Limb reduction defects in Emilia Romagna, Italy: epidemiological and genetic study in 173 109 consecutive births

Elisa Calzolari, Daniela Manservigi, Gian Paolo Garani, Guido Cocchi, Cinzia Magnani, Mario Milan

Abstract
Epidemiological and genetic variables in limb reduction defects (LRD) were analysed during the years 1978 to 1987 in a case control study in Emilia Romagna, northern Italy. During the observation period, 83 neonates out of 173 109 consecutive births had LRD (4.8 per 10 000). Cases were divided into five subgroups: transverse, intercalary, longitudinal, split, and multiple types of LRD. Of all cases, 64% were upper limb, 21% lower limb, and 15% both. Coexisting non-limb malformations were found in 10 cases (12%), five with recognised syndromes and five with other associated defects. About 7.2% of first degree relatives had defects involving the skeletal system. In two cases the mother had the same type of LRD (a split). No recurrence among sibs was observed. Risk factors correlated with LRD were found to be low birth weight (2500 g or less), vaginal bleeding, and threatened abortion.

Limb reduction defects (LRD) are severe anomalies occurring in about one in every 2000 (five per 10 000) newborn infants. Some of these anomalies are genetic in origin, resulting from dominant and recessive genes and chromosomal abnormalities.

In addition, uterine factors (amniotic bands and mechanical and constraint problems), vascular compression, thrombosis, embolisation, and maternal diseases and infections have been shown to cause limb malformations. At present thalidomide is the only known drug that can produce severe LRD in humans and only during a very brief period of early organogenesis (22 to 36 days postconception). Warfarin and anticonvulsants may produce digital hypoplasia by causing prolonged vascular or nutritional deprivation during pregnancy. However, the aetiology of many LRD is still unknown, and it is possible that a certain percentage can be attributed to the statistical probability that problems will arise during the complicated embryonic processes. In other words, to reduce the frequency of malformations beyond a certain limit may prove impossible despite continuing studies of the aetiology of malformations and preventive measures.

From a clinical point of view, LRD can either be isolated or part of syndromes or multiple non-limb malformations.

The present study reviews infants reported to have LRD in the Emilia Romagna register (1978 to 1987). Assessments were made in terms of frequency, referral patterns, co-occurring malformations, and aetiological factors.

Materials and methods
THE REGISTER
Details of the Emilia Romagna register for congenital malformations have been published previously. In this study, data on infants with congenital anomalies (whether born alive or stillborn after the 28th week of gestation) were reported using a specially designed questionnaire filled out by the medical staff during a personal interview with the mother. Each of these registered mothers was paired with a control mother whose delivery immediately preceded hers in the maternity unit. Diagnoses were written out rather than being given code numbers.

Descriptions of the malformations were supplemented with drawings, photographs, x rays, and...
necropsy reports. All information was transmitted to the coordination centre, where it was reviewed by a
trained clinician. If the information was incomplete, the
missing data were requested from the reporting
hospital.
EUROCAT guidelines\(^{17}\) were used to describe
LRD.

CLASSIFICATION OF LRD

Limb reduction defects were defined as the absence or
severe hypoplasia of skeletal structures of the limb. Excluded
from the analysis were cases of brachydactyly and sirenomelia. The recorded cases were
classified according to Temtamy and McKusick\(^{18}\) and
EUROCAT guidelines.\(^{17}\)

The following morphological types were differenti-
ated (table 1).

1. Terminal transverse: absence of the distal
structures of the limb with the proximal structures
more or less normal. Ectrodactyly was defined as
partial or total absence of distal segments of the hand.
The malformation can involve phalanges only
(aphalangia), digits (adactylia), or the full hand
(acheiria). All are considered to be various degrees of
the same malformation.

2. Intercalary: absence or severe hypoplasia of
proximal (humerus and femur) and distal (radius and
ulna or tibia and fibula) parts of the limb with normal
or malformed hands and feet. The presence of distal
skeletal structures differentiates intercalary defects
from terminal transverse defects. The term phocomelia
was avoided in this study because it has various
meanings in published reports.

3. Longitudinal: absence or severe hypoplasia of
the lateral part of the limb. Two main subtypes were
considered: preaxial (thumb, first metacarpal, radius;
or hallux, first metatarsal, tibia) and postaxial (fifth
finger, fifth metacarpal, ulna; or fifth toe, fifth
metatarsal, fibula).

4. Split hand/foot: paraxial or axial longitudinal
terminal deficiency of rays, usually associated with
syndactyly. Typical and atypical categories are
recognised. The typical split category has two
anatomical variants: lobster claw (absence of the
central ray) and monodactyly (deficiency of radial rays
with no clefts). The anatomical classification has no
genetic significance because either type may occur in
the same family or in different limbs of the same
person. Generally, the atypical category is sporadic,
affects only one hand (central rays are absent), does
not involve the feet, and has hypoplastic digits on the
axial portion of the hand.

5. Multiple types of limb reduction: more than one
type of LRD according to the definitions given above.
The femur-fibula-ulna (FFU) complex was included
in this category.

The cases in the study were divided into two major
groups: (1) infants with only limbs affected (one or
more); and (2) infants with coexisting non-limb
malformations. In this group 'syndromes' and 'as-
sociations' were taken into consideration.

CODES OF LRD

The standard International Classification of Diseases
(ICD)\(^{19}\) code was not detailed enough to allow
classification of specific malformations and thus was
inadequate for surveillance purposes. Therefore, we
formulated a special numerical code, based on the
morphological types listed above and in table 1, to
classify each LRD more specifically.

<table>
<thead>
<tr>
<th>Defect</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Terminal transverse</td>
<td>Absence of the distal structures of the limb. The following types are considered: Ectrodactyly: total or partial absence of phalanges (aphalangia), digits (adactylia), or full hand (acheiria). Amelia: total absence of extremities. Hemimelia: total absence of forearm and hand or of foreleg and foot. Apodia: absence of foot.</td>
</tr>
<tr>
<td>Intercalary</td>
<td>Absence or severe hypoplasia of proximal (radius and ulna or tibia and fibula) part of the limb with hands and feet normal or malformed. The presence of distal skeletal structures differentiates intercalary defects from terminal transverse.</td>
</tr>
<tr>
<td>Longitudinal</td>
<td>Absence or severe hypoplasia of lateral part of the limb: Preaxial (thumb, first metacarpal, radius; or hallux, first metatarsal, tibia). Postaxial (fifth finger, fifth metacarpal, ulna; or fifth toe, fifth metatarsal, fibula).</td>
</tr>
<tr>
<td>Split hand/foot</td>
<td>Paraxial or axial longitudinal terminal deficiency of rays, usually associated with syndactyly. Two main anatomical varieties of typical split are: Lobster claw (absence of the central ray). Monodactyly (deficiency of radial rays with no clefts). Atypical split resembles typical split type 1.</td>
</tr>
<tr>
<td>Multiple</td>
<td>More than one type of limb reduction (eg FFU complex).</td>
</tr>
</tbody>
</table>
FACTORS ASSOCIATED WITH LRD
To test for an association between LRD and various epidemiological variables, syndromes with recognised Mendelian inheritance were excluded from the study.

Of the variables included on the registration form, the following were studied: infant's sex, parental age at delivery, history of selected maternal diseases, threatened abortion, maternal exposure to drugs and smoking during pregnancy, and other maternal factors including parity, previous abortions, stillbirths, and malformed infants.

The presence of LRD or other malformations in parents or sibs was recorded during interviews with the mothers.

STATISTICAL ANALYSIS
Statistical tests included the $\chi^2$ test for the presence or absence of the characteristics under study in index patients and controls and the $t$ test for comparison of means. For detection of time clusters, we used the observed/expected ratio$^{20}$ and the cumulative sum techniques.$^{21}$

Results
CHARACTERISTICS OF THE POPULATION UNDER STUDY
During the study period (1978 to 1987), 83 cases of LRD were detected among 173 109 births in the Emilia Romagna region, giving a prevalence of 4.8 per 10 000.

There were 73 (88%) isolated cases (one or more limbs affected) and 10 (12%) with coexisting non-limb malformations, five (6%) with recognised syndromes or associations (table 2) and five (6%) with other associated defects (table 3).

A case of sirenomelia was excluded, and the remaining 82 cases were divided into six groups, according to the definitions described above (table 4).

Terminal transverse defects were the most common, occurring in 2.6 cases per 10 000. Other defects and their prevalences per 10 000 included: split hand/foot 0.58, longitudinal preaxial 0.52, intercalary 0.46, longitudinal postaxial 0.35, and multiple LRD 0.23 (fig 1).

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Table 2. Coexisting non-limb malformations: recognised syndromes or associations with LRD.

<table>
<thead>
<tr>
<th>Syndrome</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fanconi</td>
<td>Bilateral radial aplasia + bilateral hypoplasia of first metacarpal and first finger.</td>
</tr>
<tr>
<td>TAR</td>
<td>Bilateral radial aplasia + thrombocytopenia + unspecified cardiac defect.</td>
</tr>
<tr>
<td>ADAM</td>
<td>Ring encircling two fingers of the right hand and two toes of the left foot + limb constriction of the left leg with partial amputation of the foot.</td>
</tr>
<tr>
<td>Moebius</td>
<td>Terminal transverse bilateral arms (acheiria bilateral) + terminal transverse unilateral leg + facial paralysis + micrognathia.</td>
</tr>
<tr>
<td>VATER</td>
<td>Intercalary unilateral limb defect + atresia of oesophagus + ventricular septal defect + urinary tract anomalies.</td>
</tr>
</tbody>
</table>

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Table 3. Coexisting non-limb malformations: other associated defects (each association occurred once).

<table>
<thead>
<tr>
<th>Defect</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Proximal intercalary defect unilateral leg + transposition of great vessels.</td>
<td></td>
</tr>
<tr>
<td>Preaxial longitudinal defect (absence of first metacarpal) + atresia of auricle + dextroversion.</td>
<td></td>
</tr>
<tr>
<td>Terminal transverse unilateral leg + spina bifida + gastroschisis + ambiguous genitalia.</td>
<td></td>
</tr>
<tr>
<td>Terminal transverse unilateral (amelia) + encephalocele + gastroschisis.</td>
<td></td>
</tr>
<tr>
<td>Terminal transverse unilateral (apodia) + atresia of rectum.</td>
<td></td>
</tr>
</tbody>
</table>

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Table 4. Distribution of types of LRD in 82 reported cases.

<table>
<thead>
<tr>
<th></th>
<th>Upper limb</th>
<th>Lower limb</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Unilateral</td>
<td>Bilateral</td>
</tr>
<tr>
<td>Terminal transverse</td>
<td>29</td>
<td>2</td>
</tr>
<tr>
<td>Intercalary</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Longitudinal preaxial</td>
<td>6</td>
<td>3</td>
</tr>
<tr>
<td>Postaxial</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Split</td>
<td>7</td>
<td>2</td>
</tr>
<tr>
<td>Multiple</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Total</td>
<td>45</td>
<td>8</td>
</tr>
</tbody>
</table>
MALFORMATION FREQUENCY
To study trends over time, as well as space and time clusters, all cases of LRD, including those associated with other defects, were considered. No significant trend was detected (analysis computed for successive and constant sets of 10 000 births by De Wals and Lechat). No marked intraregional differences for prevalence of LRD were noted ($\chi^2=8.76$, df 7, NS).

No time clusters were found with the cumulative sum technique.

FACTORS ASSOCIATED WITH LRD
Cytogenetics
Karyotypes were obtained in 15 of 83 cases; all were normal.

Consanguinity and family data
There was no instance of parental consanguinity among the affected cases or controls. In the affected group, six first degree relatives (about 7%) had a musculoskeletal anomaly (two cases of congenital hip dislocation, two of polydactyly, two of split hand). In the control group, four first degree relatives had congenital hip dislocation.

Environmental factors
Of the several factors considered, only threatened abortion (bleeding or pain or both), treatment for abortion, and birth weight of 2500 g or less were related to LRD. Threatened abortion was diagnosed in 21·6% (1674) of affected pregnancies as compared to 6·0% (583) of controls ($\chi^2=6.42$, df 1, $p<0.05$).

Treatment for threatened abortion or history of previous abortion or both was given in 31·1% of index pregnancies (2374) and 9·6% of controls (883) ($\chi^2=10.27$, df 1, $p<0.001$).

The mean birth weight for the index cases was 2882 g as compared to 3363 g for the control group. The difference is significant ($t=5.09$, $p<0.001$). In particular, birth weights of 2500 g or less were found in 21·5% (1779) of the LRD cases and in 2·5% (280) of controls ($\chi^2=11.92$, df 1, $p<0.001$). These findings are in keeping with those reported by Aro et al., Polednak and Janerich, and Kallen.

Discussion
The purpose of this study was to analyse limb reduction defects as reported in the Emilia Romagna register (1978 to 1987). The prevalence of LRD in this region (4·8/10 000) was in keeping with that observed elsewhere in Italy and in other countries. The rates of types of LRD (fig 2) indicate that, in accordance with published reports, terminal transverse defects are the most common (2·6/10 000), and are followed in descending order by longitudinal preaxial (0·52), intercalary (0·46), and longitudinal postaxial (0·35).

The prevalence of split hand/foot (0·58/10 000) in our study did not differ greatly from that reported in Hungary (0·4).1

The frequency of LRD with coexisting non-limb malformations was 12% (five cases with recognised syndromes and five with other associated defects), while the percentage of associated anomalies was 30% in a Swedish study2 and 53% in a Canadian study.4 This difference is because in our study the cases of LRD coexisting with other malformations of the musculoskeletal system were not considered as associated defects (for example, polydactyly or triphalangeal thumb described in association with split malformation, or talipes with absence of fibula). In addition, the simultaneous occurrence of the same type of defect (for example, terminal transverse in upper and lower limb) was not considered to be an associated defect.

No intraregional variations in prevalence were observed, and no time clusters were identified with the observed/expected ratio or cumulative sum techniques. Special codes were used for surveillance purposes.

No recurrences among sibs were observed. In two of 10 cases of split, a mother and child were affected, which may be attributable to autosomal dominant inheritance.

Among the variables considered in the present study, threatened abortion, treatment for abortion (for example, hormones), and birth weight of 2500 g or less were significantly related to LRD. These are the only three variables that were found to be consistently correlated with LRD in previous reports.2 23- 25

Vaginal bleeding and threatened abortion can be considered indicators for impaired embryogenesis, but are not specifically associated with LRD.

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