Identifying pattern in global developmental delay children: A retrospective study at King Fahad specialist hospital, Dammam (Saudi Arabia)

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Abstract

Global developmental delay (GDD) and intellectual disability are relatively common in pediatric neurology conditions. A retrospective study was designed to analyze risk factors and clinical features in children with GDD at our hospital. No previous data is available on GDD from Saudi Arabia. This study was conducted at King Fahad Specialist Hospital Dammam (KFSHD) of 134 GDD children (82, 61% males, 52, 39% females), (age 1-9 years). They were assessed by using Griffith Mental Development Scales for (0-2) years and 3-8 years old in locomotors, personal/social, communication, eye & hand co-ordination, performance and practical reasoning. Patients with ASD and non-cooperative behavior were excluded. 75% had developmental delay since birth while 84% had no problem during pregnancy. 22% had birth weight below 2.5 kg, 56% had epilepsy and 57% had family history of GDD. 51% were diagnosed cases in the present study. 40% had genetic cause, 25% had metabolic problem, 58% had neuroradiology abnormalities and 45% had EEG abnormalities. There a variety of delays in development (speech and language variant, global delay, and the motor variant) noted and are commonly seen in a clinical practice in KFSHD. Longitudinal research beginning in early development will help to understand the developmental domains and neurological comorbidities in these children at high risk for neurodevelopmental disorders.

Introduction

Global developmental delay (GDD) is term used for children under 5 years of age. It is defined as a significant delay in two or more domains of development, including activities of daily living as well as motor, cognitive, speech/language, and personal/social skills.1,2 The clear diagnosis of GDD is problematic because of its heterogeneous etiology; thus, the causes of GDD are undetermined in approximately 62% of children.3 The exact prevalence of GDD is not known precisely, however from different studies it is estimated 1-3%.1,3,4 It is probably higher in Saudi Arabia due to high rate of consanguinity. Individuals in whom genetic causes are established generally undergo very exhaustive, expensive, and often invasive diagnostic evaluations, even though the results may not change the medical or therapeutic management of the delay.

Significant controversy still surrounds the symptom complex of GDD in term of diagnosed in children under the age of 5 years. Since formal psychometric testing often cannot be reliably undertaken in the young child a clear-cut, objective threshold for diagnosis, as in the case of cognitive delay, often cannot be clearly established.4,5 Therefore, GDD diagnosis relies on the summation of clinical findings in several developmental areas. Another cause for controversy is the apparent wide variability of clinical presentations that fall under the GDD diagnostic umbrella. Term Learning difficulty/disability is used for children older than 5 years of age. Mild learning difficulty (IQ 50-70) affects 1-2% of the child population. Severe learning difficulty (IQ < 50) affects 0.3-0.5% and may be subdivided into moderate (IQ 35-49), severe (IQ 20-34) and profound learning difficulty (IQ < 20).6

Although on established guidelines, studies were conducted in North America, Europe and Turkey.7,8 Not much data is available about the GDD in the Gulf or local populations. This study was conducted to determine the etiologic yield for GDD in children attending a tertiary care hospital in this part of the world. To address these questions, we evaluated clinic referred subject in KFSHD by correlating detail clinical history, examination, developmental assessment, behavioral assessment, and results of relevant investigations in a developmental clinic.

Materials and Methods

A detailed retrospective chart review of 134 children diagnosed with GDD in pediatric neurology and development assessment clinic at King Fahd specialist hospital, Dammam (KFSHD-D) between 1 Feb 2016 and 1 April 2018 was undertaken (i.e. 1-9 years). Permission was obtained from the local ethic committee of KFSHD. Children were identified by review of the charts of all patients assessed in the Developmental Progress Clinic during the time period studied. Referrals to the clinic derive mainly from pediatricians, family physicians and pediatric neurology in the eastern province region. The population seen reflects a wide range of local ethnic and socio-economic backgrounds. All children referred to the clinic undergo a formal multidisciplinary developmental assessment (complete history, and developmental pediatrics examination). A diagnostic conclusion from these evaluations is drawn after the first or second visit to the clinic.

A diagnosis of GDD was ascribed to all children presenting with significant delays (defined as performance is below the mean on age-appropriate, standardized, norm-referenced testing) in two or more developmental domains. As part of standard clinical protocol, all children found to have GDD undergo a formal cognitive assessment with a trained pediatric physician or clinical psychologist.

Assessment scale

These children were developmentally assessed by using Griffith mental development scales for 0-8 years old in Locomotor, personal/social, Hearing and language, eye

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and hand co-ordination, performance and practical reasoning scales.

Patients with autism and non-cooperative were excluded.

A clinical sheet was designed by the research group to get detailed history, particularly birth, developmental, family, and relevant past medical including epilepsy and medication history.

A considerable part of sheet was assigned for clinical examination particular neurological examination and development assessment in locomotor, personal/social, hearing and language, eye & hand co-ordination, performance and practical reasoning scales. Severity of development delay in each domain was labelled as mild, moderate and severe categories according to the developmental age assessed by developmental pediatrician using Griffith scale compared with actual chronological age (i.e.; mild: 67-100%; moderate: 33-66%; and severe <33%).

The results of investigations like genetic, metabolic, neuro-imaging and EEG were collected from medical records.

Inclusion and exclusion criteria

Inclusion criteria: children of 1-9 years of age presenting with development delay.

Exclusion criteria: patients with autism and non-cooperative for development assessment. Children whose development level was above 8 years of age as Griffith scale measures development age only up to 8 years. Absence of the complete data set of the patient as designed in the clinical evaluation sheet.

Statistical analysis

Developmental scores were recorded as centiles for the following variables fine motor performance, expressive vocabulary, and receptive language. All data gathered were analyzed using SPSS software version 22.0 (SPSS Inc., Chicago). A P-value of 0.05 or less was considered to be statistically significant.

Results

A total of 134 children initially diagnosed with GDD were identified over the study inclusion criteria. The age range of children was between 1-9 years (Figure 1). Of these, 134 children (82 males, 52 females, Figure 2). Among all 89% had developmental delay/learning difficulties including 11% had behavioral problems.

The data derived from their mother pregnancies were normal (n=113, 84%), while rest (n=21, 16%) showed (n=6, 4.7%) bleeding, (n=3, 2.2%) decreased fatal movements, (n=3, 2.2%) polyhydromnitos, (n=3, 2.2%) diabetes mellitus, (n=3, 2.2%) hypertension/pre-eclampsia and (n=2, 1.4%) epilepsy problems. During the labor (n=128, 96%) had spontaneous onset delivery. The data showed (n=112, 84%) normal vaginal delivery, (n=20, 15%) had caesarean section, while (n=2, 1%) had ventouse delivery.

These kids (n=30, 22%) had birth weight below 2.5 kg, (n=18, 13%) needed resuscitation and (n=14, 10%) were intubated for few days. During the neonatal period, 11 patients had neonatal seizures, 6 had sepsis, 18 had severe jaundice and 14 of them needed phototherapy. Only (n=20, 15%) children had feeding difficulties and needed

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<th>Table 1. Underlying etiology in global developmental delay.</th>
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![Figure 1. Number of cases by age.](image)

![Figure 2. Number of cases by gender group.](image)
feeding assistance. Family history showed 57% had consanguinity (Figure 3). 25% had epilepsy in close relatives, 16% had learning disability in family, 20% had relatives went to special school, 7% had mental health problems in the family and 6% had hearing difficulties in family. GDD kids’ group (56%) showed various types of epilepsy (Figure 4).

**Developmental assessment**

The developmental assessment showed GDD particularly in speech and locomotors, while 21% had development regression and 79% had development delay since birth.

Overall severe development delay was 47%, moderate delay 18%, mild delay 16% and normal 6% in this group data. In locomotor assessment task- severe development delay was (n=37, 28%), moderate delay (n=22, 16%), mild delay (n=24, 18%) and (n=11, 8%) showed normal function of locomotor in study group (Figure 5).

For the personal/social assessment, the severe development delay was in (n=32), moderate delay (n= 27) mild delay (n=21) and normal (n=12) in this study group (Figure 5). For the hearing and language assessment, (n=33) were severely delayed, (n=25) moderately delayed, (n=21) mild delay and normal (n=7) in task.

The data showed (n=32) were severely delayed, (n=23) had moderate delay, (n=23) had mild delay and normal (n=8) in Eye and hand coordination assessments (Figure 5).

Performance task showed (n=30) had severe developmental delay, (n=24) had moderate delay, (n=20) were mildly delayed and (n=8) were within normal range during performance task (Figure 5).

Growth parameters showed 23% had head circumference below 3rd. centile. While 42% of our patients had some form of dysmorphic features. 15% of our patients showed neurocutaneous abnormalities.

**Diagnosis**

Sixty-nine (51%) out of 134 were clinically diagnosed by neurologist (Figure 6). Genetic disease being the most common 41%, Metabolic abnormalities were found in 25%, Neuro radiological abnormalities were found in 57% and EEG abnormalities were found in 45% cases (Table 1).

**Discussion**

This study was hospital based and does not represent the community-based study. There is possibility of bias towards working up of more affected children. The 51% children with GDD, which was within the range
reported from previous studies. Three previous studies from a single center in North America had reported a yield of 63%, 55%, and 38%. In a retrospective study from Turkey on 247 patients with GDD, etiology was found in 63%.[10] In a study from Jordan, the rate was identified in 44.5% only.[11] These differences in the yield are explained by the differences in the population, criteria used, and the extent of investigations. Another factor which has a major impact on detection of the underlying cause is the high incidence of consanguineous marriages resulting in familial cases in this part of the world.[11] Few studies showed half of the children with delays, especially language-related delays, are not detected before starting school and do not use a standardized instrument for neurodevelopmental assessment.[12-13] Recent guidelines strongly recommend the systematic screening of developmental delay in young children, in order to provide early beneficial interventions to the child and family.[7-10]

The diagnostic work-up with regard to perinatal history, perinatal data, and clinical examination was complete in 94 to 99%, and laboratory investigations in 80 to 93% of the 134 children ascertained for analysis. Perinatal asphyxia, Genetic, metabolic diseases and cerebral digenesis were the common causes of GDD, while familial consanguinity, dimorphism and non-specific white matter changes in MRI guided more than 50% of the etiology of GDD. This distribution was similar to reports from previous studies.[7-9]

Developmental delay is a symptom not a diagnosis. It is very important to listening to parents, paying special attention to their concerns, and doing a meticulous history makes the greatest contribution to a neurodevelopmental diagnosis.[12,14] It is clear that establishing a diagnosis enables us to answer questions on: why it has happened (etiology), what does it mean for our child (prognosis), what treatments might be available (precision medicine) and whether it can be prevented in the future (prenatal and preimplantation genetic diagnosis). Major malformations for syndrome identification should be obvious to the clinician on physical examination.[15-17] Careful neurodevelopmental examination should include observations of minor neurological signs, associated movements, fine motor skills, and gait.

A diagnosis guides the clinician in looking for rare but treatable diseases and helps predict the risk of a similar condition occurring in future pregnancies.[16-18] Helping a family organize a plan to address children developmental delay is essential. Directing the child to early intervention services is a sound first start. Any child less than 3 years of age who has a physical or mental condition that has a high probability of resulting in developmental delay is eligible for services, so a neurodevelopmental diagnosis helps qualify a child for early intervention services.[19,20] Services are mandated for infants, toddlers and preschoolers. Although eligibility criteria vary by jurisdiction, common measures of delay are 25% and/or two standard deviations in one or more of the developmental domains.[20] Recognizing developmental patterns that guide a neurodevelopmental assessment is essential to help families get the support services they need.[20]

Conclusions

The study gives a baseline data about the spectrum of GDD in the region. In addition, specific screening investigations should be obtained on the basis of clinical suspicion in order to improve the yield. All patients with developmental delay/learning disability need a comprehensive medical evaluation by complete medical history with particular detail of birth, early development and family history. Physical examination with particular emphasis on growth parameters, dysmorphism, neurocutaneous stigmata, systemic and neurological examination. Development assessment by using standardized development assessment scales. If the clinical diagnosis is obvious or suspected, confirm the diagnosis with appropriate genetic testing. If diagnosis is unknown and no clinical diagnosis is strongly suspected, begin in stepwise evaluation by: chromosomal microarray; Fragile X testing; for females complete MECP2 gene study; metabolic testing; brain MRI if clinically indicated; EEG if epilepsy.

Whether diagnosed or not, results and their implications should be carefully explained to the parents/care givers. Appropriate support and referral to appropriate support services should be made.

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