researchers, policy makers, and the pharmaceutical industry.

The EMA framework for interaction with Patients

The European Medicines agency involves patients in several of

her activities. Patients are members of the Management Board and scientific committees; are consulted on disease-specific requests by the committees and working parties; take part in discussions on the development and authorisation of medicines and in the preparation of guidelines. The EMA has developed Training & Resources for patients, like video's or workshops.

Asterix methods: the POWER model

In the ASTERIX project, we have developed a model where patient representatives are involved in the choice of outcomes in the design stage of a trial, the POWER model (Patient participation in Outcome measure WEighing for Rare diseases). Patients and researchers decide together which aspects of a disease are relevant to patients and which outcomes we will measure to evaluate an experimental treatment.

The POWER model has four steps

- Create the right circumstances for communication between researchers and patient representatives
- 2 Prepare a consultation, for example with training on research methods or communication skills
- 3 Consultation, where patient representatives and researchers interact, for example trough focus groups or interviews and where they make decisions on how to progress
- 4 Follow-up, for example an evaluation with the patient representatives and researchers and agreements how to keep each other up-to-date on the progress of the trial

Possible benefits for patients

When scientists involve patients in the early stages of a trial, the trial may:

- Investigate topics that are more relevant to patients
- Become more appealing for patients to take part in

When the trial is tailored more towards patients, the efficacy of a drug may be demonstrated in endpoints that are relevant to patients.

Possible downsides

- It may be possible that patient representatives who have been involved in the design stage of a trial cannot take part in the trial themselves, because they do not fit the inclusion criteria. It is important to manage these expectations, as patient representatives often put in a lot of time and effort and hope to find a cure.
- The dedication and time that patient representatives put in research projects is not always acknowledged.

More information

EMA/Partners&Networks/Patients&Consumers

EUPATI: https://www.eupati.eu/

Eurordis: http://www.eurordis.org/content/patient-advocates-involvement

PatientPartner: Guide for Patient Representatives

Participation ladder: Arnstein SR. A ladder of citizen participation. Journal of the American Institute of planners. 1969;35(4):216-24.

There are many more national initiatives on patient involvement in research. However, in this leaflet, we focus only on the European collaborations.

Contact details





ASTERIX - Advances in Small Trials d sign for Regulatory Innovation and excellence Intis leaflet is developed together with the Patient Phink Tank and is part of the Asterix project, funded by the EU FP-7 program. For more information, see our website: www.asterix-fp7.eu













Thank You

Contact Details:

Kerry Leeson-Beevers

National Development Manager

Alström Syndrome UK

Tel: 01709 210151 / 07716135940

Email: Kerry.leeson@alstrom.org.uk

Web: www.alstrom.org.uk

www.breaking-down-barriers.org.uk

Facebook: <u>www.facebook.com/alstromsyndromeuk</u>

Twitter: @AS_UK



Certified member

This organisation has been certified as a producer of reliable health and social care information. www.theinformationstandard.org