

BeSHG Interuniversity course

Yves Sznajer

Center for Human Genetic



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Development Genetics and Birth defects



Refer to Thompson / Thompson Chapter 14 7th Ed.

Framework

- Rationale in medical genetics
- Tool I: Registry Epidemiology Database
- Tool II: Definitions and Nosology
- Tool III: Distinct approaches
 - DysmorphologySyndromology





Rationale in medical genetic

Clinical geneticist seeks a consistent approach to lead to the diagnosis in a patient with birth defect

Aimsdiagnostic assessmentmore precise delineation on natural historyfollow-up and prognosisopen to precise genetic counselling

- Evidence derived from

Embryology: chronologic and stepwise mechanisms leading to ab-/normal development Epidemiology/Population Genetics Animal models Bioinformatics prediction Registries and Databases

Tool I - Congenital Anomalies Registry



Specific public health problem indicators **EUROCAT**

European network for the surveillance of congenitalanomaliestribute to Professor Yves Gillerot

Central Registry – Project Management Committee **Reliable, Available and Comparable on quantitative or qualitative parameters** - Dataset Registries

Tool I - Congenital Anomalies (CA) Registry

The Objectives of EUROCAT

- provide essential epidemiologic information on CA in Europe
- facilitate the early warning of new teratogenic exposures
- evaluate the effectiveness of primary prevention
- assess the impact of developments in prenatal screening
 act as an information and resource center for the population health
 professionals and managers regarding clusters or exposures or risk
 factors of concern
- -provide a ready collaborative network and infrastructure for research related to the causes and prevention of congenital anomalies and the treatment and care of affected children
- act as a catalyst for the setting up of registries throughout Europe collecting comparable and standardised data



http://ec.europa.eu/health/indicators/echu/index_eu.html Frontpage

eurecat

Member Registries

Information about the EUROCAT member regsitries, coverage and how to apply for membership



Data and surveillance

Prevalence tables, prenatal detection rates and reports on statistical monitoring of congenital anomalies









Prevention and risk factors

Research

EUROCAT Network

Overview of the EUROCAT network, its organisation and committees

European Commision link

← → C 🔒 publications.jrc.ec.europa.eu/repository/

🔘 An official website of the European Union 🛛 How do you know? 🗸



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publications.jrc.ec.europa.eu/repository/handle/JRC120236

European Commission > JRC > JRC Publications Repository > European Monitoring of Congenital Anomalies

2020 Technical reports Health and consumer protection

European Monitoring of Congenital Anomalies: JRC-EUROCAT Report on Statistical Monitoring of Congenital Anomalies (2008 - 2017)

Abstract: Worldwide, congenital anomalies are a leading cause of fetal death, infant mortality and morbidity in childhood. According to the EUROCAT estimates, of the 5.1 million births in the European Union (EU) each year approximately 127,000 (2.5%) have a congenital anomaly. EUROCAT is a European network of population-based registries whose objectives are to provide essential epidemiologic information on congenital anomalies in Europe, to facilitate the early warning of new teratogenic exposures and to evaluate the effectiveness of primary prevention. Each year, EUROCAT performs statistical monitoring for both trends and clusters in time on 84 anomaly subgroups. The results of the statistical monitoring are the basis for instigating possible further investigations at the local registry level. The present report shows the results of the monitoring performed on data for the birth years 2008-2017 by the JRC-EUROCAT Central Registry. Cases of congenital anomaly among livebirths, fetal deaths from 20 weeks gestational age and terminations of pregnancy for fetal anomaly following prenatal diagnosis at any gestational age were included. We report both the statistical results and, where available, the outcome of the preliminary investigations conducted by registries.



eurocai

european surveillance of

http://www.eurocat-network.eu

EUROCAT Prevalence Data Tables

Concentral anomalies Cases and prevalence (per 10,000 births) of all congenital anomaly subgroups for all registries, from 2011 - 2015

						Excluding Gen	etic Conditions
Anomaly	LB N	FD N	TOPFA N	LB+FD+TOPFA N	LB+FD+TOPFA Rate	LB+FD+TOPFA N	LB+FD+TOPFA Rate
All Anomalies	59179	1433	14619	75231	253.31 (251.51 - 255.13)	60392	203.35 (201.73 - 204.98)
Nervous system	3402	270	4040	7712	25.97 (25.39 - 26.55)	6558	22.08 (21.55 - 22.62)
Neural Tube Defects	621	102	2334	3057	10.29 (9.93 - 10.66)	2862	9.64 (9.29 - 10.00)
Anencephalus and similar	67	58	1074	1199	4.04 (3.81 - 4.27)	1155	3.89 (3.67 - 4.12)
Encephalocele	82	8	249	339	1.14 (1.02 - 1.27)	291	0.98 (0.87 - 1.10)
Spina Bifida	472	36	1011	1519	5.11 (4.86 - 5.38)	1416	4.77 (4.52 - 5.02)
Hydrocephalus	815	66	711	1592	5.36 (5.10 - 5.63)	1327	4.47 (4.23 - 4.72)
Microcephaly	658	36	97	791	2.68 (2.49 - 2.87)	640	2.16 (2.00 - 2.34)
Arhinencephaly/holoprosencephaly	67	17	364	448	1.51 (1.37 - 1.65)	271	0.91 (0.81 - 1.03)
Еуе	1036	15	99	1150	3.87 (3.65 - 4.10)	935	3.15 (2.95 - 3.36)
Anophthalmos/micropthalmos	195	8	61	264	0.89 (0.78 - 1.00)	188	0.63 (0.55 - 0.73)
Anophthalmos	35	2	28	65	0.22 (0.17 - 0.28)	53	0.18 (0.13 - 0.23)
Congenital cataract	361	1	4	366	1.23 (1.11 - 1.37)	312	1.05 (0.94 - 1.17)
Congenital glaucoma	95	0	0	95	0.32 (0.26 - 0.39)	89	0.30 (0.24 - 0.37)
Ear, face and neck	420	21	84	525	1.77 (1.62 - 1.93)	409	1.38 (1.25 - 1.52)
Anotia	55	0	4	59	0.20 (0.15 - 0.26)	53	0.18 (0.13 - 0.23)
Congenital heart defects	19889	380	2440	22709	76.46 (75.47 - 77.47)	19309	65.02 (64.10 - 65.94)
Severe CHD §	5267	193	1489	6949	23.40 (22.85 - 23.95)	5523	18.60 (18.11 - 19.09)
Common arterial truncus	141	9	60	210	0.71 (0.61 - 0.81)	151	0.51 (0.43 - 0.60)
Double outlet right ventricle §	327	19	116	462	1.56 (1.42 - 1.70)	387	1.30 (1.18 - 1.44)
Transposition of great vessels	840	15	136	991	3.34 (3.13 - 3.55)	953	3.21 (3.01 - 3.42)
Single ventricle	123	5	111	239	0.80 (0.71 - 0.91)	210	0.71 (0.61 - 0.81)
Ventricular septal defect	10098	123	637	10858	36.56 (35.88 - 37.25)	9543	32.13 (31.49 - 32.78)
Atrial septal defect	4587	30	100	4717	15.88 (15.43 - 16.34)	3983	13.41 (13.00 - 13.83)
Atrioventricular septal defect	913	55	381	1349	4.54 (4.30 - 4.79)	612	2.06 (1.90 - 2.23)
Tetralogy of Fallot	825	22	181	1028	3.46 (3.25 - 3.68)	825	2.78 (2.59 - 2.97)
Tricuspid atresia and stenosis	133	10	70	213	0.72 (0.62 - 0.82)	198	0.67 (0.58 - 0.77)
Ebstein's anomaly	108	9	14	131	0.44 (0.37 - 0.52)	124	0.42 (0.35 - 0.50)
Pulmonary valve stenosis	1140	6	49	1195	4.02 (3.80 - 4.26)	1073	3.61 (3.40 - 3.84)
Pulmonary valve atresia	218	1	73	292	0.98 (0.87 - 1.10)	266	0.90 (0.79 - 1.01)
Aortic valve atresia/stenosis §	364	7	66	437	1.47 (1.34 - 1.62)	393	1.32 (1.20 - 1.46)
Mitral valve anomalies	337	5	55	397	1.34 (1.21 - 1.47)	351	1.18 (1.06 - 1.31)
Hypoplastic left heart	393	36	381	810	2.73 (2.54 - 2.92)	710	2.39 (2.22 - 2.57)
Hypoplastic right heart §	86	9	67	162	0.55 (0.46 - 0.64)	153	0.52 (0.44 - 0.60)

 LB = Live Births

 EUROCAT Website Database: http://www.eurocat-network.eu/ACCESSPREVALENCEDATA/PrevalenceTables (data uploaded 07/04/2017)

 University of Ulster, 2012

 FD = Fetal Deaths / Still Births from 20 weeks gestation

 Source: Copyright: University of Ulster, 2012

TOPFA = Termination of pregnancy for fetal anomaly following prenatal

7

diagnosis - - = Data not available

§ = Incomplete or missing specification of ICD 9 codes



European Commission > EU Science Hub > European Platform on Rare Disease Registration > EUROCAT > EUROCAT data > Prevalence

Prevalence charts and tables

			_		
Country/Registry	Anomaly	Years		Case with genetic conditions	How to read the data
All full registries	All Anomalies	2011 to 2017		Include	Export the raw data

https://eu-rd-platform.jrc.ec.europa.eu/eurocat/eurocat-data/prevalence

Selected categories

Country/Registry All full registries	Anomaly All Anomalies		Years 2011 to 2017	
 All countries and regis All registries All full registries All associate registries All past registries (P) Manual selection 	tries ; (A)	 Belgiun Antw Antw Haina Denmar Denmar Oder France France Auve Auve France Auve Auve France Auve Auve Auve France Auve France Auve Auve France Auve <	n verp aut k se ergne any ch West Indies de la Reunion he-Alps (A) ral East France (P sbourg (P)	 Emilia Romagna Tuscany Campania (P) North East Italy (P) Sicily (P) eland Cork and Kerry Dublin SE Ireland Galway (P) L N Netherlands

Example of extractable data



Another example – Selected categories

Chromosomal	43.58 (42.99 - 44.17)	16.67 (16.30 - 17.04)	1.54 (1.43 - 1.66)	25.37 (24.92 - 25.83)
- Down Syndrome	24.14 (23.71 - 24.59)	9.74 (9.46 - 10.03)	0.50 (0.44 - 0.56)	13.91 (13.57 - 14.24)
 Patau syndrome/trisomy 13 	2.18 (2.05 - 2.31)	0.30 (0.25 - 0.35)	0.12 (0.09 - 0.15)	1.76 (1.64 - 1.88)
 Edward syndrome/trisomy 18 	5.94 (5.72 - 6.16)	0.69 (0.62 - 0.77)	0.45 (0.39 - 0.52)	4.79 (4.59 - 4.99)
- Turner syndrome	2.48 (2.34 - 2.63)	0.57 (0.51 - 0.65)	0.16 (0.12 - 0.20)	1.75 (1.64 - 1.87)
 Klinefelter syndrome 	0.65 (0.58 - 0.73)	0.44 (0.38 - 0.50)	0.01 (0.00 - 0.03)	0.20 (0.16 - 0.24)

Tool I - Congenital Anomalies Registry(ct'd)



Networks of action for 'Rare diseases'*

- have been supported under the Program for Community Action on Rare Diseases in 1999-2003; the EU Public Health Program 2003-2007 and the second EU Health Program 2008-2013

* Rare diseases define conditions with a prevalence <1/2.000 livebirths

The aims

- improve exchange of information via existing European networks
- promote better classification, develop strategies and mechanisms for exchanging information between people affected by a rare disease volunteers and professionals
- define relevant health indicators
- develop comparable epidemiologic data at EU level
- support and exchange of best practice develop measures for patient groups
- network and infrastructure for research into the causes and prevention of congenital anomalies and treatment/ care of affected children
- survey policies and practices with regard to periconceptional folic
 CLouvain
 acid supplementation

Rare diseases - definition



Rare diseases impact

- >80% with genetic origin
- Early onset , lifelong burden
- Difficult to diagnose and treat
- Often multisystemic alter QOL
- Impact on social environment
- Fragmented support and research

Rare diseases - basics



Manolio et al. Nature 461, 2009

Rare diseases – not limited to a concept



Kidney International (2017) 92, 796-808;

Belgisch plan voor zeldzame ziekten



Plan belge Maladies rares / Zeldzame ziekten

TABLE DES MATIERES

Introduction

Domaine 1: Diagnostic et information au patient
Action 1: Remboursement des tests nécessaires au diagnostic et au suivi des maladies rares, effectués en Belgique ou à l'étranger
Action 2: Système de qualité au sein des centres de génétique humaine
Action 3: Introduction d'une consultation de conseil génétique dans les centres d'expertise pour maladies rares
Action 4: Consultation multidisciplinaire
Action 5: Communication centrée sur le patient
Action 6: Europlan

Domaine 2: Optimalisation des soins

Action 7: Concentration de l'expertise et renforcement des centres de référence pour pathologies rares spécifiques existants : introduction d'un coordinateur de soins Action 8: Création de centres d'expertise pour l'hémophilie Action 9: Fonction maladies rares Action 10: Réseaux maladies rares Action 11: Nouveaux centres d'expertise Action 12 : Alimentation médicale pour maladies rares Action 13: Communication rapide des besoins médicaux : utilisation du dossier patient multidisciplinaire informatisé Action 14: Unmet medical need Action 15: Inventaire des besoins non couverts

Domaine 3: Connaissances et information Action 16: Registre central des maladies rares Action 17: Orphanet Belgium Action 18: Formation des prestataires de soins Action 19: Codification et terminologie

Domaine 4: Gouvernance et durabilité Action 20: Evaluation et monitoring du Plan

Rare diseases : information sources

https://www.orpha.net/consor/cgi-bin/index.php



... informations



Accueil > Associations de patients > Associations

Rechercher une association de patients

A	SS	50	CI	a	tI	0	n	s

Fédérations/Alliances

Lignes directes d'information

Nom de maladie		Chercher
) Champ obligatoire		
Tous les pays	~	Autre(s) option(s) de recherche 🔺
		> Chercher par nom d'association

Aide

Lata collection and registration of patient organisations in Orphanet

Orphanet fournit des informations sur les associations de patients, les regroupements d'associations et les alliances dédiées à une maladie rare ou à un groupe de maladies rares.

Entrez le nom de la maladie recherchée pour accéder à cette information.

Les résultats peuvent être triés géographiquement (par pays, région et ville, par ordre alphabétique) ou par pertinence (les résultats spécifiques de la maladie apparaissent en premier, ceux relatifs aux groupes de maladies en second).



Patient support

EDITORIAL

Patient Organizations and Research on Rare Diseases

Julie R. Ingelfinger, M.D., and Jeffrey M. Drazen, M.D. N Engl J Med 2011; 364:1670-1671 April 28, 2011 DOI: 10.1056/NEJMe1102290





orphanhealthcare

Foundation for Rare Disease



The scientific knowledge sharing and crowdfunding platform for rare diseases















BASIC PRE-CLINICAL TRANSLATIONAL & SOCIAL

Written by the members of the working group PENREP Guide first published in July 2020 on www.ejprarediseases.org

Patient Engagement in Biomedical Research Projects





Portail - http://www.institutdesmaladiesrares.be



Bienvenue sur le site de l'Institut des Maladies rares des Cliniques universitaires Saint-Luc !

ine multidie est dire tare longu ville alleste mains de 1 personne sur 2000. Entre 6000 à 8000 maladies tares sont identifiées. En Belgique, environ 700 000 personnes en sont atteintes. Souvent, les patients et leur médecin traitant se enrouvent démunis face aux pathologies et au



- Affections pancréatiques de l'enfant
- Malformations vasculaires (angiomes)
- Centre labio-palatin Albert de Coninck
- Hémopathies cellulaires héréditaires ou congénitales
- Maladies autoimmunes et autoinflammatoires de l'enfant
- Maladies cardiagues rares
- Maladies endocriniennes rares
- Maladies hépatiques de l'enfant
- Maladies neuro-cutanées congénitales
- Maladies neurogénétiques et neurodégénératives de l'enfant
- Maladies rhumatismales systémiques
- Pneumopathies infiltrantes diffuses





- Spécifiques aux maladies rares
- Hémophilie
- Maladies métaboliques héréditaires
- Maladies neuromusculaires
- Maladies rénales rares
 Mucoviscidose
- Non spécifiques aux maladies rares
 - Epilepsie réfractaire
- Infirmité motrice cérébrale (IMOC)
- Spina bifida
- Troubles du spectre autistique



Centre de génétique



Chiari type 1

- Dysplasie fibromusculaire
- Lupus Clinic
- Malformations vasculaires complexes et/ou syndromiques
- Naevus géant
- Ostéogénèse imparfaite (maladie des os de verre)
- Pathologies malformatives de l'œsophage
- Pied de Charcot
- Sclérose systémique
- Sclérose tubéreuse de Bourneville
- Syndrome de Marfan
- Syndrome de microdélétion 22
- Syndrome de Prader-Willi
- Syndrome de Williams
- Trisomie 21

Laboratoire de référence Screening prénatal

> 20.000 patients

Rare diseases: european umbrella



>2016: 24 ERNS involving 25 European countries, over 300 hospitals with over 900 healthcare units and covering all major disease groups. <u>http://ec.europa.eu/health/ern/policy_en</u>







EUROPEAN REFERENCE NETWORKS							
Share. C	are. Cure.						
े	European Reference Networks						

European Commission Heolth

ERN – Cliniques Saint-Luc

ERN confirmés en 201	14 & 2017	
EuroBloodNet	haematological diseases	Cédric Hermans
Endo-ERN	endocrine conditions	Dominique Maiter
ERKNet	kidney diseases	Olivier Devuyst
EURO-NMD	neuromuscular diseases	Peter Van den Bergh
RARE-LIVER	hepatological diseases	Etienne Sokal
ReCONNET	connective tissue and musculoskeletal diseases	Frédéric Houssiau
TRANSPLANT-CHILD	transplantation in children	Etienne Sokal
MetabERN	hereditary metabolic disorders	Marie-Cécile Nassogne
VASCERN	multisystemic vascular diseases	Laurence Boon

Candidatures 2019

ERN CRANIO	craniofacial anomalies and ENT disorders	Chr Raftopoulos	Geraldo Vaz
ERN EpiCARE	epilepsies	Roberta Cilio	Nicolas Gaspard
ERN EURACAN	adult cancers (solid tumours)	Jean-Pascal Machiels	Xavier Geets
ERN ITHACA	congenital malformations and rare intellectual disability	Yves Sznajer	Isabelle Maystadt
ERN LUNG	respiratory diseases	Antoine Froidure	Sophie Gohy
ERN PAEDCAN	paediatric cancer (haemato-oncology)	Bénédicte Brichard	Ann Van Damme
ERN RITA	immunodeficiency, autoinflammatory and autoimmune diseases	Cécile Boulanger	Bénédicte Brichard
ERNICA	inherited and congenital anomalies	Isabelle Scheers	Pierre Deprez



Tool I - Congenital Anomalies Registry (ct'd)

Networks of action for Rare diseases

- The EU EUROCAT project surveillance of congenital anomalies in Europe is coordinated by the University of Ulster Northern Ireland
- European network of **51 registries in 28 countries**
- > 1 million births/year in Europe are surveyed



Birth defect - congenital anomaly

Prevalence – 3%

Genetically determined highly penetrant

Monogenic subtypes

Incompletely penetrant

Polygenic

Multifactorial

Environmental

Environmental effect

Genetic load

Birth defect – 'main categories'



Tool II - Birth defect - Nosology

• <u>Deformation</u> consequence of extrinsic factors that modify/alter physical devlpt (histology nl)

• <u>Disruption</u> destruction of irreplaceable fetal tissue (vascular, trauma, teratogen)

• <u>Dysplasia</u> abnormal shape, structure, composition and/ or histology during devlpt ectoderm structure and derivates leading to a wide range of phenotype bone/skelettal

 Malformation result from intrinsic abnormalities in one or more genetic program operating during development (e.a polydactyly)

Congenital malformation, sequence, syndrome, deformation, 'disruption'



Box figure 14.1 – Clinical photographs of the main types of dysmorphic features.

(a) Cleft lip, a malformation representing failure of fusion of components of upper lip.
(b) Meningomyelocele, talipes and hydrocephalus, a malformation sequence due to failure of closure of the neural tube and consequent effects. (c) Trisomy 13, a baby with a malformation syndrome consisting of holoprosencephaly, midline cleft lip and palate, polydactyly and heart defects. (d) Talipes, abnormal position of the feet, a deformation due to extreme lack of liquor *in utero*. (e) Amniotic bands, disruption of a normal hand by constriction with strands of amnion leading to amputation and secondary fusion of finger tips (syndactyly). (f) Femur bones with multiple fractures and abnormal modeling due to osteogenesis imperfecta, a skeletal dysplasia.

Nosology (ct'd)

<u>Association</u>:

similar birth defects in different embryologic fields and inability so far to identify a genetic cause 'V.A.T.E.R', 'VACTER', 'VACTERL', cervico auriculo vertebral (coined 'Goldenhar' association)

• <u>Sequence</u>:



phenotype description of alteration of
structures inside an embryonic field during
development - pleiotropic: effect on a single
organ/system - precise moment
NOT a diagnosis ! 'Pierre Robin', Potter',

Nosology (ct'd)

• **Syndrome:** combination of birth defects that occur secondarily to cytogenetic and/or a gene anomaly

• **Spectrum**: monogenic or cytogenetic anomaly leading to a modification in a signaling pathway during devlpt - possibly responsible for a wide range of signs that may be overlooked as distinct



Holoprosencephaly Sonic Hedgehoc pathway - SHH; ZIC2; PATCH genes

Nosologie (ct'd)

 Association: malformation congénitale analogue similaire touchant différents territoires embryologiques sans cause identifiée/connue à ce jour
 exemples: V.A.T.E.R, VACTER, VACTERL

- **Syndrome**: combinaison d'anomalie(s congénitale(s secondaire(s à l'effet pléiotropique : une anomalie chromosomique et/ou génique

Tool III Clinical approach Dysmorphology

David Smith's contribution in 1966

- defines the clinical approach to look for distinct and/or minor signs in a patient to possibly obtain a diagnosis and orientate strategy to complete genotype definition – cytogenetic and/or molecular

- 'dysmorphologist' diagnoses a child with a birth defect, suggests appropriate work-up, guarantees follow-up and integrates pedigree and family history to published clinical reports to basic science literature



Path for reasoning







Dysmorphologist's 'textbook' "Elements of Morphology: Standard Terminology" John C. Carey*

International group of clinicians working in dysmorphology

Aims:

- initiate standardization of terms used to describe human morphology
- reach consensus regarding their definitions
- increase the utility of descriptions of the human phenotype
- facilitate reliable comparisons of findings among patients
- improve discussions with other related workers (pathologists, devlpt biology, molecular genetics) which will become more precise

Dysmorphology 'textbook' "Elements of Morphology: Standard Terminology" John C. Carey*

« These six articles provide **recommendations for the description and definition of human phenotypic variations** *... in the same way that the International Standing Committee on Human Cytogenetic Nomenclature accomplished this for human cytogenetics* [ISCN, 2005] the Nomenclature Working Group proposed the *description of human sequence variations* [den Dunnen and Antonarakis, 2001

AMERICAN JOURNAL OF Medical genetics A Standard Terminology

For Head and Face: pg 1-23



FIG. 1. An antero-posterior view of the cranium and face shows bony landmarks.



FIG. 2. A lateral view of the cranium and face shows bony landmarks.

AMERICAN JOURNAL OF medical genetics A 2009;149A(1):1-23 Standard Terminology

For Head and Face





FIG. 7. *Macrocephaly*. Note the increased size of the cranium. Differences in size are difficult to appreciate but increased head size in this child is notable because of comparison with the smaller face.

FIG. 8. *Microcephaly*. Decreased size of the cranium is accompanied by marked posterior sloping of the forehead.

AMERICAN JOURNAL OF
medical geneticsPART
A2009;149A(1):1-127Standard Terminology
For Peri orbital Region – pg 29-39





Eyebrows

Orientation

- high
- horizontal

Shape

- arched
- Heavy,





Distinctive

• Synophris, absence,...



External ears sometimes pathognomonic of a syndrome





From

'Face'

То

Gene'...

From Face to Gene ?...



http://www.face2gene.com Enhanced patient evaluation with Next-Generation Phenotyping

- The Genetics Resource
- Search for Syndromes
- **Review Photos & Features**
- Up-to-date Content through Genetics Community Curation

Ct'd





FORUMS

Collaborative Case Review for Diagnostic Dilemmas

START USING FACE2GENE (HTTP://APP.FACE2GENE.COM/FORUMS)

- ! Access the renowned <u>Expert Review Panel</u> (/forums-diagnostic-dilemmas/expert-reviewpanel/)
 - detect phenotypes
 - reveal relevant facial and non-facial features
 - review relevant syndrome matches
 - access Best-in-class Resources
 - give and Receive Clinical Feedback

A second database tool

Human Phenotype Ontology

希 Home	About	📥 Downloads	¢ ₿ Tools	Documentation	📽 Users	්ට History
😧 FAQ	© License	🗞 Citation 🛛 🖂	Contact			

This page is split into:

- Introduction
- Annotation guide
- Logical definitions

An Introduction to the Human Phenotype Ontology

The Human Phenotype Ontology (HPO) intends to offer a tool that will allow large-scale computational analysis of the human phenome. The HPO currently contains over 11,000 terms, each of which describes an individual phenotypic anomaly. The terms are arranged in a directed acyclic graph and are connected by **is-a** (subclass-of) edges, such that a term represents a more specific or limited instance of its parent term(s). All relationships in the HPO are **is-a**

Human Phenotype Ontology



HPO Browser

The HPO Browser has a separate page for every term in the HPO. The following page, for instance, is for the root term: http://www.human-phenotype-ontology.org/hpoweb?id=HP:0000118.

PhenExplorer: The PhenExplorer has been superceded by the new HPO browser, which has all the functionalities of PhenExplorer and has additional features such as Excel-Exports

Clinical diagnostics using the HPO

Phenomizer:

The Phenomizer is available at http://compbio.charite.de/phenomizer.

It is a web-based application for clinical diagnostics in human geneti searches in ontologies Köhler et al., AJHG, October 2009. The Manual for the

mantic similarity omizer can be

Catalog – Features on keywords

Insert/select keywords based on features

Facture: Description Problem: Patient's Feature: Prod 01074 1 22 thes productivy Prod01075 1 22 thes productivy Prod01076 1 35 thes productivy Prod01076 1 45 thes productivy Prod01076 1 45 thes productivy Prod01076 1 56 thes productivy Prod01076 2 36 thes productivy Prod01076 2 36 thes productivy Prod01076 2 36 thes productivy Prod01076 2 46 thes productivy Prod01776 2 46 thes productivy Prod01776 2 46 thes productivy Prod01786 2 46 thes productivy Prod01786 2 46 thes productivy Prod01786 2 46 thes productivy	Menu. 🔹 Support t	the Phenomizer. Help.				
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'Syndromology' derived from syndrome: clinical approach to occurrence multiple congenital anomalies



Ped Cardiology clinic First description

9 patients with pulmonary valve stenosis, short stature
Hypertelorism, mild intellectual disability and ptosis cryptorchidism, 'skelettal malformations'



Noonan J and Ehmke D. J Pediatr 1963;63:468-470

Noonan JA. and Lexington K. Am J Dis Child 1968;116:373-380

Variability in phenotype

Large front, hypertelorism; down slanting palpebral fissures, low set posteriorly rotated ears

pectus carinatum

Remnant of prenatal nuchal translucency and hygroma

11 02 2022

Distinct patient...<u>similar</u> dysmorphic features Large front, Hypertelorism; down slanting palpebral fissures,low set posteriorly rotated ears

and variable degree of lymphedema (unconstant)

pectus carinatum

...similar phenotypes

Same syndrome...

8 months

4 years



Congenital heart defect « pulm atresia» IUGR Hypertelorism Macrocephaly (relative) Curly hair Failure to thrive **Evolving phenotype with age**

...Knowledge on natural history

...to adulthood

...to adult

- Sporadic
- Pulmonary valve stenosis
- Post natal growth retardation
- GH deficiency therapy





		Highest RASopathy	Specific Conditions				
	Gene	Association Achieved	Noonan	CFC	Costello	NSML	NS/LAH
	BRAF	Definitive	Moderate (10.5)	Definitive (13.5)	Disputed	Limited (5.5)	-
	HRAS	Definitive	-	-	Definitive (14.5)	-	-
	KRAS	Definitive	Definitive (14)	Strong (12.5)	Disputed	-	-
	MAP2K1	Definitive	Limited (3.5)	Definitive (12.5)	Disputed	Limited (2.5)	-
	MAP2K2	Definitive	Limited (1)	Definitive (12.5)	-	-	-
	NRAS	Definitive	Definitive (13.5)	Limited (0.5)	Limited (1)	Limited (3.5)	-
	PTPN11	Definitive	Definitive (13.5)	Disputed	Disputed	Definitive (15)	-
	RAF1	Definitive	Definitive (13.5)	Disputed	Disputed	Limited (4.5)	-
	RIT1	Definitive	Definitive (13.5)	-	-	-	-
	SHOC2	Definitive	Disputed	Disputed	Disputed	-	Definitive (12.75)
	SOS1	Definitive	Definitive (12.5)	Limited (1.5)	Disputed	-	-
	$LZTR1^{\dagger}$	Strong	Strong (12)	-	-	-	-
	PPP1CB	Strong	Strong (12.5)	-	-	-	Strong (12)
	SOS2 [†]	Moderate	Moderate (9.5)	-	-	-	-
	LZTR1 (AR)	Limited	Limited (8.75)	-	-	-	-
	MRAS	Limited	Limited (4.5)	-	-	-	-
	RRAS	Limited	Limited (3.25)	-	-	-	-
From	RASA2	Limited	Limited (1.5)	-	-	-	-
	A2ML1	Disputed	Disputed	-	-	-	-
IVI. Zenkel	RASA1	Disputed	Disputed	-	-	-	-

Management of Noonan Syndrome

A Clinical Guideline

Noonan Syndrome Guideline Development Group





Recommended baseline investigations in Noonan Syndrome



Remaining questions...

on bench side:

caveats: level of evidence from whole exome sequencing for variants in 'Rasopathy'

at bedside – genetic counselling Index patient and family How to explain penetrance ?

How to explain intra familial variable expressivity ?

c.922A>G (p.N308D) *PTPN11 Prevalent gene inside RASopathies- 40% Prevalent mutation – heart defect 60%* <u>Genetic counselling...</u> <u>in patients with very mild phenotype</u> <u>'de novo' autosomal dominant condition</u> <u>50% transmission</u> <u>Variable expressivity</u>

access to PGD

. . .



Birth defect to genetic counselling

Building evidence since 2020

Clinical Geneticists are far less pertinent then thought Human Hardware/brain memory on 4.000 mendelian disorders and phenotypes ?

Integrate targeted/non selective NGS gene panel approach / WES

Filtering on variants of unknown significance

When genetics approach become reverse ('unusual phenotype'), genotype identification ...« From Gene to Face » and vice-versa



Suggested Readings





Thank you for your attention

Open to your question



yves.sznajer@uclouvain.be