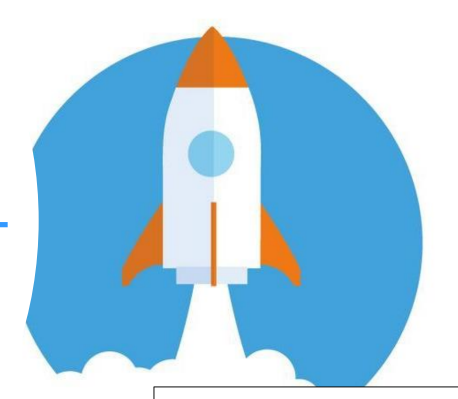






LAUNCH OF THE FIRST EUROPEAN REGISTRY FOR ALL NEUROMUSCULAR DISEASES



EURO-NMD Registry Consortium:

- Assistance Publique–Hôpitaux de Paris (AP-HP)
- University Medical Centre Freiburg (UKLFR)
- Radboud University Medical Center (Radboudumc)
- World Duchenne Organization (WDO)
- Duchenne Data Foundation (DDF)
- Institute of Myology (AIM)
- French Muscular Dystrophy Association (AFM-Téléthon)

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REVIEW Open Access



EURO-NMD registry: federated FAIR infrastructure, innovative technologies and concepts of a patient-centred registry for rare neuromuscular disorders

Antonio Atalaia^{1*†}, Dagmar Wandrei^{2†}, Nawel Lalout^{3,4}, Rachel Thompson⁵, Adrian Tassoni², Peter A. C. 't Hoen³, Dimitrios Athanasiou⁶, Suzie-Ann Baker⁶, Paraskevi Sakellariou⁷, Georgios Paliouras⁷, Carla D'Angelo⁸, Rita Horvath⁹, Michelangelo Mancuso¹⁰, Nadine van der Beek¹¹, Cornelia Kornblum¹², Janbernd Kirschner¹³, Davide Pareyson¹⁴, Guillaume Bassez¹⁵, Laura Blacas¹⁶, Maxime Jacoupy¹⁶, Catherine Eng¹⁷, François Lamy¹⁷, Jean-Philippe Plançon¹⁸, Jana Haberlova¹⁹, Esther Brusse¹¹, Janneke G. J. Hoeijmakers²⁰, Marianne de Visser²¹, Kristl G. Claeys²², Carmen Paradas²³, Antonio Toscano²⁴, Vincenzo Silani²⁵, Melinda Gyenge¹⁵, Evy Reviers²⁶, Dalil Hamroun²⁷, Elisabeth Vroom⁶, Mark D. Wilkinson²⁸, Hanns Lochmuller^{5,13} and Teresinha Evangelista²⁹





HOW TO GET STARTED?

01

SIGN

THE DATA SHARING
AGREEMENT (ACCESSION
FORM ATTACHED AS
APPENDIX 8 TO THE DSA)

02

TRANSLATE

THE ENGLISH VERSION OF THE PATIENT INFORMATION AND CONSENT FORM (APPENDIX 3 TO THE DSA) 03

ASSESS

IF ETHICAL/ GOVERNANCE APPROVAL IS REQUIRED

OBTAIN

NECESSARY APPROVALS

04

RETURN

SIGNED DSA AND
TRANSLATED ICF TO
COORDINATION TO INITIATE
CENTRE'S ENROLLMENT





HCP ENROLLMENT

▶ Sign-up process:

- activated by the HCP Rep (ERN Board representative)
- ► HCP rep will nominate the registry users & access rights and roles
- ▶ HCP rep will indicate the need (or not) to create different "DATA ACCESS GROUPS" for units or sites

▶ Definitions:

- > HCPs = legal entities that have applied to become members of the ERN (individual hospitals or hospital consortia) -> Goal: onboard 82 ERN HCPs
- > An HCP may consist of several <u>UNITS</u> = services/departements belonging to the same hospital (e.g. paediatric and adult)
- > An HCP may act as a Consortium reporting on several <u>SITES</u> = different hospitals (legal entities) -> to access the registry, sites need to sign the DSA



REGISTRY ROLL OUT

Onboarding

- Enrolling an HCP into the registry happens on a rolling basis
- > Pilot roll out: one-to-one meetings with HCP teams to assist with the onboarding based on needs, capabilities and data management practices (EURO-NMD Registry helpdesk: registryhelpdesk.euronmd@outlook.com)

▶ Where we stand?

- ✓ 1st meeting with Radboudumc (8 Feb)
- Collab w/ Belgium NMD Registry (BNMDR) & French National Registry for RDs (in progress)
- Meeting with APHP planned (1 Mar)
- ✓ Meeting with Norwegian registry for hereditary neuromuscular diseases (NORNMD) (6 Mar)
- EOI from HCPs in Pisa (AOU Pisana) & Dublin (Beaumont Hospital)



DATA ENTRY INCENTIVE PILOT

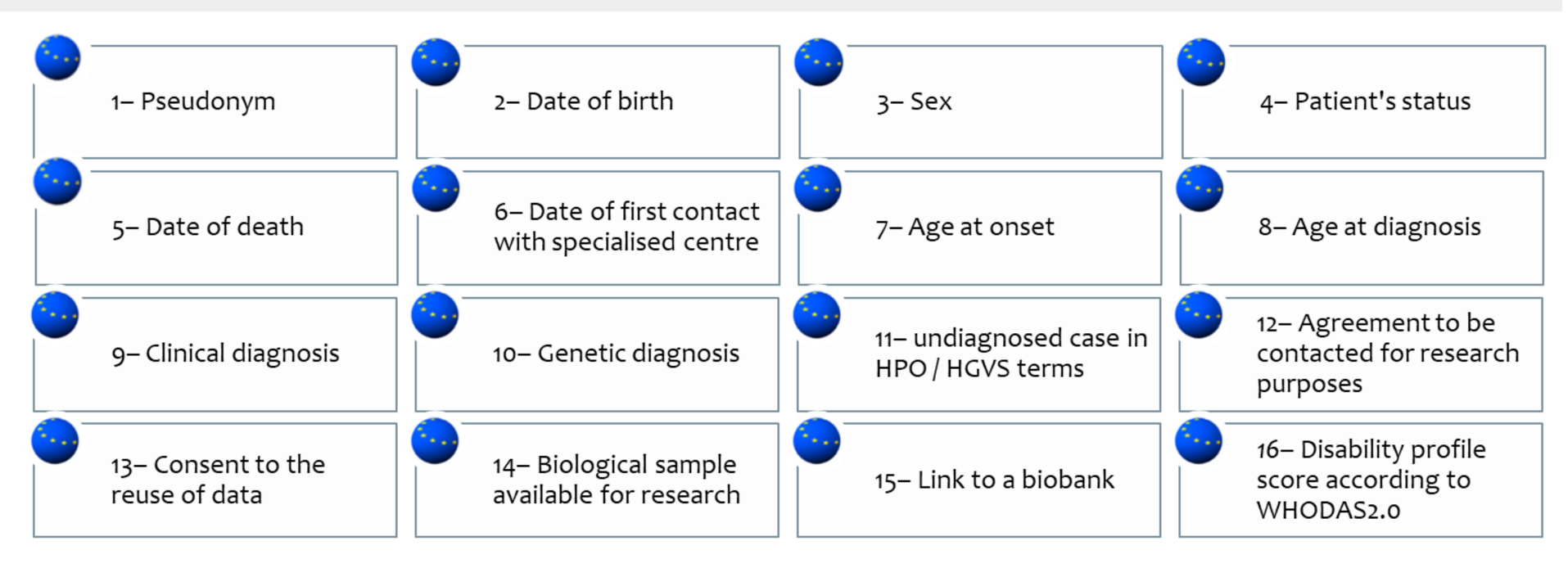
► Financial compensation 1 500 € per ERN HCP for the first 100 patients enrolled EURO-NMD 23-27 GA ≠101156434 (EUR 123 000)

▶ EURO-NMD Registry Reimbursement Protocol

- > ERN HCPs will need to sign a collaboration agreement with the HCP Network Coordinator (APHP)
- Each HCP receives a flat amount of 1 500 € upon receipt of invoice only if they have completed the minimum mandatory data set (next slide \rightarrow)
- Where a single HCP represents a consortium of different clinical sites, <u>only</u> the ERN HCP who submitted the Network Application (ERN Member) is eligible



Set of CDEs for RD Registration (EU RD Platform)







DATA SHARING AGREEMENT

WHAT?

- ▶ Relations between <u>Parties</u>: responsibilities (under GDPR), terms and conditions of participation, rights & obligation
- ► Each of the Parties are Joint Controllers: share responsibility for compliance with GDPR & are jointly liablility!!

WHO (Parties to the DSA)?

- ► EURO-NMD Registry Consortium
- ▶ Registry coordinator: APHP is the coordinator of the EURO-NMD Registry as well as a data provider
- Registry host: registry is hosted by the University Medical Centre Freiburg (UKLFR), which is also a data provider
- HCPs participating in the Registry (data providers)



DATA SHARING AGREEMENT

Specific responsibilities of HCPs as joint controllers (Appendix I)

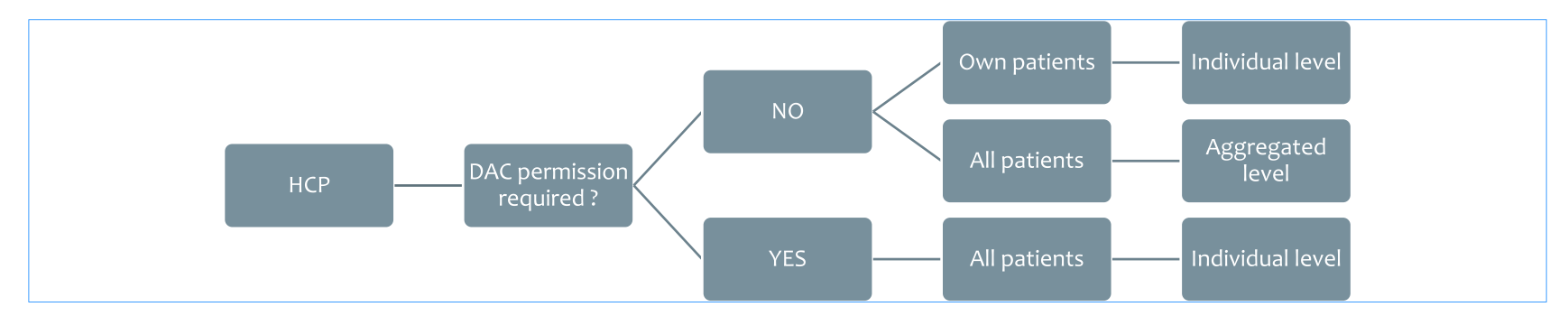
- 1. Implementation of security measures (pseudonymisation, users access)
- 2. Dissemination of the information to the Patients
- **3.** Collection of the consent for Patients
- 4. Recording of consent of Patients
- 5. Receiving and answering requests of Patients
- 6. Informing the other joint controllers of a Data Breach



DATA ACCESS POLICY (Appendix 6 to DSA)

WHAT?

- ► Third parties entitled to request data, data access authorization levels & procedures
 - > HCP agrees that HCP Data may be shared with third parties according to the conditions defined in the DAP
 - > Only aggregated (anonymised) or individual-level (pseudonymised) data will be shared by the Registry
 - > DAC approval is always required for access to individual-level (pseudonymised) data







AUTHORSHIP POLICY (Appendix 7 to DSA)

WHEN?

- ► Any form of publication produced using data from the Registry
- ► EXCEPTION: contributing hospitals are free to publish their own data (with acknowledgement of the Registry)

Authorship criteria:

- ▶ Will be granted to at least one representative from HCPs that have contributed with data for the study
- ▶ Number of co-authors will depend on the size of the study and number of HCPs:
 - ▶ the criteria for defining the number of co-authors per HCP will be the number of included patients
 - ▶ In special cases, only centres that contributed data for substantial numbers of patients meeting the eligibility criteria for a study will be invited to become co-author

PARTICIPATION REQUIREMENTS

To ensure the success of the Registry, HCPs must:

- ensure adequate resources and accurate information, including:
 - named staff who will have access to the EURO-NMD Registry
 - > contact person who will be responsible for all registry-related communications
- ▶ inform patients about the Registry and obtain patient informed consent
- ► For a dataset to be considered complete:
 - collect the mandatory set of data known as the Common Data Elements (CDEs)
- complete follow-up questionnaires annually after the baseline visit
- assist with data quality assurance processes









