Procedural document
Inventory of genes related to rare diseases

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I. Introduction

1. Purpose/objectives

Orphanet has developed and maintains the Orphanet nomenclature of rare diseases (RDs), a unique and multilingual standardised system aimed at providing a specific terminology for RDs. Each clinical entity within the Orphanet nomenclature of RDs is enriched by scientific and textual information. In order to better define RDs of genetic origin, Orphanet provides information on genes related to a RD. This information is structured around the international nomenclature of genes, provided by the HUGO Gene Nomenclature Committee (HGNC), and comprises the gene typology, the chromosomal location, cross-mappings with international genetic databases and semantic relationships between genes and their related RDs, along with the evidence-based publications supporting this relationship.

This procedural document describes the production, validation and update process of the Orphanet inventory of genes related to RDs.

2. Range of application

The present procedure applies to all genes listed in the Orphanet database.

The Orphanet inventory of genes is managed by an information scientist with a genetic background, under responsibility of the Orphanet nomenclature manager. The Orphanet Advisory Board on Genetics and international experts are regularly consulted to ensure the accurate representation of the current knowledge. The Orphanet Advisory Board on Genetics is also consulted on this procedure.

3. Disclaimer

- This publication is part of the OrphaNetWork Direct Grant (831390) which has received funding from the European Union’s Health Programme (2014-2020).
- The content of this publication represents the views of the author only and is his/her sole responsibility; it can not be considered to reflect the views of the European Commission and/or the Consumers, Health, Agriculture and Food Executive Agency or any other body of the European Union. The European Commission and the Agency do not accept any responsibility for use that may be made of the information it contains.
- Candidate genes and biomarkers are excluded from the inventory unless a diagnostic test is made available in one of the Orphanet network’s countries.

4. References

For more information on the management of the inventory and classification of RDs, see the procedural document: The Orphanet nomenclature and classification of rare diseases.
5. Definitions

a. Orphanet gene-related dataset

<table>
<thead>
<tr>
<th>Gene dataset</th>
<th>Definition</th>
<th>Data coming from</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gene nomenclature</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gene name</td>
<td>HGNC-approved name</td>
<td>HGNC</td>
</tr>
<tr>
<td>Symbol</td>
<td>HGNC-approved symbol</td>
<td>HGNC</td>
</tr>
<tr>
<td>Synonyms</td>
<td>HGNC alias symbols and names</td>
<td>HGNC</td>
</tr>
<tr>
<td>Previous symbols and names</td>
<td>HGNC previous symbols and names</td>
<td>HGNC</td>
</tr>
<tr>
<td><strong>Gene information</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gene Type:</td>
<td>Basic unit of heredity, consisting of a segment of DNA arranged in a linear manner along a chromosome that is transcribed in RNA and translated in a protein.</td>
<td>HGNC</td>
</tr>
<tr>
<td>Disorder-associated locus</td>
<td>Chromosomal region associated with a single heritable disorder. The heritable disorder may be mapped to a chromosome but generally has not been associated to a specific gene.</td>
<td>HGNC</td>
</tr>
<tr>
<td>Non-coding RNA</td>
<td>RNA encoded by a gene but not translated in protein. ie: Transfer RNA.</td>
<td>HGNC</td>
</tr>
<tr>
<td>Chromosomal location</td>
<td>Cytogenetic location on the chromosome of a gene with protein product, non-coding RNA or disorder-associated locus.</td>
<td>HGNC</td>
</tr>
<tr>
<td><strong>Genetic database cross-mappings</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HGNC</td>
<td>Non-profit making body which is jointly funded by the US National Human Genome Research Institute (NHGRI) and the Wellcome Trust (UK). The HGNC is responsible for approving unique symbols and names for human loci.</td>
<td>Orphanet curation</td>
</tr>
<tr>
<td>OMIM</td>
<td>Online Mendelian Inheritance in Man is a compendium of human genes and genetic phenotypes.</td>
<td>HGNC</td>
</tr>
<tr>
<td>UniProtKB</td>
<td>Central hub for the collection of functional information on proteins, with accurate, consistent and rich annotation.</td>
<td>HGNC</td>
</tr>
<tr>
<td>Ensembl</td>
<td>An EBI database that maintains automatic annotation on selected eukaryotic genomes.</td>
<td>HGNC</td>
</tr>
<tr>
<td>IUPHAR</td>
<td>The International Union of Basic and Clinical Pharmacology portal.</td>
<td>HGNC</td>
</tr>
<tr>
<td>Reactome</td>
<td>An open-source, open access, manually curated and peer-reviewed pathway database.</td>
<td>HGNC</td>
</tr>
<tr>
<td>GenAtlas</td>
<td>A database of genes and phenotypes. Only the objects with a known cytogenetic location are retained. This cross reference is still present in Orphanet database but no longer updated.</td>
<td>HGNC</td>
</tr>
<tr>
<td>LOVD</td>
<td>Leiden Open Variation Database is an online gene-centered collection and display of DNA variants.</td>
<td>HGNC</td>
</tr>
</tbody>
</table>

1 Matches with Gene with protein product; immunoglobulin gene; T cell receptor gene; Protocadherin; Readthrough (from HGNC locus type)
2 Matches with RNA, long non-coding; Pseudogene; Virus integration site; Endogenous retrovirus; Immunoglobulin pseudogene; RNA, micro; RNA, ribosomal; RNA, transfer; RNA, cluster; RNA, misc; RNA, small nuclear; RNA, small nucleolar; RNA, small cytoplasmic; RNA, Y; T cell receptor pseudogene; RNA, vault; unknown (from HGNC locus type)
3 Matches with region; Complex locus constituent; Fragile site (from HGNC locus type)
<table>
<thead>
<tr>
<th>Gene dataset</th>
<th>Definition</th>
<th>Data coming from</th>
</tr>
</thead>
<tbody>
<tr>
<td>Disease-causing germline mutation(s) in</td>
<td>A gene mutation in a germ cell that is sufficient to produce the disorder and that can be passed on to offspring.</td>
<td>Orphanet curation</td>
</tr>
<tr>
<td>Disease-causing germline mutation(s) (gain of function) in</td>
<td>A gene mutation in a germ cell that provides a new function of the corresponding protein and that is sufficient to produce the disorder and that can be passed on to offspring.</td>
<td>Orphanet curation</td>
</tr>
<tr>
<td>Disease-causing germline mutation(s) (loss of function) in</td>
<td>A gene mutation in a germ cell that impairs the function of the corresponding protein and that is sufficient to produce the disorder and that can be passed on to offspring.</td>
<td>Orphanet curation</td>
</tr>
<tr>
<td>Disease-causing somatic mutation(s) in</td>
<td>A gene mutation in a somatic cell that is sufficient to produce the disorder but that cannot be passed on to offspring.</td>
<td>Orphanet curation</td>
</tr>
<tr>
<td>Major susceptibility factor in</td>
<td>A gene mutation in a germ cell that predisposes to the development of a disorder, and that is necessary but not sufficient to develop the disorder.</td>
<td>Orphanet curation</td>
</tr>
<tr>
<td>Modifying germline mutation in</td>
<td>A gene mutation in a germ cell that modifies the clinical presentation of the disorder and that can be passed on to offspring.</td>
<td>Orphanet curation</td>
</tr>
<tr>
<td>Part of a fusion gene in</td>
<td>A coding or regulatory DNA sequence from a gene that has fused with another coding and/or regulatory DNA sequence from a different gene.</td>
<td>Orphanet curation</td>
</tr>
<tr>
<td>Role in the phenotype of</td>
<td>A gene included in a chromosomal rearrangement of which the mutation results in a phenotype related to the given chromosomal rearrangement, therefore proving its influence in a particular manifestation.</td>
<td>Orphanet curation</td>
</tr>
<tr>
<td>Biomarker tested in</td>
<td>A gene in which a variation is used to monitor disease activity and/or patient outcome.</td>
<td>Orphanet curation</td>
</tr>
<tr>
<td>Candidate gene tested in</td>
<td>A gene in which a mutation is suspected, but not yet proven, to be responsible for a disorder, and that is tested for in a clinical setting.</td>
<td>Orphanet curation</td>
</tr>
<tr>
<td>Reference</td>
<td>Reference(s) for a given source associated with a disease-gene relationship.</td>
<td>Orphanet curation</td>
</tr>
<tr>
<td>Validation status</td>
<td>Indication that the curation is done accordingly to Orphanet procedures and is scientifically valid.</td>
<td>Orphanet curation</td>
</tr>
</tbody>
</table>

b. **Other definitions**

**Experts**: a medical doctor or researcher with prominent experience in a rare disease or a group of rare diseases, and identified by Orphanet based on published articles (particularly reviews and guidelines), involvement in expert centers, expert networks, and/or in dedicated research activities including clinical trials.

**Familial cases**: At least two patients in the same family affected by a rare disease suspected to be of genetic origin.

**Genetic functional study**: Genetic functional studies aim to explain the physiological and pathophysiological roles of a gene. The functional studies used to validate the association of a gene with a disorder can be carried out in animal models like knock-out, cell experiments such as restoration of a gene/protein activity and/or in vitro experiments like testing the gene expression level and/or protein activity.

**Information scientist**: a member of the Orphanet team with a scientific and/or medical background in charge of collecting, producing and updating information provided in the Orphanet database.

**Orphanet advisory board on genetics**: This board is composed either by geneticists, members of the Orphanet management board and joining the advisory board on a volunteer basis, or by geneticists not
belonging to the Orphanet management board and invited under proposal of the Orphanet Management Board. The membership, mandate and role of the Orphanet Advisory Board on Genetics is available here.

**Orphanet gene manager:** an information scientist of the Orphanet coordinating team in charge of updating gene information and their relationships to the Orphanet nomenclature of rare diseases.

**Orphanet nomenclature manager:** an information scientist of the Orphanet coordinating team in charge of producing and updating the nomenclature and classification of rare diseases.

**Orphanet national team:** An Orphanet team based in one of the member countries of the Orphanet Network as per the Orphanet Network Agreement, and responsible for the collection of data on national expert resources. Some of the national teams are also in charge of the translation of the Orphanet nomenclature into one of the languages of translation of the database (German, Italian, Spanish, Portuguese, Polish, Czech and Dutch).

**Orphanet Rare Disease Ontology (ORDO):** an open access, structured, and machine-readable vocabulary for rare diseases derived from the Orphanet database, capturing relationships between diseases, genes and other relevant features, and forming a useful resource for the computational analysis of rare diseases. ORDO was jointly developed by Orphanet and the European Bioinformatics Institute (EMBL-EBI).

**Orphadata:** a platform developed by Orphanet to provide the scientific community with comprehensive, high-quality and freely accessible datasets related to rare diseases and orphan drugs, in a reusable format.

**Panel of genes:** Collection of targeted genes thought to be relevant for particular diseases or conditions that are analysed together in a single diagnostic test. Genes present in a panel are usually linked by common biological pathways, or known disease associations.

**Penetrance:** the proportion of individuals that expresses an associated phenotype within a population carrying a particular variant of a gene.

**Rare disease:** a disease that affects less than five in 10,000 persons in Europe, as defined by the European Regulation on orphan medicinal products (Regulation (EC) No 141/2000 of the European Parliament and of the Council of 16 December 1999 on orphan medicinal products). In order to be registered in Orphanet, the disease must be described in at least two independent individuals, confirming that it is not an incidental association of clinical signs.

6. **Filing and updates**

This procedural document is updated annually and as often as necessary by the data manager. The most up-to-date version is available on the Orphanet website: [https://www.orpha.net/orphacom/cahiers/docs/GB/Orphanet_Genes_inventory_R1_Ann_gen_EP_02.pdf](https://www.orpha.net/orphacom/cahiers/docs/GB/Orphanet_Genes_inventory_R1_Ann_gen_EP_02.pdf)
II. Methodology

1. Flowchart

Figure 1: Flowchart of the procedure
Figure 2: Diagram of decision: gene-disease relationship for a germline point mutation in a gene

Procedural document on Orphanet inventory of genes related to rare diseases. June 2021 – Number 02.
Figure 3: Diagram of decision: gene-disease relationship for a somatic point mutation in a gene.

1. Somatic point mutation in a single gene
2. How many isolated cases?
   - Multiple isolated cases
     - Are there controls? (No)
       - Is the mutation present in the control/healthy tissue? (Yes)
         - Is there a positive functional study? (No)
           - Is it a biomarker? (No)
             - Is there a diagnostic test referenced in Orphanet? (No)
               - Link the gene with the relationship: Biomarker tested in
             - Is there a diagnostic test referenced in Orphanet? (Yes)
               - Link the gene with the relationship: causing somatic (assessed)
           - Is it a biomarker? (Yes)
             - Is there a diagnostic test referenced in Orphanet? (No)
               - Link the gene with the relationship: Biomarker tested in
             - Is there a diagnostic test referenced in Orphanet? (Yes)
               - Link the gene with the relationship: causing somatic (assessed)
   - One isolated case
     - Are there controls? (Yes)
       - Is the mutation present in the control/healthy tissue? (No)
         - Is there a positive functional study? (Yes)
           - Link the gene with the relationship: causing somatic (assessed)
         - Is there a positive functional study? (No)
           - Is it a biomarker? (No)
             - Is there a diagnostic test referenced in Orphanet? (No)
               - Do not link the gene.
Figure 4. Diagram of decision: gene-disease relationship for a chromosomal anomaly

Chromosomal anomaly

Chromosomal rearrangement/translocation resulting in a fusion gene*

How many isolated cases?

Several isolated cases

Are there controls?

No

Yes

Is the mutation present in the controls?

No

Yes

Link the gene with the relationship: fusion gene (assessed)

One isolated case

Is this anomaly identified as causal?

No

Yes

Does the described gene cause a similar phenotype in an isolated manner?

No

Yes

If there is a diagnostic test referenced in Orphanet, link the gene with the relationship tested candidate gene

Link the gene with the relationship: role in the phenotype (assessed)

*see definition part of the procedure
2. Description

The update process of the Orphanet inventory of genes related to RDs is subject to a validation workflow designed to ensure that the information and data provided by Orphanet have a satisfactory level of exhaustivity, accuracy, pertinence, and reliability.

a. Requests for modification of the Orphanet inventory of genes

Different sources point the need for creation or update of a gene entry:
- literature monitoring;
- requests from experts;
- requests from national information scientists of the Orphanet consortium;
- cross-referencing with medical and/or genetic databases (such as OMIM).

b. Analysis of scientific resources

Regarding every update request, the Orphanet gene manager gathers information through a review of the scientific literature in order to determine the most appropriate action in view of the current state of knowledge and the Orphanet database rules (see below). Only peer-reviewed publications are consulted. Decisions are based only on the publications establishing the gene-disease relationship.
As an exception, candidate genes and biomarkers can be recorded without peer-reviewed publication upon an expert demand regarding a diagnostic test in use.

c. Identification of the related rare disease

Based on the publication(s), the RD is identified in the Orphanet rare disease nomenclature. Orphanet rare diseases are clinically defined. Accordingly, the decision to establish a gene-disease relationship is clinically oriented. It can differ from other genetically defined database for which Orphanet provides cross-referencing in order to enhance interoperability of the dataset.
If needed, addition of a new disease to the Orphanet rare disease nomenclature is studied according to The Orphanet nomenclature and classification of rare diseases procedure.

d. Identification of the gene-disease relationship

The gene-disease relationship is qualified according to the pathogenesis of the described mutations. Thus, genes are annotated as causative (from germline or somatic mutations), modifiers, major susceptibility factors or playing a role in the phenotype (for chromosomal anomalies). When causative mutations are of germline origin, loss or gain of protein function are documented if available.

Two relationships are specifically dedicated to candidate genes and biomarkers, that are registered only if a diagnostic genetic test is made available in one of the Orphanet network’s countries.

Criteria to link a gene to a disease differ from a relation to another. Decisional diagrams (figure 2, 3 and 4), validated by the Orphanet Advisory Board on Genetics, help the data manager to standardise this step. However, the scientific knowledge couldn’t be easily represented in a simple diagram and number of the decisional steps depends on the data manager expertise.

Scientific validation is sometimes necessary to confirm the identification of the related disease and/or its gene
relationship. These particular cases are first submitted for scientific validation to the Orphanet nomenclature manager. If necessary, experts on a specific disease or members of the Orphanet Advisory Board on Genetics are also consulted.

The main decisional criteria are described below:

**Disease-causing germline mutation(s) in**
- The mutation segregates in one or more family(ies) or is identified in several isolated cases. Segregation of the putative pathogenic mutation is assessed case by case. The structure of the family, the mode of inheritance and the penetrance of the disease are carefully examined.
- The mutation is not found in the general population (absent from control cohort and/or genetic variants databases). In case of low penetrance, mutation’s occurrence in control is permitted. Then, causality is assessed according to the degree of penetrance, the frequencies of the mutation in patient’s and control’s populations. In that case, functional studies may help to confirm the relationship.
- If there is only one isolated case, functional studies are needed to confirm the association.
- If the inheritance is digenic (or oligogenic more generally), functional studies are needed to confirm the association.

Functional studies are assessed case by case.
If available, loss or gain of the protein function is documented.

**Disease-causing somatic mutation(s) in**
- The mosaic mutation is found in several patients.
- The mosaic mutation is absent from controls (persons or healthy tissues).
- Functional studies confirm the association.

Functional studies are assessed case by case.
If there is only one isolated case, the gene is not registered in the database.
Concerning cancers, the causality of the mutation is considered if the mutation is an early event in oncogenesis and if an *ex vivo* or *in vivo* study confirms its oncogenicity.

**Major susceptibility factor in**
- The variant segregates in a family or is present in several isolated cases.
- The variant is usually found in the general population and/or the disease is known to be multifactorial.
- If the variant has been identified by a Genome wide association study (Gwas), only meta-analysis and a p-value less than 5.10(-8) are considered.

Currently, there is no accepted consensus for other types of association studies (ie: Whole exome sequencing). Consequently, these studies are assessed case by case and gene is registered only if every evidence converged (replicated studies, functional studies…).
Isolated cases are not considered.

**Modifying germline mutation in**
A modifying germline mutation is considered only if the causative gene is already identified in the patients described in the study which fulfill the following criteria:
- The variant is present in several cases.
- The variant is absent from the general population or don’t lead alone to a similar phenotype.
- The consequences of the mutation are identical from a patient to another.

Isolated cases are not considered.
Part of a fusion gene in
Genes fused by chromosomal rearrangement are registered only if the molecular study show evidence of the causality of the fusion and if they fulfill the following criteria:

- The fusion is observed in several cases.
- The fusion is absent from controls.

Role in the phenotype of
A gene with a major role in the phenotype of a chromosomal rearrangement is linked to a disease only if:

- A mutation hitting only this gene gives rise to similar phenotypic consequences when it occurs in isolation.

Mutations or epimutations in an imprinting center are also considered as having a major role in a phenotype as the phenotypic consequences are similar to the one of a chromosomal rearrangement.

Candidate gene tested in or Biomarker tested in
Candidate genes or biomarkers are registered only if a genetic test is available in at least one of the Orphanet network’s countries.

Depending on the purpose of the test, either for diagnosis or to monitor disease activity, the candidate gene or biomarker relationship is attributed.

If any of these relationships can’t be implemented, the gene is not introduced in the Orphanet inventory of genes. As an exception, genes claimed in a Panel indicated for a diagnostic test can be registered in the Orphanet inventory of genes without any relationship to rare disease.

e. Update of the Orphanet database

When a request fullfills all the inclusion criteria, the gene is introduced in the Orphanet inventory of genes. Computational cross-reference with international genetic databases is completed and allows to inform on:

- Gene nomenclature, including main name, symbol, synonyms and previous names and symbols according to HGNC.
- Chromosomal location.
- Mappings with HGNC, OMIM, UniProtKB, Genatlas, Ensembl, Reactome and IUPHAR-DB, LOVD.

Every gene-disease relationships that have been curated and fulfill our inclusion criteria is mentioned as «assessed». Without this note, the gene-disease relationship is still waiting for expert validation.

Candidate genes are never «assessed» since there is no scientific evidence to attest the gene-disease relationship.

f. Quality control

To guarantee that the Orphanet genetic database is consistent with the rules and process described in this document, quality controls are performed regularly.

An automatic mapping with HGNC database is monthly performed to ensure the interoperability with other genetic international databases.
III. Availability of data

Orphanet inventory of gene is released at a variable frequency depending on the channel of dissemination, daily for the website (note that references are not displayed on the website) - www.orpha.net -, monthly for the Orphanet download platform - www.orphadata.org - and annually for the Orphanet Rare Disease Ontology – ORDO.