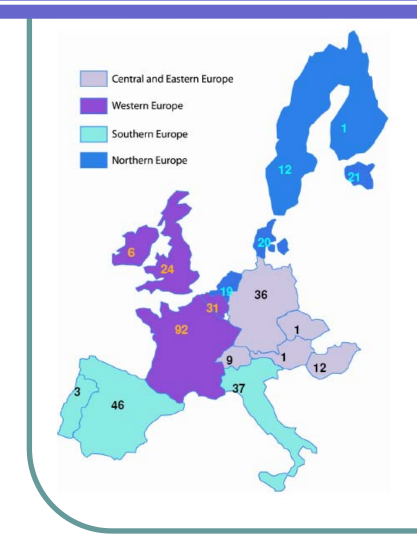
The Information Centre for Rare Diseases and Orphan Drugs as a Promoter of Research of Rare Diseases

Dr. Rumen Stefanov, MD, PhD Information Centre for Rare Diseases and Orphan Drugs (ICRDOD) Bulgarian Association for Promotion of Education and Science

Information Services: a European survey





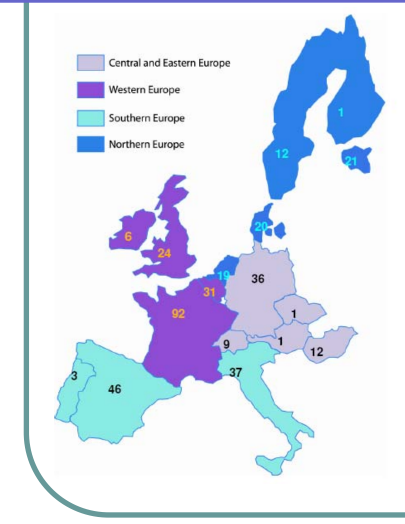
Analysis of a questionnaire to 372 information services on rare diseases

- From September to December 2003
- A questionnaire sent to 1402 information services in 20 EU countries
- A survey of organisations, their needs, sources, tools, services and expectations

EURORDIS PARD III: A PROJECT SUPPORTED BY THE RARE DISEASES PROGRAMME OF DIRECTORATE C OF THE EUROPEAN COMMISSION, AND ASSOCIATION FRANÇAISE CONTRE LES MYOPATHIES

Source: EURORDIS. Complete report available from www.eurordis.org

Information Services: a European survey

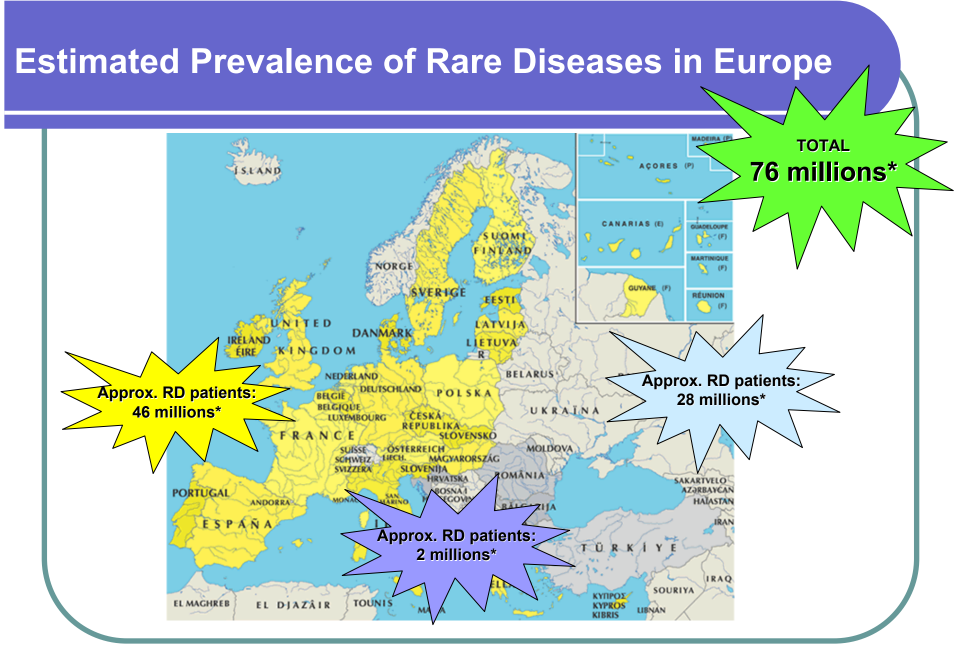




Key results:

- >372 respondents from 18 countries >62% dealing with a single rare disease 5% (~18) dealing with >50 diseases
- > 5% (~18) dealing with >50 diseases
- 26% with national funding
- \succ only 2% (~7) with EC funding

Source: EURORDIS. Complete report available from www.eurordis.org



Extrapolations based on the assumption of 9.19% prevalence rate of rare diseases. Source: http://www.wrongdiagnosis.com/r/rare_diseases/stats-country.htm

The Number of Specialized Centers for Rare Diseases in Europe is Extremely Limited

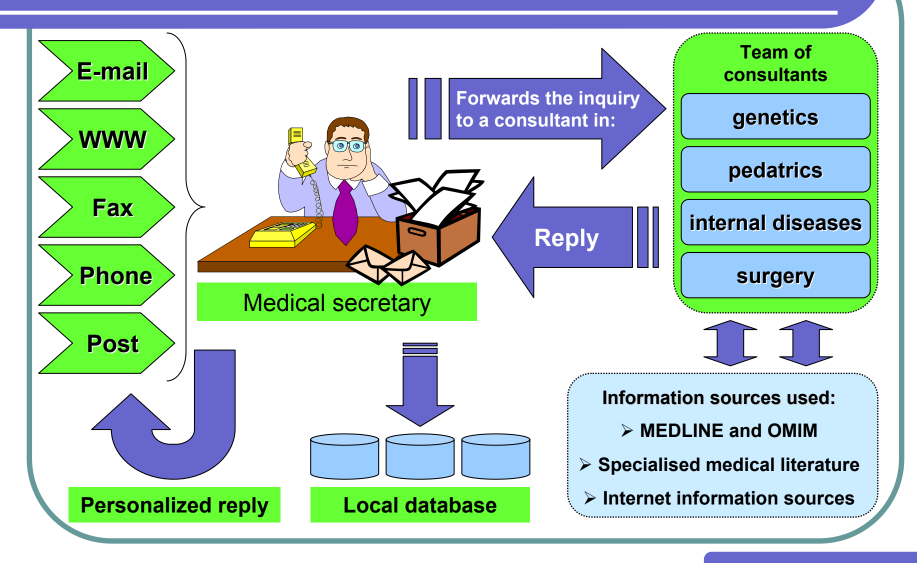
However, their impact on the public health is enormous:

- *First*, they provide patients, relatives and doctors with up-to-date high-quality information about the conditions of interest.
- Second, they systemize and create databases of hospital settings, genetic and clinical laboratories and prominent medical teams, occupied with a given rare disease research or study.
- *Third*, they provide evidences about the epidemiology of rare diseases.
- *Fourth*, these centres serve as a bridge between the patients with rare diseases and the public health care systems by advocating for the basic human right to receive contemporary and equal medical care.

Information Centre for Rare Diseases and Orphan Drugs (ICRDOD)

- Unique for Eastern Europe (40% of the European population)
- Operates in Bulgarian and English languages (projected to increase the number of languages with Russian and Greek)
- Provides free information to patients and medical professionals with educational purposes
- Builds databases of doctors, associations, clinical centers and clinical trials
- Provides data on the epidemiology of rare diseases
- Workshops, conferences and research on rare diseases
- Increases the public awareness on rare diseases
- Lobbying for adequate national health policy for rare diseases

Organization of Work



An example of reply to a medical professional - I

DISORDER: Parry–Romberg syndrome (PRS)

SYNONIMS: progressive hemifacial atrophy

DEFINITION: This syndrome is characterized with slowly progressive atrophy of the soft tissues of essentially half the face, accompanied usually by contralateral Jacksonian epilepsy, trigeminal neuralgia, and changes in the eyes and hair. It was first described by Parry (1825) and Romberg (1846) and it has been known for at least 2000 years. Because of its rarity, the literature on Parry–Romberg syndrome (PRS) largely consists of case reports. There is a prevalence of affected women towards men (M/F = 2/3).

CAUSES: They are not known yet. Autoimmune factors have been suggested but their role is not proved as yet. There is no data to support a direct heritability. A study which set up an internet interview among 205 patients with Parry–Romberg syndrome found that none of them has affected relatives, 3% report one relative with a pronounced face asymmetry but without a hemifacial atrophy (Stone, 2003). There was suggested a primary involvement of connective tissue in this disease (Bandello F at al., 2002). Some authors hypothesize that it could be related to a neural crest migration disorder, from which both fronto-nasal mass and cranial vessels take origin (Pichiecchio A at al., 2002).

GENETIC COUNSELING: Recomended.

An example of reply to a medical professional - II

AVAILABLE THERAPIES: At the moment there is no reliable therapy which would influence the progress of the disease. A reconstructive surgery is suggested for the advanced atrophy cases. Surgical intervention has to be planed in advance and performed at least two years after the disease becomes stable. There is a disease acceleration during or immediately after pregnancy.

EXPERIMENTAL THERAPIES: N/A

OMIM code: 141300

URL OMIM: http://www.ncbi.nlm.nih.gov/entrez/dispomim.cgi?cmd=entry&id=141300

ICD 10: L94.8

PATIENT ASSOCIATIONS:

http://www.geocities.com/rombergs/ http://www.geocities.com/HotSprings/1018/index.html

SPECIALIZED CLINICS: The surgical treatment is done in specialized clinics for plastic surgery. Accompanying symptoms are treated in the neurology, endocrinology or rheumatology departments as needed.

An example of reply to a medical professional - III

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AUTHOR OF THE REPORT: Assoc. Prof. Dr. Margarita Stefanova, PhD

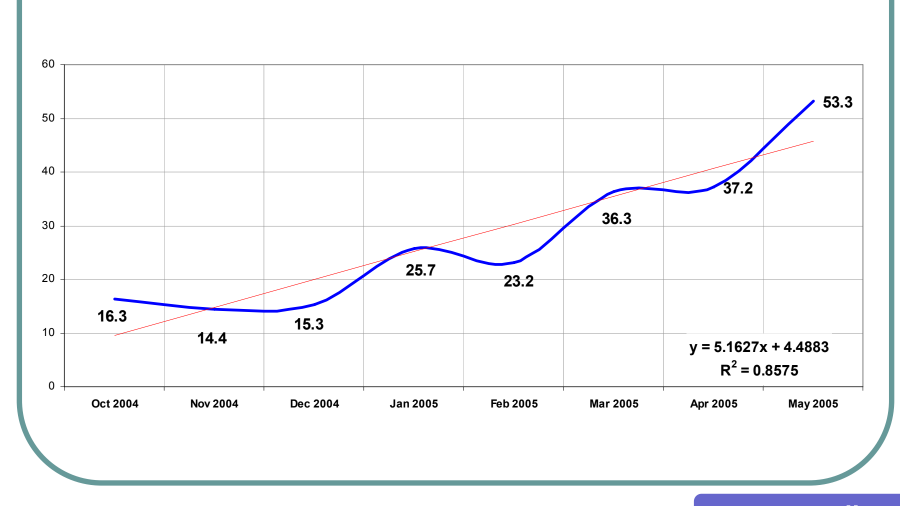
DATE OF LAST UPDATE: 31.10.2004

DISCLAIMER: The information in this reply is provided completely free of charge for educational purposes and should not be used for self diagnosis and treatment. In the case of a health problem, please contact your doctor!

The Internet Portal: www.raredis.org

Select your language Image: The 1-st Eastern European Continue on Rare Diseases and Orphan Drugs Search Home About Us Image: The 1-st Eastern European Countries Register Nov! Forums Image: The 2-st Eastern European Countries Register Nov! Advanced Search Rare Diseases Orphan Drugs Refister Nov! Image: The 2-st Eastern European Countries Register Nov! Corban Drugs Refister St PULMONARY ALVEOLAR MICROLLINITABING Register Nov! Image: The 2-st Eastern European Countries Refister Nov! Links Contact Us Image: Anarcterized by informer on 2005/4/19 22-47:18 (17 cadd) Refister Nov! Username: Dested by informer on 2005/4/19 22-47:18 (17 cadd) Refister Nov! Image: The 2-st Eastern European Countries Username: Dested by informer on 2005/4/19 22-47:18 (17 cadd) An ex disease profile is added to our database - Pulmonary alveolar reactioned the severe in a cation degree of dyspnees with a productive cough may occur together with a sporadic Image: The 2-st Eastern European Countries Username: Dested by informer on 2005/4/19 22-47:18 (17 cadd) An ex disease profile is added to our database - Pulmonary alveolar reactioned the severe in cadding of dyspnees with a sporadic Image: The 2-st Eastern European Countries Password? A new disease profile is added to our database -		Associate member of European Organisation for Rare Diseases - EURORDIS www.erordis.org	
Foruma Rare Diseases Orphan Drugs Ragistration Forms Links Contact Us Login Jsername: -acriatiodsreach characterized by infra-alveolar acloum deposits. The aetiology of the disease is still unknown. In most cases patients have mild clinical symptoms, contrasting with the severe ratiographic appearance: this is a typical feature that should raise the suspicion of PAM. However, a criatin degree of dyspneae with a productive cough may occur together with a sporadic haemoptysis and thoracic pains. To receive an e-mail with more information about the disease completely free of charge, please click here to send us a request. The profile was written by our consultant in internal disease) Dr. Nikolay Botushanov, MD. VertLogin Lost Password? Register now! Who's Online Q user(s) are online (2) the senders: profile is added to our database - the familel form of Cushing's syndrome. It is characterized histologically by small to normal-iszed glands with cortical micronodules (average 2) to a mm). About half of the cases are sporadic. The reminformation about the disease completely free of args, please click here to send us a request. The profile was written by our consultant in internal disease): o mm. About half of the cases are sporadic. The reminform of Cushing's syndrome. It is characterized histologically by small to normal-iszed glands with cortical micronodules (average 2) to a mm). About half of the cases are sporadic. The reminformation about the disease completely free of charge, please click here to send us a request. The profile was written by our consultant in internal diseases - Dr. Nikolay Botushanov, MD. Who's Online 0 mm. About half of the cases are sporadic. The reminformation about the disease completely fr	Menu Home	The 1-st Eastern European Conference on Rare Diseases and Orphan Drugs Promotion of Healthcare and Research on Rare Diseases and Orphan Drugs in Eastern European Countries 27 May 2005, Plovdiv, Bulgaria Register Now! Acc	Search Ivanced Search
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Links A new disease profile is added to our database - Pulmonary alveolar microlithiasis (PAP). If Rare Diseases Contact Us is a rare diseases, characterized by intra-alveolar calcium deposits. The aetiology of the disease is still unknown. In most cases patients have mild clinical symptoms, contrasting with the severe radiographic appearance: this is a typical feature that should raise the suspicion of PAM. However, a certain degree of dyspnoea with a productive cough may occur together with a sporadic haemoptysis and thoracic pains. To receive an e-mail with more information about the disease completely free of charge, please click here to send us a request. The profile was written by our consultant in internal diseases Emarphic appearance in the second is a disease is support to send us a request. The profile was written by our consultant in internal diseases User Login Lost Password? Rare Diseases : FAMILIAL FORM OF CUSHING'S SYNDROME Who's Online 2 user(s) are conline (2 user(s) are browsing fibe) Sum about half of the cases are spradic. The reminders occur in association with a variety of other abnormalities, including myxomatous masses of the heart, skin or breast; blue nevi and other endocrine disorders. To receive an e-mail with more information about the disease completely free of charge, please click here to send us a request. The profile was written by our consultant in internal diseases CEKLING NO XMANAGA W TMANAGA W TMAN		Posted by informer on 2005/4/19 22:41:18 (1 reads)	HEALTH
User Login Lost Password? Register now! who's Online 2 user(s) are online (2 user(s) are online (2 user(s) are browsing fibé/e) Members: 0 Guests: 2 Members: 0 Guests: 2 Remeters: 0 Comments?	Contact Us Login Username:	A new disease profile is added to our database - Pulmonary alveolar microlithiasis (PAM). It is a rare diseases, characterized by intra-alveolar calcium deposits. The aetiology of the disease is still unknown. In most cases patients have mild clinical symptoms, contrasting with the severe radiographic appearance: this is a typical feature that should raise the suspicion of PAM. However, a certain degree of dyspnoea with a productive cough may occur together with a sporadic haemoptysis and thoracic pains. To receive an e-mail with more information about the disease completely free of charge, please click <u>here</u> to send us a request. The profile was written by our consultant in internal diseases	Rare Diseases "Aldo e Cele Dacco", Aario Negri Institute, Italy Българско дружество по неврология
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The Internet portal – Unique Visitors per Day



Activities for Nov 2004 – April 2005

- Building and running the Internet service www.raredis.org
- Building team of consultants
- Creating databases of medical professionals, associations and clinical settings
- Established contacts with major national and international scientific and patient associations
- Deposited an official proposal for establishment of National program for rare diseases and orphan drugs at the Bulgarian Ministry of Health
- Organized a workshop on Hereditary Angioedema (11 March 2005) together with the Foundation for the Prevention and Treatment of Fatal Angioedematous Disease, Hungary
- Established and equiped office of the Information Centre

Results from the Workshop

on Hereditary Angioedema - 11 March 2005

- Created initiative group for improvement of diagnosis, treatment and healthcare of HAE patients in Bulgaria
- Establishment of HAE center in Plovdiv, referring for all Bulgarian HAE patients.
- Specialized training for two clinicians in Hungary.
- Dr. Farkas and Varga will provide **immunological diagnostic tests** for HAE to the Department of immunology at the Medical University of Plovdiv.
- Increasing the awareness and knowledge about HAE among medical specialists and patients in Bulgaria – papers, conferences.
- Genetic counseling to HAE patients
- Initiatives for reimbursement of diagnosis and treatment of HAE patients.







Plan for 2005

- Including rare diseases as a priority area in the National Health Strategy of Bulgaria
- Lobbying for reimbursement of the prophylaxis, diagnosis and treatment of all rare diseases
- Working towards adequate legislation, regarding easier access and registration of orphan drugs
- Close collaboration and integration with similar European organizations and services
- The First Eastern European Conference on rare diseases and orphan drugs – 27 May 2005 г.



Established Contacts, Support and Collaboration

