Cerebral Palsy Among Kurdish Children in the City of Dohuk: A Case-series Study

Khalaf H. Hassan*1

Abstract

Background: Cerebral palsy (CP) is defined as non-progressive disorder of movement and posture due to brain insult or injury occurring in the period of early brain growth (generally under three years of age). The clinical characteristic of this common problem had not been studied before in any of the Kurdish cities of the north of Iraq.

Objectives: This is a descriptive study of the clinical characteristics of cerebral palsy in the city of Dohuk at the north of Iraq.

Patients and Methods: This is a case series descriptive study conducted in the children rehabilitation unit of Azadi general hospital and Dohuk children's rehabilitation center in the city Dohuk on children with Cerebral palsy. The following information were studied, the first presenting compliant, age distribution, the antenatal illnesses, antenatal drug use, mode of delivery, gestational age, the state at birth, types of CP, associated congenital abnormalities, any other associated problems and the treatment used.

Results: The total number of patients were 100, males were outnumbering females (1.3:1), spastic CP was the most common type (57%). Most of the patient presented between the age of 7-12 months (66%), the most common first presenting complaints were the delayed milestones (50%), most of the patients came from a vaginal delivery (68%), the associated problems were: swallowing problems (68%), walking problems (65%), epilepsy and seizure disorders (60%), speech problems (46%), neonatal jaundice which was developed in 95% of cases, and finally, most of the patient came to parents who got married as a result of consanguineous marriage(68%).

Conclusions: This is the first descriptive study among children with CP from the Kurdish population highlighting certain demographic and clinical characteristics of them.

Keywords: Cerebral palsy, Kurdish children, Dohuk

Introduction

Cerebral palsy (CP) is a diagnostic term used to describe a group of motor syndromes resulting from disorders of early brain growth (generally under 3 years of age). The manifestation of a given lesion may change as the nervous system

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mature, but the insult that caused the lesion is no longer present and there is no active disease at the time of diagnosis, although it has historically been considered as a static encephalopathy. This term is not entirely accurate because of the recognition that the neurological features of CP often change or progress over time.\textsuperscript{1,2,3,4} This disease was first described by Dr. Little in 1826.\textsuperscript{5}

CP is the most common cause of severe childhood disability affecting 1 in 400 children,\textsuperscript{6} the incidence varies from one country to another but it is around 1-2/1000 live birth, and; prevalence is about 500/100000 population.\textsuperscript{7,8,9,10}

Traditionally, spastic CP has been the most frequent type, accounting for approximately 50% of cases followed by Athetoid CP seen in approximately 20%.\textsuperscript{11,12}

The clinical findings in some affected children appear to evolve and affected children may exhibit a somewhat different distribution of involvement at different ages.\textsuperscript{13}

A team of physicians from various specialties, as well as occupational and physical therapists, speech pathologists, social workers, educators, and developmental psychologists provided important contribution to the treatment of these children.\textsuperscript{1}

This study was conducted to assess the clinical and demographic characteristics of CP children among the Kurdish population at the North of Iraq.

**Patients and Methods**

The period of the study extended for one year from May 2002 till April 2003 on patients who were visiting the Dohuk children's Rehabilitation Center (DCRC) and Children Rehabilitation Unit of Azadi General Hospital (CRU) for regular physiotherapy and for treatment of any other associated problem. A special data collection sheet was developed for the purpose of this study. One hundred children with a diagnosis of CP were retrospectively studied.

Information was obtained by the means of direct questionnaire of parents and/or from the registration record of DCRC and the birth records of Azadi general hospital. The author had examined all the patients, regarding the other problems associated with CP. The APGAR scoring system (14) was used to assess the newborn delivery score, being good if more than 8, average 3-7, and low if below 3. The ethical committee of azadi general hospital had approved conducting the study. Consent was taken from all children guardians. Data were analyzed using simple statistical analysis packages (Epi info 6).

**Results**

One hundred children with CP were studied, male accounting for 57% and females for 43%. The most common first presenting compliant was the delayed milestones (50%) as shown in table (1).

<table>
<thead>
<tr>
<th>Complaint</th>
<th>No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Delayed milestones</td>
<td>50</td>
</tr>
<tr>
<td>Convulsion</td>
<td>14</td>
</tr>
<tr>
<td>Difficulty in swallowing</td>
<td>14</td>
</tr>
<tr>
<td>Recurrent LRTI</td>
<td>9</td>
</tr>
<tr>
<td>Hand preference</td>
<td>8</td>
</tr>
<tr>
<td>FTT</td>
<td>5</td>
</tr>
<tr>
<td>Total</td>
<td>100</td>
</tr>
</tbody>
</table>

Majority of the patients presented between the age of 7-12 months (66%), other age distributions are shown in figure (1).

**Figure (1): Age of presentation.**
Seventy six percent of mothers were attending the antenatal clinics, about 21% of them had pre-eclamptic toxemia.

Sixty eight percent of the patients were delivered by normal vaginal delivery and the other modes of deliveries are shown in figure (2).

Figure (2): The mode of delivery.

The Apgar score ranged between: good in 48%, average in 39% and low in 13%. History of asphyxia was reported in 48 cases, 45 cases of them had been admitted previously to the NICU; while the remaining 3 cases were home delivery and they had no access to the medical care. Neonatal jaundice developed in 95% of cases, 60% received no treatment, 34% were treated with physiotherapy and 6% required exchanging transfusion.

Spastic CP was the most common type among the studied cases (57%), among the spastic CP; the spastic quadriplegia was the most common (72%); the other types of spastic CP are shown in table (2).

Table (2): Types of CP.

<table>
<thead>
<tr>
<th>Type</th>
<th>No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spastic</td>
<td>57</td>
</tr>
<tr>
<td>Quadriplegia</td>
<td>41</td>
</tr>
<tr>
<td>Hemiplegia</td>
<td>8</td>
</tr>
<tr>
<td>Diplegia</td>
<td>7</td>
</tr>
<tr>
<td>Monoplegia</td>
<td>1</td>
</tr>
<tr>
<td>Hypotonic</td>
<td>19</td>
</tr>
<tr>
<td>Chorio-arthetoid</td>
<td>6</td>
</tr>
<tr>
<td>Ataxic</td>
<td>5</td>
</tr>
<tr>
<td>Mixed</td>
<td>13</td>
</tr>
<tr>
<td>Total</td>
<td>100</td>
</tr>
</tbody>
</table>

Table (3) details the associated congenital abnormalities that were found in 16% of cases and table (4) lists the associated problems.

Table (3): Associated congenital anomalies.

<table>
<thead>
<tr>
<th>Anomaly</th>
<th>No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Musculoskeletal</td>
<td>8</td>
</tr>
<tr>
<td>Eye</td>
<td>4</td>
</tr>
<tr>
<td>Ear</td>
<td>2</td>
</tr>
<tr>
<td>Heart</td>
<td>1</td>
</tr>
<tr>
<td>Kidney</td>
<td>1</td>
</tr>
<tr>
<td>Total</td>
<td>16</td>
</tr>
</tbody>
</table>

Table (4): The other associated problems.

<table>
<thead>
<tr>
<th>Problem</th>
<th>No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Swallowing</td>
<td>68</td>
</tr>
<tr>
<td>Walking</td>
<td>65</td>
</tr>
<tr>
<td>Epilepsy</td>
<td>60</td>
</tr>
<tr>
<td>Speech</td>
<td>46</td>
</tr>
<tr>
<td>Intellect</td>
<td>42</td>
</tr>
<tr>
<td>Vision</td>
<td>32</td>
</tr>
<tr>
<td>DDH</td>
<td>31</td>
</tr>
<tr>
<td>Behavior</td>
<td>29</td>
</tr>
<tr>
<td>Hearing</td>
<td>10</td>
</tr>
</tbody>
</table>

Discussion

Similar to many previous reports, there is slight predominance of males compared to females (57% vs. 43%). The main reason of first presentation to any of the health centers in this study was the delayed motor developmental skills (50%). Certain developmental observations were often made first by the parents; suggesting a possible diagnosis of CP, the most frequent one being the delayed motor development often with a dissociation between motor and intellectual development.

Similar to previous reports, the commonest type of CP is the spastic quadriplegia that has been reported to occur in up to 80%. Both hand preference and difficulty in walking signify the underlying type of spasticity. One hand preference (especially below one year of age) signifies one side of the body involvement and the later signify spastic hemiplegia.

The last presenting complain in this study was the head lag which is the most common presenting feature of chorio-athetosis complicating bilirubin
encephalopathy, as hypotonia predominate in the first year of life then abnormal movements, as athetosis, occur later as myelination will have completed in the next 2 years. \(^{19}\)

Majority of the patients were children of parents who got together by consanguineous marriage; which may explain a genetic or familial predisposition. \(^{20}\)

Early diagnosis of CP had been done in 18% of cases below the age of 6 months because they were severe enough in addition to the availability of neuroimaging. \(^{21}\)

The vast majority of patients were diagnosed between the ages of 7-12 months (66%) because most of the CP patients are diagnosed by the end of the first year of life. \(^{16}\)

Improvement of the antenatal care and follow up of mother with high risk pregnancies did not reduce the total number of CP suggesting a non-obstetrical cause for CP. \(^{16}\) This also had been observed here because 76% of cases attended the antenatal clinic but still they had ended with babies with CP which may be explained by many reasons; the low availability of equipments for the sake of early diagnosis and detection, mismanagement during intra-partum period because of the lack of facilities necessary for fetal management, or because most of the CP cases are developmental in nature.\(^{1}\)

Majority of patients were delivered by normal vaginal delivery, while 15% were products of prolonged and difficult labor which is a known factor for the development of CP due to oxygen deprivation in the birth canal, and the possibilities of complications especially intracranial hemorrhage; which is also a risk factor for those 5% products of forceps delivery. \(^{17}\) 13% of patients were delivered by the cesarean section which did not prevent the future development of CP, this may be explained in part that in about 70% of cases of CP brain damage occurs before birth, although in the majority of cases it occurs around the time of delivery, or in the first month of life. In many cases, the cause of CP is not known yet. \(^{22}\)

Thirty six percent of patients were premature, which is an important risk factor especially when associated with low birth weight, primarily because of intra-cerebral hemorrhage and peri-ventricular leukomalacia.\(^{1}\)

Six patients only were term; of whom 5 were babies of diabetic mother’s. CP occurrence in these patients may be related to many factors such as macrosomia and related birth injuries, placental insufficiency, attacks of hypoglycemia and the increased incidence of congenital malformations in these babies.\(^{1}\)

Forty eight patients sustained asphyxia, of which 45 underwent resuscitation and were admitted to NICU. The APGAR score does not predict the neonatal mortality or subsequent CP and indeed the score is normal in most patients who subsequently developed CP although a score of 0-3 at 20 minutes is associated with high mortality and morbidity.\(^{1}\) Still, 20% of children with CP who had a normal birth weight and no recorded congenital malformation at birth had a 5 minute APGAR score below 7. \(^{23}\)

The high prevalence of birth asphyxia in out locality may have an effect on the growth and the development of the child. \(^{24, 25}\)

Majority of patients had history of neonatal jaundice which may develop in 60% of term and 80% of premature infants.\(^{1}\)

The most common type of CP encountered in the clinical practice is the spastic one ranging from 40% (14) up to 60% \(^{17}\) followed by the mixed type 9-22%. \(^{13}\)

The high prevalence of quadriplegia may be due to the high prevalence of birth asphyxia. \(^{19, 12, 26, 25}\)

Associated congenital malformations were comparable to other studies. \(^{27}\)

Other problems associated with CP were comparable to the published studies with the hearing problem being less.\(^{3}\) This is the first study highlighting the clinical profiles of CP children in Kurdish population.
References


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الشلل الدماغي لدى الأطفال الأكراد في مدينة داهوك: سلسلة من الدراسات

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الملخص

الهدف: تقييم الوضعيات الشعاعية لدى المرضى المصاصبين بالشلل الدماغي، وتقديم وجوه علامات شعاعية نوعية تميزها لكل طيف من أنماط مرض الشلل الدماغي.

طريقة الدراسة: أجريت الدراسة على عينيتين مكونتين من 158 مريضاً يعانون شللًا دماغياً، ويراجعون عيادات أمراض الأعصاب والتطور العضلي في مستشفى الملك المؤسس عبد الله الجامعي على مدى عامين (2007-2008)، حيث أجريت دراسة راجعة بالاعتماد على سجلات العيادات، والربط مع الوضعيات الشعاعية في تصوير الطيفي المختفي والرنين المغناطيسي.

النتائج: تم تحمل عينة مكونة 158 حالة بالدماغ في الدراسة كالآتي:
- 84 حالة (53%) من العينة كانوا من الذكور.
- 74 حالة (47%) من العينة كانوا من الإناث.
- 41 حالة (26%) من العينة المتشابحة بالدراسة ولدوا قبل الحمل (Preterm).
- 117 حالة (74%) من العينة المشمولة بالدراسة ولدوا بتمام الحمل (Full term).

وشكل الدراسة:
1- 112 حالة من الشلل الدماغي التشنجي (Spastic CP) ونسبة 70.8%، مع ملاحظة كون النمط التشنجي الرحبى صور
2- 63 حالة من الشلل الدماغي التشنجي الاشعة (Spastic quadriplegia) ونسبة 40% من العينة.
3- 22 حالة من الشلل الدماغي الرحبى الرحمي (hypotonic ataxic) ونسبة 14% من العينة.
4- 15 حالة من الشلل الدماغي التشنجي ذو طيف عضوية (Dyskinetic CP) ونسبة 9% من العينة المختلطة (Mixed CP).
5- 9 حالات من النمط المختلطة (1.5%).

وقد تم ملاحظة تغيرات شعاعية مرتبطة باستخدام الرنين المغناطيسي في 113 حالة، ونسبة 79%، باستخدام التصوير الطيفي في 113 حالة، ونسبة 26%، وكما تم ملاحظة وجود تغريزات شعاعية نوعية تشير إلى:
1- تقسيم الأكسسوات الدماغية (Brain Asphyxia) في 40 حالة، ونسبة 25%. ونسبة 14%.
2- الأماكن الدماغية الواقعة في 22 حالة، ونسبة 14%.
3- التغريز داخل الحصين في 10 حالات، ونسبة 6%.
4- اسكتما دماغية وجسمية في 29 حالة، ونسبة 88.5%.
5- تحسن الدماغ حو حل ]). ونسبة 20%.
6- أسباب غير معروفة: مثل: الضرور الدماغي المعرض، في 26 حالة، ونسبة 16.5%.

وبناءً عليه، فإن العامل المسبب الأكثر شيوعًا في الشلل الدماغي هو نقص الأكسجة الدماغية (Brain Asphyxia). يشبه تليف الدماغ
(periventricular encephalomalacia).

أضاف إلى ما تقدم أنه تم في هذه الدراسة اعتبار الضمور الدماغي موجودًا غير نواعي، علماً أنه يُشاهد بكثرة لدى المرضى المصابين بنمط عسرة الحركة (Dyskinetic CP)، وشملت الدراسة حزمة مرضى من أصل خمسة عشر، بنسبة 33%، كما شملت في النمط الشنحالي الرباعى 15 مريضًا من أصل 63، ونسبة 24% مقترنة بالأعمال الأخرى.

النتيجة: تأكيد أهمية دور التشخيص الشعاعي في فهم العوامل المسببة والأكلية الأمراضية، والنسب من العوامل المؤثرة التي يكون لها دور في الاستشارة الوراثية (مثل الشوهات).

البيانات الدالة: الشلل الدماغي، الموجودات الشعاعية، الأكراد، داهوك.

الكلمات المفتاحية: الشلل الدماغي، الموجودات الشعاعية، الأكراد، داهوك.