Screening for Neurodevelopmental Delay for Preterm Very Low Birth Weight Infants at Tertiary Care Center in Saudi Arabia

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Abstract

Background

Preterm infants are more susceptible to death, short-term complications, and long-term complications such as neurodevelopmental impairments. However, definitive assessment tools are not available in a resource-limited setting. Hence a screening tool is needed the Arabic-speaking population.

Method

Infants born at a gestational age of <32 weeks or a very low birth weight (VLBW) of less than 1500 g were recruited into a cross-sectional study. We identified infants (n ~ 61) admitted to the neonatal ICU at King Abdulaziz Medical City and reached 18 up to 24 months of corrected gestational age (CGA). The developmental assessment was done at 18, 20, 22, and 24 CGAs using the Ages and Stages Questionnaire third edition - Arabic version (ASQ3-A). The primary outcomes are early detection rate of neurodevelopmental delay (NDD), defined as a delay in one or more of the following: communication, gross motor, fine motor, problem-solving, and personal-social skills as per ASQ3-A.

Results

Sixty-one out of 92 eligible infants (36 excluded) completed the sufficient assessment. Twenty-six infants (42.6%) had at least one NDD in one of the following domains: communication skills: (11.5%), gross motor: (11.5%), fine motor: (19.7%), problem-solving skills: twelve infants (19.7%), and personal-social skills: twenty infants (23%). Perinatal events and periventricular leukomalacia (PVL) were significant independent predictors for the NDD.

Conclusion

This single-center study in Saudi Arabia screened preterm, VLBW infants based on ASQ3-A, twenty infants (42.6%) had an abnormal NDD at a corrected age of 18-24 months. Perinatal events and PVL were independent predictors of NDD. We recommend that all preterm VLBW infants in Saudi Arabia be evaluated by a neurodevelopmental screening tool, ASQ3-A, especially in resource-limited settings to start early intervention. Also, more extensive multicenter studies are to be carried out with definitive diagnostic tools to have a national benchmark for the long-term neurodevelopmental impairment.

Introduction

The WHO reported that 15 million babies are born prematurely, born before 37 weeks, worldwide every year [1]. Preterm is classified into three subcategories, moderate or late preterm (from 32 to less than 37 weeks), very preterm (from 28 to 32 weeks), and extremely preterm, which is <28 weeks [1-4]. Moreover, they are born with very low birth weight (VLBW), under 1,500 gm, or extremely low birth weight (ELBW), below 1,000 gm [5-7]. Preterm births and VLBW have been associated with risk factors such as maternal infections, multiple pregnancies, genetic factors, and chronic conditions such as hypertension and diabetes [1,8]. Death is the most severe outcome, which is inversely proportional to the infants' gestational age and birth weights [3,9]. This approximately accounts for 1 million premature deaths each year [1]. As more preterm infants survive, they are more susceptible to short-term morbidities acquired during their stay in neonatal intensive care units (NICU) and long-term complications [9,10]. Such maternal or short-term morbidities may affect the long-term complications, namely neurodevelopmental impairments (NDI), which may have lifelong
impacts on preterm infants [5,10-12]. Neurodevelopment, defined as the development of the CNS and the neurological processes, affects tasking and normal performance like gross motor development [15,14]. Thus, NDI mainly affect vision, hearing, social skills, learning ability, behaviors, and other aspects of life [15]. Such impairments might be aggravated by developmentally supportive therapies access, socioeconomic status of the family, and lifelong complications that necessitate feeding tubes and oxygen supplementation [3,16].

The routine neurodevelopmental assessment for preterm VLBW infants is carried out around the world. It includes multidisciplinary, longitudinal, and extensive evaluation which is the Bayley Scales of Infant Development [17]. It is costly, lengthy, and requires trained personnel; hence, the Ages and Stages Questionnaire, version 3 (ASQ3), is a useful alternative tool that can effectively screen developmental delay in limited-resource settings [18]. It is quick, inexpensive, easy to administer and interpret, and is accepted by the American Academy of Pediatrics as a valid developmental screener tool [19]. A new Arabic version was developed (ASQ3-A) and being used [20].

There is a lack of studies that have been published in Saudi Arabia on monitoring preterm infants and conducting their long-term neurodevelopmental outcomes. Moreover, research is required to develop standardized values for health care workers to understand the local incidence of neurodevelopmental delay (NDD) among preterm children. This study aims to measure the rates for NDD in infants born preterm, VLBW using the ASQ5-A tool in a local tertiary care center in Saudi Arabia. In addition, this research will provide neurodevelopmental values that can assist parents with preterm infants in perinatal counseling and assist health care workers.

Materials And Methods

Method

This study is a cross-sectional design conducted at King Abdulaziz Medical City (KAMC), Jeddah, Saudi Arabia, seen between June 1, 2019 and December 30, 2020. It included all infants born at the gestation of <32 weeks or VLBW, admitted to NICU at the KAMC, and reached the age of 18 to 24 months corrected gestational age (CGA). Infants who had documented chromosomal abnormalities, significant congenital anomalies, inherited inborn errors of metabolism, or severe perinatal asphyxia, were excluded from the study.

The specific objectives were to assess the applicability of the Ages and Stages Questionnaire (ASQ3- A) to early NDD, defined as a score of less than one SD (1SD) of one or more of the following: communication, gross motor, fine motor, problem-solving, and personal-social domains as per ASQ3-A scoring. The screen of NDD is carried out at 18, 20, 22, and 24 months CGA in the respective cohort. The secondary objective was to examine for possible confounding factors to neurodevelopmental outcomes. The patients were identified from the NICU logbook of admission, and the maternal and infant data were extracted from the electronic health records (BESTCare). Consents were obtained from one of the legal guardians prior to data collection. The study was approved by the local IRB at King Abdullah International Medical Research Center with a reference number SP18/189/1.

Data collection

The involved staff was trained to administer the ASQ3-A to help parents fill up the questionnaire and answer any queries. A senior specialist and a neonatologist supervised them. We interviewed the parents to collect developmental milestone changes as per the ASQ3-A. ASQ3-A is a questionnaire that is used effectively to screen any developmental delay in the first five and half years of a child’s life. The five domains of the ASQ3-A developmental screening areas are communication, problem-solving, fine motor, gross motor, and personal-social. The ASQ3-A was divided into two parts. The first part was done by interviewing the parents via phone call, and the second part was done through an online survey after parents observed fine motor and problem-solving skills. After completing the questionnaires, the scores were recorded. Then, raw data was entered into the data collection sheet.

ASQ3-A is a valid and simple screening tool used confidently to distinguish and exclude neurodevelopmental impairment as its sensitivity, and the negative predictive value approaching 100%, its specificity was acceptable (76%) [21]. However, sensitivity to detect infants with impairments ranged from 20 to 60% [22]. ASQ3-A is a suggested screening tool in routine monitoring of low-risk children due to its psychometric properties. Also, it is recommended to use ASQ3-A in the follow-up programs for preterm infants and children who have biological risk factors [23]. ASQ3-A was translated and verified by experts from different Arabic-speaking countries, including Saudi Arabia, Egypt, and Kuwait, to assure that the translation is accurate and accessible [24].

Data analysis

Data were entered into SPSS version 27 or SADA version 15.1. Frequency, percentages, and bar charts were used to represent the qualitative data. The quantitative data were described using the mean and SD or median and quartiles as per normality. The reason why we defined NDD as a score of less than 1SD of one or more of the five domains of ASQ3-A is to detect more premature children with delays as suggested by
Simard MN et al. [22]. For univariate analysis, we used Chi-squared, Fisher’s Exact, or t-test accordingly. We used Firth’s logistic regression as a standard approach for analyzing binary outcomes with small samples to assess for significant predictors. Whereas it reduces the bias in maximum likelihood estimates of coefficients, bias towards one-half is introduced in the predicted probabilities [25]. A p-value of 0.1 is considered statistically significant.

**Results**

The study screened 128 preterm infants of less than 32 weeks gestational age or VLBW. Thirty-six preterm infants were excluded; 23 infants died before discharge, and 13 reported chromosomal abnormalities or major congenital anomalies. After exclusion, 92 infants were eligible; 13 did not respond, while 18 lost to follow-up either because they did not complete the online survey or exceeded the age limit due to the COVID-19 pandemic. There were no reported deaths after NICU discharge. Finally, 61 preterm infants completed sufficient assessment to be included in the study’s primary outcomes (Figure 1). There were 18 (10%), 16 (26.2%), 12 (19.7%), and 23 (37.7%) who completed the assessment around 18, 20, 22, and 24 months corrected age, respectively.

![Patient's flow](image)

**FIGURE 1: Patient’s flow.**

VLBW: Very low birth weight; NICU: Neonatal intensive care units.

Based on ASQ3-A, at the corrected age of 18-24 months; 35 infants (57.4%) had a normal neurodevelopmental screen, and 26 infants (42.6%) had at least NDD (<1SD) in one of the five domains which are communication skills, gross motor, fine motor, problem-solving skills, and personal-social skills.

A summary of the maternal and neonatal demographics of the final sample is shown in Tables 1 and 2, respectively. Of the maternal characteristics, only perinatal events (defined as any preterm premature rupture of the membranes [PPROM], antepartum haemorrhage [APH], or placental abruption) were statistically significant (4 [11] vs. 10 [39%] in children without vs. with NDI, respectively, p-value: 0.05). Of the neonatal characteristics, there were almost similar demographics in terms of gender: boys, 33 (54.1%) and girls, 28 (45.9%), gestational age at birth, and birth weight. During NICU stay, the included infants had similar morbidities except for the development of periventricular leukomalacia (PVL) (0 vs. 6 [23%], p-value: 0.03) (Table 2).
| Maternal characteristics                  | No NDD 35  | n (%) | With NDD 26 | n (%) |
|-------------------------------------------|------------|-------|-------------|-------|
| Antenatal care                            | 21 (62)    |       | 19 (73)     |       |
| Antenatal steroids                        | 30 (86)    |       | 23 (88)     |       |
| Hypertension                              | 7 (20)     |       | 2 (8)       |       |
| Preeclampsia                              | 7 (20)     |       | 2 (8)       |       |
| Diabetes                                  | 5 (14)     |       | 4 (5)       |       |
| C-S                                       | 27 (77)    |       | 19 (73)     |       |
| Perinatal events (PPROM, Abruptio, APH)   | 4 (11)     |       | 10 (39)*    |       |

**TABLE 1: Maternal characteristics.**

PPROM: Preterm premature rupture of the membranes; C-S: Caesarean section; APH: Antepartum haemorrhage.

* P-value <0.05
## Neonatal characteristics

|                  | No NDD 35 | n (%) | With NDD 26 | n (%) |
|------------------|-----------|-------|-------------|-------|
| **Male**         | 21 (60)   |       | 12 (46)     |       |
| **Female**       | 14 (40)   |       | 14 (54)     |       |
| **GA mean (SD)** | 29.5 (2.1)|       | 28.6 (2.6)* |       |
| **GA =>28 weeks**| 27 (77)   |       | 16 (62)     |       |
| **GA <28 weeks** | 8 (23)    |       | 10 (39)     |       |
| **BW mean (SD)** | 1281 (311)|       | 1150 (401)  |       |
| **BW =>1000g**   | 27 (77)   |       | 15 (58)     |       |
| **BW <1000g**    | 8 (23)    |       | 11 (42)     |       |
| **Apgar Score at 5 min** | 8 (7-9) |       | 8 (7-9)**   |       |
| **Cord pH**      | 7.24 (0.08) |     | 7.24 (0.09)* |       |
| **SGA**          | 3 (9)     |       | 0 (0)       |       |
| **Any IVH**      | 4 (11)    |       | 5 (19)      |       |
| **IVH (>grade II)** | 0 (0) |       | 4 (9.7)     |       |
| **Ventriculomegaly** | 1 (3) |       | 0 (0)       |       |
| **Hydrocephalus**| 0 (0)     |       | 1 (4)       |       |
| **PVL**          | 0 (0)     |       | 6 (23)      |       |
| **Sepsis**       | 17 (49)   |       | 10 (39)     |       |
| **PDA**          | 13 (37)   |       | 9 (35)      |       |
| **NEC**          | 1 (3)     |       | 4 (15)      |       |
| **ROP**          | 0 (0)     |       | 2 (8)       |       |
| **Surgery**      | 6 (17)    |       | 3 (12)      |       |
| **BPD**          | 4 (11)    |       | 7 (27)      |       |
| **Postnatal steroids** | 3 (9) |       | 5 (19)      |       |
| **Intubation**   | 21 (60)   |       | 19 (73)     |       |
| **MV**           | 11 (31)   |       | 6 (23)      |       |
| **Weight at discharge** | 1969 (451) |   | 1969 (369)* |       |
| **CGA at discharge** | 36 (35-39) |   | 36 (35-39)** |       |
| **LOS**          | 37 (29-78)|       | 57 (36-95)** |       |

### TABLE 2: Neonatal characteristics and in-hospital morbidities.

NDD: Neurodevelopmental delay; GA: Gestational age; BW: Birth weight; SGA: Small gestational age; IVH: Intraventricular hemorrhage; PVL: Periventricular leukomalacia; PDA: Hemodynamically significant patent ductus arteriosus; NEC: Necrotizing enterocolitis > stage I; ROP: Retinopathy of prematurity more than stage II or plus disease; BPD: Bronchopulmonary dysplasia with the requirement of oxygen by 28 days of life; MV: Mechanical ventilation.

* Mean (SD)  ** Median (IQR)

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From a total score of 500, each domain accounts for 60 points. Out of 300, the lowest total score was 35, and the highest total score was 275. Children with a normal screen had a mean (SD) of 230 (31.2) compared to children with delay 137.3 (59), p-value <0.001. The delay in communication skills was detected in seven infants (11.5%), gross motor: seven infants (11.5%), fine motor: 12 infants (19.7%), problem-solving skills: twelve infants (19.7%), and personal-social skills: 14 infants (23%). In regards to the number of domains affected, in infants with delays, 15 (24.6%) had only one domain affected, while the rest 11 (18.%) had more...
than one domain affected. Of notice, all infants who had three or more domains affected corresponded to<br><br>1SD of the total score, while those who had all five domains affected corresponded to <2 SD. In univariate<br>analysis, only perinatal events were statistically significant of the maternal factors 4 (11%) versus 10 (39%)<br>in infants without NDD and with NDD, respectively. While in the analysis of neonatal factors, PVL was<br>statistically significantly higher 0 (0%) versus 6 (23%) (Table 2). Using Firth’s logistic regression model that<br>included the significant variable in the univariate analysis, PVL and perinatal events were independently<br>significant predictors for NDD, where PVL coefficient: 2.9, p-value: 0.05 and perinatal events coefficient:<br>1.34, p-value: 0.04, penalized log-likelihood: -33.2 and Wald ch2: 7.5, p-value: 0.02.

Discussion

This study provides information on the NDD rates of Saudi preterm, VLBW infants born in the KAMC-WR<br>using the ASQ3-A screening tool in a cross-sectional design, accounting for 42.6%. The prevalence of NDD<br>in preterm infants is well-documented, according to multiple studies [3,26,27].

Since ASQ3-A has a high negative predictive value, it would decrease the workload and the need for<br>extensive neurodevelopmental testing. In addition, ASQ3-A is a cost-effective screening tool as it selects<br>only smaller groups of high-risk children who require further assessment. Whereas a larger group of high-risk<br>children like moderate and late preterm-born children, children with congenital defects or any other<br>severe malformation who require neonatal surgery can be assisted by other essential available<br>neurodevelopmental testing [21]. Moreover, a retrospective study compared Bayley and ASQ3, and they<br>found no statistically significant difference in the performance of Bayley (81.1%) compared to ASQ3-A<br>(85.3%). Furthermore, ASQ3-A had a kappa score of (k – 1.00). As a result, ASQ3 scores seemed satisfactory<br>in conducting multidisciplinary team referral decisions and general proficiency in all situations [28]. As seen<br>in this study, all 61 preterm infants initially recruited had complete demographic data with all morbidities,<br>which is an excellent source to record the prevalence of common morbidities known to be associated with<br>less than 32 weeks or VLBW preterm infants.

A study in France using ASQ3 on 224 very preterm born children at the corrected age of 2 reported that 27%<br>of children had abnormal ASQ3 scores, whereas our study showed that 42.6% of the children had abnormal<br>ASQ3-A scores. The difference could be explained by the low number of samples collected or the chances of<br>high false-positive results [21]. A local retrospective study demonstrated that 14.2% of a total of 567 preterm<br>infants aged 24-36 months had abnormal cognitive function according to Bayley Infant<br>Neurodevelopmental Screener (BINS). Such screening tools are old and not widely used and are based on an<br>older version of Bayley-II [17,29].

Compared to a prospective study done by Agarwal PK et al. to evaluate VLBW and preterm at 9,12,18,24<br>months without congenital malformation by using ASQ3, The NDI in 18 and 24 months in the overall<br>domain was 45% which was comparable to our study, 42.6%. Moreover each domain was; communication<br>accounted for 12.5% vs. 11.5%, gross motor 15% and 11.5%, fine motor 13.8% vs. 19.7%, problem-solving<br>12.5% vs. 19.7%, personal-social 14.1% and 23%, respectively. Such comparable figures are reassuring. Of<br>note, they used 2SD below the mean as a cut-off for NDD; however, with sensitivity and specificity averaging<br>70%, and net present value (NPV) of 94% compared to Bayley–III, our approach of choosing 1SD is<br>considered more conservative [30]. Such an approach in a country still lacking standardized definitive<br>diagnostic psychomotor and cognitive testing is considered acceptable.

The study that was done in King Khalid University Hospital in Riyadh showed that the male gender is a<br>significant predictor of poor outcomes. On the other hand, our study showed that NDD was almost<br>comparable in males [29]. Such male predominance is known in the literature; however, our results could be<br>affected because of high loss to follow-up. A study of children at 24 months showed that abnormal<br>neurodevelopmental outcomes were more prominent in children with risk factors as pathologies of<br>neonates, pregnancy or delivery, and birth before 57 weeks of gestation. A strong correlation was found<br>between adverse neurodevelopmental outcomes and the combined effect of risk factors and neonatal<br>pathologies, particularly complicated neonatal sepsis and CNS malformation [31,32]. This was no different<br>from what we got as prenatal events and PVL were strong independent predictors for NDD.

In addition, a meta-analysis of 11 studies found that 18.6% of patients with isolated severe bilateral<br>ventriculomegaly have mild/moderate disability, and 59.6% have a severe disability [33]. In Portugal, a study<br>published in 2020 reported that severe neurodevelopmental deficit was diagnosed in 30.4% of premature<br>infants born with severe peri-intraventricular hemorrhage. These studies support our results that concluded<br>that CNS abnormalities, including IVH grade 3 and 4, ventriculomegaly, hydrocephalus, and periventricular<br>leukomalacia, are major risk factors for NDD in preterm infants [54].

In our study, other risk factors were comparable between the groups, especially GA and BW, hence predictors<br>were insignificant. However, a larger sample size could have more robust results that concur with the<br>literature.

This study screened for more detailed domains related to NDI. Most of the other similar studies focused on
either significant or fewer outcomes. In addition, ASQ3-A is a very reliable screening method to be used in limited settings. This study is the first study to use ASQ3-A to screen for NDI in Saudi Arabia; further validation studies using this tool in Arabic-speaking populations are required.

We collected data of 61 children, while a higher number of data is recommended to show more accurate results. Low infants' numbers could be explained by the delay caused by the COVID-19 pandemic. Also, this study is conducted in a single center, which might not reflect the actual prevalence in Saudi Arabia. We recommend a national multi-center study with multidomain factor assessment, including socioeconomic and antenatal variables [35,36]. The follow-up should be comprehensive, considering particular risks such as preterm infants have, like increased risk of respiratory diseases and health care utilization, especially during the COVID-19 pandemic [37-39]. National guidelines should be developed based on country-specific resources and parents' values [36].

Conclusions
This single-center study in Saudi Arabia screened preterm, VLBW infants based on ASQ3-A. Twenty infants (42.6%) had an abnormal NDD at a corrected age of 18-24 months. Perinatal events and PVL were independent predictors of NDD. We recommend that all preterm VLBW infants in Saudi Arabia be evaluated by a neurodevelopmental screening tool, ASQ3-A, especially in resource-limited settings to start early intervention. Also, more extensive multicenter studies are to be carried out with definitive diagnostic tools to have a national benchmark for the long-term NDI.

Additional Information
Disclosures
Human subjects: Consent was obtained or waived by all participants in this study. Institutional Review Board (IRB) at King Abdullah International Medical Research Center issued approval SP18/189/J. Animal subjects: All authors have confirmed that this study did not involve animal subjects or tissue. Conflicts of interest: In compliance with the ICMJE uniform disclosure form, all authors declare the following: Payment/services info: All authors have declared that no financial support was received from any organization for the submitted work. Financial relationships: All authors have declared that they have no financial relationships at present or within the previous three years with any organizations that might have an interest in the submitted work. Other relationships: All authors have declared that there are no other relationships or activities that could appear to have influenced the submitted work.

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