[Press release]

Stars of the rare disease community recognised at 2019 EURORDIS Black Pearl Awards Ceremony

#EURORDIS2019 #BlackPearl

12 February 2019, Brussels – The recipients of the EURORDIS Black Pearl Awards 2019 are being presented with their awards at a ceremony held this evening in Brussels.

Watch online via blackpearl.eurordis.org/live

EURORDIS Young Patient Advocate: Laëtitia Ouillade, France
EURORDIS European Rare Disease Leadership Award: Prof. Till Voigtländer, Austria
EURORDIS Policy Maker Award: Dr Edmund Jessop, UK
EURORDIS Scientific Award: Prof. Philip Van Damme, Belgium

EURORDIS Volunteer Awards: Richard West, Behçet's Syndrome Society, UK
Russell Wheeler, Leber's Hereditary Optic Neuropathy (LHON) Society, UK

EURORDIS Members Award: ALS Liga Belgium
EURORDIS Company Award for Innovation: Chiesi, Italy
EURORDIS Company Award for Patient Engagement: The HERCULES Project (Pfizer Inc, PTC Therapeutics, Roche, Sarepta Therapeutics, Solid Biosciences, Summit Therapeutics, Wave Life Sciences, Duchenne UK)

EURORDIS Company Award for Health Technology: Air Liquide Medical Systems, France
EURORDIS Visual and Audio Media Award: ‘Et Les Mistrals Gagnants’, Anne-Dauphine Julliand, France
EURORDIS Written Media Award: ‘Word for Life’, Bojana Mirosvljević, Serbia

EURORDIS Lifetime Achievement Award: Michael Griffith, Ireland

The EURORDIS Black Pearl Awards Ceremony takes place in February to mark the occasion of Rare Disease Day.

The Black Pearl Awards recognise the major achievements and outstanding commitment of patient advocates, patient organisations, policy makers, scientists, companies and media who strive to make a difference for the rare disease community.

There were hundreds of nominations from over 37 different countries worldwide.

The EURORDIS Photo Award

The EURORDIS Photo Award is open to everyone around the world. It is an opportunity to visually express the reality of living with a rare disease, the joy and the challenges. After online submissions closed on 7 January, world-renowned National Geographic photojournalist Marcus Bleasdale shortlisted five photos for the public vote.

The winner will be announced live (watch online via blackpearl.eurordis.org/live) at the EURORDIS Black Pearl Awards this evening. The three finalists with the most votes will each receive a prize.
EURORDIS-Rare Diseases Europe

EURORDIS-Rare Diseases Europe is a unique, non-profit alliance of over 800 rare disease patient organisations from more than 70 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. Follow @eurordis or see the EURORDIS Facebook page. For more information, visit eurordis.org.

Rare diseases

The European Union considers a disease as rare when it affects less than 1 in 2,000 citizens. Over 6,000 different rare diseases have been identified to date, affecting an estimated 30 million people in Europe and 300 million worldwide. Due to the low prevalence of each disease, medical expertise is rare, knowledge is scarce, care offering inadequate and research limited. Despite their great overall number, rare disease patients are the orphans of health systems, often denied diagnosis, treatment and the benefits of research.

Press contact

Eva Bearryman, Communications Manager
EURORDIS-Rare Diseases Europe
eva.bearryman@eurordis.org
+33 1 56 53 52 61