Looking for information and collaborative opportunities?

ECRD PROGRAMME
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WHAT IS ECRD?

› The European Conference on Rare Diseases & Orphan Products is the unique forum across all rare diseases, across all European countries, bringing together all stakeholders - patients’ representatives, academics, researchers, health care professionals, industry, payers, regulators and policy makers.

› ECRD provides the state-of-the-art of the rare disease environment, monitoring and benchmarking initiatives.

› ECRD covers research, development of new treatments, health care, social care, information, public health and support at European, national, regional and international levels.
Hear from patients first-hand about their needs and experiences and get insight into how to work more efficiently with patient organisations

See the full picture of how different elements are coming together at global, regional and country level to support orphan medicinal products discovery, development and availability

Dialogue with all of the stakeholders shaping your environment: patients, policy makers, regulators, member state representatives, academia, researchers, clinicians, HTA evaluators, payers and industry colleagues – together in one venue

Understand the state-of-the-art for orphan development as well as trends for the future

Evaluate new pathways for collaboration and partnership at all stages of the development chain

Explore the implications of emerging pricing, reimbursement and access trends together with those responsible for the markets

Discover areas of common interest and establish collaborations

Enlarge your network and be part of a community that is focused on improving the lives of rare disease patients

Voice your own point of view, be seen as a thought leader in the rare disease community and contribute to its development

LANGUAGES

Simultaneous interpretation and live streaming of the Opening and Plenary sessions on 11 May 2018 will be available in:

ENGLISH  FRENCH  GERMAN

For more information, please visit www.rare-diseases.eu

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