EUROCAT - Epidemiological studies

Fabrizio Bianchi
National Research Council
Institute of Clinical Physiology, Dept of Epidemiology
Rare Diseases Registry of Tuscany region

(on behalf of Professor Helen Dolk, Eurocat Project Leader)
Eurocat

European Surveillance of Congenital Anomalies

Funded by the Public Health Programme of the European Commission

WHO Collaborating Centre for the Epidemiological Surveillance of Congenital Anomalies
What is EUROCAT?

• European network of population-based registries for the epidemiologic surveillance of congenital anomalies.
• Started in 1979
• More than 1 million births surveyed per year in Europe
• 39 registries in 19 countries
• 25% of European birth population covered
• High quality multiple source registries, ascertaining terminations of pregnancy as well as births.
Map of Registries

- Full Member
- Associate Member
- < 10,000 births per year
- 10,000 – 40,000 births per year
- > 40,000 births per year
EUROCAT Objectives

• Provide essential epidemiologic information on congenital anomalies in Europe

• Co-ordinate the establishment of new registries throughout Europe collecting comparable, standardised data

• Co-ordinate the detection and response to clusters and early warning of teratogenic exposures
EUROCAT Objectives cont.

- evaluate the effectiveness of primary prevention

- assess the impact of developments in prenatal screening

- provide an information and resource centre and ready collaborative research network to address the causes and prevention of congenital anomalies and the treatment, care and outcome of affected individuals
Welcome to EUROCAT - European Surveillance of Congenital Anomalies

Welcome to eurocat

What is EUROCAT?
Contact Us
Member Registries
Surveillance
Publications and Data
Announcements
Membership Only

Useful Links
Selection Criteria

- A1: Total number of cases, number of cases by type of birth (liveborn, fetal death, induced abortion), and total prevalence rate per 10,000 births for congenital anomaly subgroups in selected registries (registries combined), selected time period (available for 28 full member registries).
- A5: Selected congenital anomaly: Total number of cases, number of cases by type of birth (liveborn, fetal death, induced abortion), population and total prevalence per 10,000 births per year and per registry in selected registries, selected time period (available for 32 registries).
- B3: Selected congenital anomaly: Total number of cases, number of cases by type of birth (liveborn, fetal death, induced abortion); total, birth, and livebirth prevalence rates per 10,000 births, per registry, in selected registries; selected time period (available for 32 registries).
- FI: Selected congenital anomaly: Line graph of total, birth and livebirth prevalence rate per 10,000 births per year, in selected registries, selected period.

(Note: Italicised type indicates that choice is available from an option menu.)

Date From: [ ] (eg. 1997)
Date To: [ ]

Continue

Click Here to view table and definitions

Programming by BioMedical Computing
### Eurocat

**European Surveillance of Congenital Anomalies**

#### Registries:
- Styria (Austria)
- Antwerp (Belgium)
- Hainaut (Belgium)
- Bulgaria
- Croatia
- Odense (Denmark)
- Strasbourg (France)
- Paris (France)

- Select All Full Members

#### Anomaly:
- Cleft lip with or without palate
- Cleft lip with or without palate
- Cleft palate
- Coarctation of aorta
- Common arterial truncus
- Congenital absence, atresia and/or stenosis of other specified parts of small intestine
- Congenital absence, atresia and/or stenosis of the duodenal
- Congenital absence, atresia and/or stenosis of the small intestine
- Cystic kidney disease
- Diaphragmatic hernia
- Down Syndrome (trisomy 21)
- Edward syndrome (trisomy 18)

- Select All Associate Members
(E3) - Cleft lip with or without palate (prevalence per 10,000 births) for the following registries: Galway, Dublin, Mersey, North Thames (West), Glasgow, From 1980 - 1999

<table>
<thead>
<tr>
<th>Registry</th>
<th>LB N</th>
<th>FD N</th>
<th>IA N</th>
<th>LD+FD+IA N</th>
<th>LB Rate</th>
<th>LD+FD Rate</th>
<th>LB+FD+IA Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dublin (Ireland)</td>
<td>347</td>
<td>20</td>
<td>0</td>
<td>367</td>
<td>8.31</td>
<td>8.73</td>
<td>8.73</td>
</tr>
<tr>
<td>Galway (Ireland)</td>
<td>31</td>
<td>5</td>
<td>0</td>
<td>36</td>
<td>5.44</td>
<td>6.28</td>
<td>6.28</td>
</tr>
<tr>
<td>Mersey (UK: England)</td>
<td>110</td>
<td>2</td>
<td>5</td>
<td>117</td>
<td>6.56</td>
<td>6.84</td>
<td>6.94</td>
</tr>
<tr>
<td>North Thames (West) (UK: England)</td>
<td>338</td>
<td>19</td>
<td>74</td>
<td>431</td>
<td>7.14</td>
<td>7.55</td>
<td>9.11</td>
</tr>
<tr>
<td>Glasgow (UK: Scotland)</td>
<td>191</td>
<td>13</td>
<td>22</td>
<td>226</td>
<td>7.49</td>
<td>7.96</td>
<td>8.81</td>
</tr>
<tr>
<td>Wales (UK: Wales)</td>
<td>61</td>
<td>1</td>
<td>10</td>
<td>72</td>
<td>5.25</td>
<td>6.32</td>
<td>7.34</td>
</tr>
<tr>
<td>Total (full member registries):</td>
<td>1078</td>
<td>60</td>
<td>111</td>
<td>1249</td>
<td>7.34</td>
<td>7.72</td>
<td>8.47</td>
</tr>
</tbody>
</table>

LB = Live Births  
FD = Fetal Deaths / Still Births from 20 weeks gestation  
IA = Induced Abortions following prenatal diagnosis

For total live birth rates where the live birth denominator is not available, the denominator has been substituted by the total number of births (live births + still births). This applies to the following registries: North Thames (West)

<table>
<thead>
<tr>
<th>Description Of Anomaly</th>
<th>ICD9</th>
<th>ICD10</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clefting of the upper lip with or without clefting of the maxillary alveolar process and hard and soft palate</td>
<td>7491, 7492</td>
<td>Q36-Q37</td>
<td></td>
</tr>
</tbody>
</table>
Why European collaboration?

• Pooling of data
• Comparison of data
• Common response to public health questions
• Sharing of expertise and resources
EUROCAT Special Reports and Coding Guides 2005-2006


A full list of EUROCAT publications and links to Reports can be found on: www.eurocat.ulster.ac.uk/pubdata/publist.html
8. ..............................
15. ............
EUROCAT News

• Further funding has been obtained from the European Commission for the period September 2007 to August 2010

• Prevalence data updated to 2006 on website.
(F1) - Neural Tube Defects (total, birth and livebirth prevalence per 10,000 births) for the following registries: Barcelona, France.

The graphs on this page require the Java2 runtime environment.
Click here if you cannot see the graph below.

Legend:
- Total Prevalence
- Birth Prevalence
- Live Birth Prevalence

(F1) - Neural Tube Defects (total, birth and livebirth prevalence per 10,000 births) for the following registries: Basque Country, Asturias, Barcelona, From 1990 - 2002

The graphs on this page require the Java2 runtime environment. Click here if you cannot see the graph below.

LEGEND
Total Prevalence
Birth Prevalence
Live Birth Prevalence

Asturias (Spain)  Barcelona (Spain)  Basque Country (Spain)
Surveillance
This web site makes use of Adobe's Portable Document Format (PDF). With the free Adobe Acrobat Reader software, you can view and print Adobe PDF files across a broad range of hardware and operating systems.

For further information or to download the free reader click here
This web site makes use of Adobe's Portable Document Format (PDF). With the free Adobe Acrobat Reader software, you can view and print Adobe PDF files across a broad range of hardware and operating systems.

For further information or to download the free reader click here
Guide 1.3 - Instructions for the Registration and Surveillance of Congenital Anomalies

Annex Documents

- EUROCAT Guide 6: Definition and Coding of Syndromes
- ICD10-BPA Extension Codes
EUROCAT activity on rare congenital anomalies

- routine collection, registration, analysis

- ad-hoc studies on rare
ADOPTED AND FINALISED: Commission Proposal for a New Health and Consumer Strategy and Programme

Useful Links

Rare Diseases Task Force

Rare Diseases www.rdtf.org

DG Sanco Rare Diseases http://www.europa.eu.int/comm/health/ph_threats/non_com/rare_diseases_en.htm

Orphanews www.orpha.net

Rare Diseases Spain http://repier.relics.net/repier/home.aspx

Orphanet Journal of Rare Diseases www.ojrd.com
ACTIVITIES ON RARE CONGENITAL ANOMALIES: FIVE EXAMPLES

1. Epidemiology of rare syndromes in Europe
2. Registration and surveillance of Sentinel Phenotypes
3. Study of Gastrochisis
4. Survey of confidentiality and consent issues in EUROCAT registries
5. Folic acid and congenital anomalies
Main objectives are to:

- produce prevalence rates and descriptive epidemiology data on 9 rare malformation syndromes recognisable at birth
- assess the possible impact of prenatal diagnosis on the prevalence rates
- look for possible regional differences regarding the time of diagnosis of selected syndromes
EPIDEMIOLOGY OF RARE SYNDROMES IN EUROPE

cont..

• study type of birth (TOP, SB, LB) and survival

• describe clinical presentation and associated malformations in relation to “pure” or “minimal diagnostic criteria” definition of selected syndromes in non-Mendelian conditions

• look into geographic and time distribution for possible trends and clusters

• investigate the possible teratogenic exposures
EPIDEMIOLOGY OF RARE SYNDROMES IN EUROPE

cont …

• assess the accuracy of the diagnosis in registries of congenital malformations

• determine the role of medical geneticists in local registries in the diagnosing of syndrome cases

• evaluate the process of coding to improve the coding system
## Epidemiology of Rare Syndromes in Europe

<table>
<thead>
<tr>
<th>Syndrome</th>
<th>Reference</th>
<th>Prevalence (*)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fraser</td>
<td><em>Martinez-Frias et al 1998</em></td>
<td>4.3</td>
</tr>
<tr>
<td>Cornelia de Lange</td>
<td><em>Beck et al 1976</em></td>
<td>6.0</td>
</tr>
<tr>
<td>Holt-Oram</td>
<td><em>Elek et al 1991</em></td>
<td>9.5</td>
</tr>
<tr>
<td>Fryns</td>
<td><em>Ayme et al 1989</em></td>
<td>7.0</td>
</tr>
<tr>
<td>Goldenhar</td>
<td><em>Gorlin et al 1990</em></td>
<td>18.0</td>
</tr>
<tr>
<td></td>
<td><em>Morrison et al 1992</em></td>
<td>22.0</td>
</tr>
<tr>
<td>Meckel-Gruber</td>
<td><em>Salonen and Norio 1984</em></td>
<td>11.1</td>
</tr>
<tr>
<td></td>
<td><em>Martinez-Frias et al 1994</em></td>
<td>8.0</td>
</tr>
<tr>
<td>Ellis van Creveld</td>
<td><em>Mahony 1994</em></td>
<td>7.0</td>
</tr>
<tr>
<td>Beckwith-Wiedemann</td>
<td><em>Hamel et al 1981</em></td>
<td>66.7</td>
</tr>
<tr>
<td>Treacher-Collins</td>
<td><em>Jahrsdoerfer et al 1995</em></td>
<td>20.0</td>
</tr>
</tbody>
</table>

(*) per 10,000 births
Based on 23 years of epidemiologic monitoring (8,558,346 births in the 1980-2002 period), we found the prevalence of the classical form of CdLS to be 12.4/10,000 and estimated the overall CdLS prevalence at 16-22/10,000.

- live born 91.5%
- fetal deaths 2.8%
- termination of pregnancy 5.7%
The most frequent associated congenital anomalies were:

- limb defects (73.1%)
- congenital heart defects (45.6%)
- central nervous system malformations (40.2%)
- cleft palate (21.7%)
Descriptive epidemiology of Cornelia de Lange syndrome in Europe

• In the last 11 years, as much as 68% of cases with major malformations were not detected by routine prenatal US
• Live born infants have a high first week survival (91.4%)
• All patients were sporadic
• Maternal and paternal age did not seem to be risk factors
• Almost 70% born after the 37th week of gestation weighed ≤ 2,500 g.
• Low birth weight correlated with a more severe phenotype.
• Severe limb defects were significantly more often present in males.
THE PREVALENCE AND SURVEILLANCE OF “SENTINEL PHENOTYPES”

• Surveillance of “sentinel phenotypes” has been proposed as a way of monitoring the rate of new mutations in a population


• For a successful monitoring, SP should be well defined and diagnosed prenatally or soon after birth. Since information is not available to verify familial inheritance for each case, syndromes can be chosen which are always or usually due to new mutations, including those lethal in early life where familial inheritance is not possible.
The EUROCAT subcommittee on Multiple Malformations recommends five syndromes, caused by new mutations (rather than familial inheritance) in more than 90% of cases.

- Achondroplasia 1.3 - 15.0 per 100,000 births
- Thanatophoric dysplasia 0.7 - 3.8 “
- Osteogenesis imperfecta 3.6 - 4.0 “
- Campomelic dysplasia 0.5 “
- Apert Syndrome 0.7 “
Gastroschisis is a rare abdominal wall anomaly which has been increasing in prevalence in Europe and worldwide.

It is well established that young maternal age is a strong risk factor for this anomaly. Smoking, recreational drug use and some therapeutic drugs are also risk factors.

A working group for investigating trends in prevalence by maternal age has been activated and a paper is in press.
Study of Gastroschisis

We analysed 936 cases of gastroschisis from 25 population-based registries in 15 European countries, 1980-2002.

The maternal age standardised prevalence rose from 0.54 per 10,000 in 1980-84 to 2.12 per 10,000 in 2000-2002.
Study of Gastroschisis

The relative risk for mothers <20 years of age 1995-2002 was 7.0 [95% C I 5.6, 8.7]

There were geographical differences within Europe, with higher rates in the UK and lower rates in Italy, after adjusting for maternal age
After standardising for regional variation, our results showed that the increase in risk over time was the same for mothers of all ages.

These findings indicate that the phenomenon of increasing gastroschisis prevalence is not restricted to younger mothers only.
Survey of confidentiality and consent issues in EUROCAT registries

The EUROCAT survey was published in BMJ, 2005. It was also presented to the Network of Competent Authorities, 5-6 July 2005 in Luxembourg and the Rare Diseases Task Force.

Registries collecting personal medical data must, under EC Directive 95/46/EC, obtain consent for the processing of such data, unless national law or a national supervisory body allows for an exemption. Member states have not always taken advantage of the ability to exempt health care or disease registries.
EUROCAT experience indicates that opt-in consent poses a serious threat to the operation of congenital anomaly registries.

The debate about informed consent has eclipsed more relevant consideration of procedures to maintain confidentiality of data or ensure the ethical operation of registries.
EUROCAT activities on FOLIC ACID

Annual survey in member countries of policy changes regarding:

- periconceptional FA supplementation
- health education initiatives undertaken
- available data on FA uptake
- analysis of available data on protective effect for other congenital anomalies
EUROCAT activities on FOLIC ACID

The 2007 update documents policy changes in a few countries, especially Italy, and the extremely slow progress in fortification of food with FA.

Although fortification has been advised in a few countries, it has not yet been implemented.

This report is on the EUROCAT website.

EUROCAT continues to advocate the recommendations from the earlier report.

Country representatives supply information on policy and practice changes periodically.
Welcome to EUROCAT - Microsoft Internet Explorer

Address: http://www.eurocat.uls.ac.uk/

Eurocat
European Surveillance of Congenital Anomalies

Welcome to eurocat

What is EUROCAT?
Contact Us
Member Registries
Surveillance
Publications and Data
Antenatalcare
Membership Only

Useful Links