RD-Connect is a unique global infrastructure project that links databases, registries, biobanks, and clinical bioinformatics data used in rare disease research into a central resource for researchers worldwide. In a six-year project funded by the European Union but uniting researchers across the world, it will develop an integrated research platform in which complete clinical profiles are combined with “-omics” data and sample availability for rare disease research, in particular research funded under the International Rare Diseases Research Consortium (IRDiRC).

Mission

RD-Connect’s primary objectives are as follows:

Structure & Governance

RD-Connect is led by coordinator Hanns Lochmüller of Newcastle University.
The project’s Governing Board, consisting of one representative from each partner institution, is responsible for setting annual objectives and policy in accordance with the rules of the EC contract. The Governing Board provides legal and strategic oversight and ratifies any necessary changes to the consortium and workplan.

For operational management, the coordinator is supported by the Executive Management Committee and the coordination team in Newcastle.

**Patent Engagement**

The Patient Advisory Council (PAC) examines ethical, social, and participatory questions linked to research taking place in the context of RD-Connect, NeurOmics, and EURenOmics. This will ensure that all project activities have a patient-centric approach throughout. EURORDIS coordinates activities in respective projects with the PAC as well as invites its patient representatives to meetings and discussions.

Issues, concerns, or dilemmas brought forth by the PAC are addressed by the Rare Disease Patient and Ethics Council (RD-PEC) as well as all project partners. The PAC is composed of a multidisciplinary group of patients, parents, representatives of patients’ organizations, clinicians, legal academics, sociologists, scientists, and ethicists.

**Data Sharing**

All data- and material-sharing agreements should be ethically robust and mindful of the responsibilities owed to the donors to make best ethical use of the samples and data consistent with their consent. The international charter of principles for sharing biospecimens and data provides guidance for effective legally and ethically grounded sharing and can be found on the RD-Connect website.

The charter is inspired by the following principles:
Points of Contact

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Sponsors & Partners

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Association Francaise contre les Myopathies
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Centro Nacional de Investigaciones Oncologicas
Charles University
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Consiglio Nazionale delle Ricerche
EuroBioBank
European Bioinformatics Institute
European HD Network
European Organisation for Rare Diseases
Finovatis
Fondation maladies rares
Fondazione Telethon
Foundation for Research and Technology – Hellas
Genzyme Europe BV
Glaxo SmithKline
Innolyst, Inc. Patient Crossroads
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University of Belgrade
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LYSOGENE
Medical University of Graz
Murdoch University
National Center for Neurology & Psychiatry
National Institute of Environmental Health
National Institutes of Health, Office of Rare Diseases Research
NGRL Manchester
Office of Population Health Genomics
Orphan Europe Recordati Group
Orphanet
PhenoTips
Telethon Network of Genetic Biobanks
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University of Aveiro
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