Epileptic and cognitive changes in children with cerebral palsy: an Egyptian study

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Background: Cerebral palsy (CP) is the most frequent cause of motor handicap among children.

Aim of the study: We aim to study the relation of epilepsy in children with CP to various risk factors that affect the development of seizures.

Patients and methods: In a cross-sectional, descriptive, population-based, case-control study, 98 children with CP (48 children with CP with epilepsy, and 50 children with CP without epilepsy) were compared with 180 children without CP or seizures. The children lived in two regions in Egypt: the Al-Kharga District–New Valley and El-Quseir city–Red Sea. These cases were subjected to meticulous neurological assessment, brain magnetic resonance imaging, electroencephalography, and Stanford-Binet (4th edition) examination. Multinomial logistic regression was used to assess the risk factors.

Results: Epilepsy was diagnosed in 48.9% of all cases of CP. Mental subnormality was observed more frequently in children with epilepsy than in those without epilepsy (84.6% versus 66.7%). The frequency of epilepsy was highest in patients with the spastic quadriplegic type of illness (58.3%). Multinomial logistic regression revealed that prematurity (<32 weeks of pregnancy), low birth weight (<2.5 kg at birth), neonatal seizures, jaundice, and cyanosis were significantly associated with CP with epilepsy.

Conclusion: CP is associated with a high percentage of seizure disorders. Prematurity, low birth weight, neonatal seizures, cyanosis, and jaundice are significant risk factors among patients with CP with epilepsy compared to patients with CP without epilepsy or a healthy control group.

Keywords: cerebral palsy, epilepsy, risk factors, Egypt

Introduction

Cerebral palsy (CP) is the consequence of nonprogressive damage to the developing brain and consists of a number of clinical neurological syndromes of heterogeneous etiology. Epilepsy is one of the most common problems among patients with CP. It has been used as a marker of severity and often worsens the quality of life of patients with CP.

Epilepsy occurs in 15%–60% of children with CP; however, its clinical course is not well-defined. In our previous study at Al-Kharga District–New Valley, active epilepsy was recorded in 27 (51.9%) of 52 children with CP. The percentage of those with epilepsy was highest in patients with quadriplegia and lowest in patients with spastic diplegia and dystonic CP.

Early prediction of intractable epilepsy would be valuable because rapid identification of patients at the highest risk level would allow physicians to consider them...
for earlier treatment with recent medications and other surgical and nonsurgical treatments, such as vagus nerve stimulation.\textsuperscript{8,9}

According to our knowledge, this study was the first study carried out in Egypt. We aimed to study the frequency of epilepsy among children with CP and various risk factors that affect the development of seizures.

**Patients and methods**

**Study area**

This study was carried out in two different areas in Egypt. The first was New Valley, which represents about 44\% of the area of Egypt. It is divided into the El-Kharga, El-Farafra, and El-Dakhla districts. El-Kharga is far away from the Nile Valley and has different geographic, social, economic, and cultural characteristics. The second area was El-Quseir city, one of largest cities of the Red Sea Governorate. Its surface area is about 119,000 km\(^2\), and it lies directly on the Red Sea.\textsuperscript{6,10}

**Sample**

In a cross-sectional, descriptive, population-based and case-control study, 98 children with CP (with or without epilepsy) were compared with 180 children without CP or seizures. The children were collected from two regions in Egypt, Al-Kharga District–New Valley and El-Quseir city–Red Sea, through two large projects carried out to detect the epidemiology of major neurological disorders.\textsuperscript{11,12} All inhabitants who have been living in Al-Kharga District–New Valley (25,540) and El-Quseir city (33,285) for at least 6 months at the time of the interview were included as a sampling frame to identify the potentially eligible population of children and adults with CP in defined areas.\textsuperscript{6,13} Screening of all households door to door (including every door) was done using a simple standardized Arabic screening questionnaire designed specifically for this study to pick up different neurological disorders, including CP cases.\textsuperscript{12} The control group was chosen from same population with matched sex, age, education level, and sociodemographic\textsuperscript{14} state.

**Definitions**

CP was defined as motor disabilities caused by nonprogressive damage to the developing brain.\textsuperscript{13} CP is divided into the spastic, dyskinetic, and ataxic subtypes.\textsuperscript{16} The spastic subtype is further divided into unilateral (limbs on one side of the body are involved) and bilateral (limbs on both sides of the body are involved) types. The spastic bilateral type was further subdivided into quadriplegia and diplegia.\textsuperscript{6}

Epilepsy was diagnosed according to Commission of Classification and Terminology of the International League Against Epilepsy.\textsuperscript{17} Active epilepsy was considered when two or more unprovoked seizures occurred during the previous year.\textsuperscript{6,16}

The Standardized, Valid Arabic Version\textsuperscript{18} of the Stanford-Binet scale (4th edition)\textsuperscript{19} was used to assess intelligence quotient (IQ). Total IQ was classified according to Melika\textsuperscript{18} into mentally retarded including those with IQ $>$ 67, slow learner (IQ =68–78), below-average intelligence (IQ =79–88), average intelligence (IQ =89–110), above-average intelligence (IQ =111–120), excellent intelligence (IQ =121–131), and genius (IQ =132). The Stanford-Binet scale was done for 60.2\% (n=59) of case patients and 22.2\% (n=40) of control patients. The children with CP or from the control group were chosen randomly, as every fourth child, and if that child refused, the second child was chosen.

Interictal electroencephalography (EEG) and computed tomography (CT) head scans or cranial magnetic resonance imaging (MRI) findings were evaluated. Cranial CT was performed with 1 cm thick slices using Hitachi Radix Turbo (Hitachi Ltd, Tokyo, Japan), brain MRI was used to acquire T1- and T2-weighted axial and coronal and T1-weighted sagittal images, using Picker 1-Tesla Vista HPQ (Picker International, Cleveland, OH, USA).

The following data were ascertained: type of CP, type of epilepsy, age, sex, age of onset of seizures, and sociodemographic\textsuperscript{14} state.

A structured questionnaire also was filled out face to face, and mothers’ past medical and obstetrics histories were obtained. Available birth records of the children with CP were reviewed, including information regarding the health of the mother prior to the delivery and during pregnancy, family history of seizures, and the external factors occurring during the pregnancy and delivery. Specific complications of pregnancy encompassing abortion, hemorrhage in late pregnancy, premature rupture of membranes, and preterm labor (<32 weeks of pregnancy) were examined. Additional variables were examined in relation to labor, including prolonged labor and meconium staining of liquor. In addition, postpartum variables, including low-birth-weight babies (<2.5 kg at birth), birth asphyxia, prolonged jaundice, neonatal sepsis, other infections, neonatal seizure, and trauma, were examined.

EEG date and CT head scans or cranial MRI findings were evaluated. Cranial CT was performed with 1 cm thick slices using Hitachi Radix Turbo, and brain MRI was performed to acquire T1- and T2-weighted axial and coronal and
T1-weighted sagittal images, using a Picker 1-Tesla Vista HPQ. Then the patients were divided into three groups: epileptic patients with CP, nonepileptic patients with CP, and a healthy control group (without CP or seizures). Those groups were compared on the basis of clinical types, EEG, and cranial imaging findings.

Ethics
The Ethics Committee of Assiut University in Egypt approved the two studies (projects). Informed written consent was obtained from the Health Institute and from each patient’s family (responsible person in each family).

Statistical methods
SPSS for Windows version 16 (IBM Corporation, Armonk, NY, USA) was used for data analysis. Prevalence rates were presented with binomial exact 95% confidence intervals. The chi square test, independent sample t test, or one-way ANOVA test, followed by a post hoc test, were used to analyze differences in proportions between groups. Multivariate logistic regression was used to detect the influence of different risk factors for the development of epilepsy. Odds ratio and 95% confidence interval were calculated from the model. All P values were two-sided, and statistical significance was defined as P<0.05.

Results
Ninety-eight patients (52 boys, 46 girls) with CP were recruited. They included 48 patients (48.9%) who had CP with epilepsy and 50 (51.02%) CP patients without epilepsy compared with 180 healthy children. Other factors associated with the occurrence of epilepsy in children with CP were also analyzed (Table 1). Mental subnormality (mentally retarded, including those with IQ >67) according to the Stanford-Binet scale was observed more frequently in children with epilepsy than in those without epilepsy (84.6% versus 66.7%). Sex, abnormal neuroimaging results, and family history of seizure disorders were not associated with the development of epilepsy, but neonatal seizure was significantly higher among the patients in the CP with epilepsy group than other groups. Abnormal MRI results included brain atrophic and white matter changes.

The frequency of epilepsy was highest in patients with spastic quadriplegic type disease (58.3%; Table 2).

Multinomial logistic regression was used to assess the relationship between risk factors and CP with epilepsy (Table 3).

Discussion
Epilepsy affected 48.9% of children with CP in the present study. These results were similar to those from other studies.\textsuperscript{2,5,7,8,20} This was explained by the fact that children with CP might suffer extensive brain injury including to the cortex, deep white matter, and central nuclei, and therefore they are liable to develop epilepsy.\textsuperscript{16}

In the present study, the frequency of epilepsy varied with the different forms of CP (58.3% for spastic quadriplegic, 27% for mixed, 8.3% for spastic hemiplegic, 4.2% for dystonic type, and 2.1% for spastic diplegic). The frequency of epilepsy varied by CP subtype. When comparing and/or

| Table 1: Demographic data of children and factors associated with development among CP |
|---------------------------------|----------------|----------------|----------------|---|
| Factor                          | CP with epilepsy (n=48) | CP without epilepsy (n=50) | Control without CP or epilepsy (n=180) | P-value |
| Age, mean ± SD                  | 7.97±4.741 | 7.91±4.821 | 7.19±2.712 | 0.248 |
| Sex, M/F                        | 28/20 | 24/26 | 86/94 | 0.416 |
| Abnormal MRI                    | 22 (45.8%) | 21 (42%) | - | 0.242 |
| Positive family history         | 3 (11.1%) | 1 (4%) | - | 0.335 |
| Neonatal seizures               | 6 (12.5%) | 1 (2%) | 5 (2.8%) | 0.009 |
| Total IQ                        | 22 (84.6%) | 22 (66.7%) | 0 | 0.0001 |
| Mentally retarded (<67)         | 2 (7.7%) | 6 (18.2%) | 7 (17.5%) | 0.05 |
| Slow learners (68–78)           | 0 | 0 | 0 | 0.248 |
| Below-average intelligence (79–88) | 0 | 0 | 0 | 0.0001 |
| Average intelligence (89–110)   | 2 (7.7%) | 3 (9.1%) | 26 (65%) | 0.335 |
| Above-average intelligence (111–120) | 0 | 0 | 0 | 0.0001 |
| Excellent intelligence (121–131) | 0 | 0 | 6 (15%) | 0.0001 |

Notes: Unless otherwise indicated, data are expressed as mean ± SD or number and percentage done. IQ was performed for 26 patients with CP with epilepsy, 33 patients with CP without epilepsy, and 40 control patients. Abnormal MRI refers to brain atrophy and white matter changes.

Abbreviations: MRI, magnetic resonance imaging; M, male; F, female; CP, cerebral palsy; IQ, intelligence quotient; SD, standard deviation.
pooling data between several registers, Gururaj et al reported that spastic tetraplegia was the commonest type of CP with epilepsy, and spastic diplegia was the commonest type of CP in a group of patients with CP without epilepsy. Surveillance of Cerebral Palsy in Europe collaborators reported that children who were described as having diplegic or tetraplegic spastic CP were more likely to have epilepsy.

Epidemiological study has revealed that cognitive impairment is a very common occurrence in CP children. Epilepsy in children with CP also has been related to impaired intellectual performance. In the present study, mental retardations were reported in 84.6% in patients with CP with epilepsy versus 66.7% among patients with CP without epilepsy. That matched with other studies. Vargha-Khadem et al reported that in patients, the presence of epilepsy is clearly associated with more severe and increasing cognitive difficulties, and Süssová et al reported that mental subnormality was more common in children with epilepsy than in those without epilepsy. Again, that could be explained by the fact that children with CP might suffer extensive brain injury including cortex, deep white matter, and central nuclei, and therefore are vulnerable to mental retardation and epilepsy.

Hence, an early intervention with treatment of epilepsy prevents further progression of neuronal injury, with subsequent cognitive impairment caused by uncontrolled seizures, among those patients.

Data regarding additional perinatal factors that might contribute to the pathogenesis of seizures in patients with CP are relatively scarce, and most studies refer to either CP in general or epilepsy in the general population, rather than to epilepsy in children with CP. The presence of prematurity and low birth weight was strongly predictive for the subsequent occurrence of epilepsy among CP children in the present study. That was matched with that fact that these studies consistently show that low birth weight and prematurity are risk factors for both epilepsy and CP.

With regard to neonatal seizures, in these studies, their presence was prediagnostically associated with occurrence of epilepsy among CP cases. A strong association of neonatal seizures with epilepsy has been already reported by Nelson and Ellenberg, during the Collaborative Perinatal Project of the National Institutes of Health, summarizing 54,000 singleton pregnancies between 1959 and 1966. Subsequently, additional retrospective studies clearly demonstrated that in children with CP, a history of neonatal seizures was strongly predictive for future development of epilepsy. The presence of neonatal seizure was considered to be a factor for subsequent development of neurological disabilities (mental retardation, CP, or epilepsy). Levene reported that neonatal seizures had an adverse effect on neurodevelopmental progression and may predispose to cognitive, behavioral, or epileptic complications later in life.

In this study, cyanosis of child at birth and neonatal jaundice were found to be significant risk factors for developing epilepsy in these children with CP when analyzed using multivariate logistic regression. We considered cyanosis to be a picture of respiratory distress in children because we have no sufficient data or recorded reports for these children. However, respiratory distress was significantly associated with the occurrence of CP. In addition, apnea requiring ventilation therapy appeared to be associated with the occurrence of CP, although it was not assessed well in this study. Prolonged jaundice may be responsible for CP, although jaundice attributed to presence of meconium staining of the amniotic fluid is more likely to indicate a sign of intrapartum hypoxia and was more common in the CP cases.

These risk factors, which might affect the development of seizures, lead to more deterioration in CP cases, although it is nonprogressive disease.
Conclusion
Epilepsy among children with CP is common and represents 48.9% of children with CP. The observed relationship between epilepsy and associated impairments was as predicted. However, it requires longitudinal studies of children with CP and epilepsy to further improve our understanding of the relationship between epilepsy and CP. In addition, further studies concerning a possible risk of epilepsy development and its relationship with the EEG and cranial imaging findings may help in presenting the other risk factors involved and the factors affecting the prognosis of CP and understanding of the cause of epilepsy in CP.

Disclosure
The authors report no conflicts of interest in this work.

References