## SOME OF OUR **ACHIEVEMENTS**

- > Contribution to the adoption of EU regulations on Orphan Medicinal Products (1999), Paediatric Drugs (2006) and Advanced Therapy Medicinal Products (2007)
- > Contribution to the EU Commission Communication on Rare Diseases (2008), the EU Council Recommendation on an action in the field of rare diseases (2009) and the EU Directive on Patients' Rights in Cross-Border Healthcare (2011)
- > Promotion of rare diseases as an EU public health priority
- > Promotion of national rare disease plans in EU Member States and other countries
- > Promotion of rare diseases as a priority in the EU Research Framework Programme
- > Contribution to bringing a significant number of rare disease products to market (over 1,800 orphan designations and over 130 orphan medicinal products)
- > Organisation of the European Conference on Rare Diseases & Orphan Products (ECRD), held every two years since 2001
- > Creation and coordination of the international campaign Rare **Disease Day**, held since 2008 in over 94 countries and regions
- > Training of 400 patient representatives and researchers from 40+ countries through the EURORDIS Summer School
- > Crucial role in establishment and development of the 24 European Reference Networks (ERNs) and creation of 24 European Patient Advocacy Groups (ePAGs) involving around 1,000 patient organisations and 200 ePAG patient representatives





### **OUR MISSION**

**CC** EURORDIS works across borders and diseases to improve the lives of people living with a rare disease >>

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EURORDIS is supported by its members, the AFM-Téléthon, the European Commission, foundations and the health industry. EURORDIS was founded in 1997.







The Voice of **RARE DISEASE PATIENTS** in Europe



**G** Better lives and cures for people living with a rare disease

**EURORDIS.ORG** 

# WHO **WE ARE**

EURORDIS-Rare Diseases Europe is a unique, non-profit alliance of over 700 rare disease patient organisations from more than 60 countries that work together to improve the lives of the 30 million people living with a rare disease in Europe.

> By connecting patients, families and patient groups, as well as by bringing together all stakeholders and mobilising the rare disease community, EURORDIS strengthens the patient voice and shapes research, policies and patient services.

## **RARE DISEASES**

**OVER** 6000 distinct rare diseases

Affects between

of the population

in the course of

their lives

6 % AND

Each one affects fewer than 1 IN **PEOPLE** 

for the vast majority of

available

few treatments

All together, an estimated

diseases and

are living with a rare disease in Europe and 300 **MILLION** worldwide

MILLION PEOPLE







## **EMPOWERING PATIENTS**

By uniting the rare disease community and building the capacities of patients, we empower them to become advocates equipped with the knowledge and skills needed to fight for better lives.



### **ADVOCATING FOR PATIENTS**

We advocate to ensure that healthcare and social policies and services, research, and the development of medicines take into account the real

needs of people living with a rare disease and their families. Our advocacy is fuelled by real-life experiences of the rare disease community.









## **ENGAGING PATIENTS**

We make possible the engagement of patients in decision-making processes so that the patient voice is active, amplified and meaningful in research and medicines development, as well as in both healthcare and social policies and services.

**OUR INITIATIVES** 



Expert Patient and Researcher **EURORDIS Summer School** 



**EURORDIS** survey initiative



The online network for rare disease patients & families



The European Conference on Rare Diseases & Orphan Products



The global campaign to raise awareness of rare diseases





The global alliance of people living with a rare disease

Making rare diseases a global public health priority