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Abstract

Background: Although inborn errors of metabolism (IEM) are rare individually, collectively IEM cause substantial morbidity and mortality and the diagnosis is challenging.

Aims: To analyse epidemiological and clinical data, final diagnosis and clinical outcomes of patients with a suspected diagnosis of IEM (small molecule disorders type) admitted to a paediatric intensive care unit (PICU).

Methods: We collected and analysed medical records data of all patients admitted to the PICU at Alexandria University Children’s Hospital from January 2010 to December 2014 with a suspected or confirmed diagnosis of small molecule disorders, including clinical presentations, laboratory results and clinical outcomes.

Results: A total of 34 patients had a suspected or confirmed diagnosis of small molecule disorders at PICU admission. Diagnosis was confirmed in 22.7% of suspected cases at admission and in 25% of suspected cases during PICU stay. Consanguineous marriage was found in 50% of cases with confirmed small molecule disorders.

Conclusions: A high index of suspicion is important for diagnosing and categorizing small molecule disorders in screening of high-risk individuals in developing countries.
Introduction

Inborn errors of metabolism (IEM) are a diverse heterogeneous group of inherited disorders that usually present in the paediatric population as varied clinical manifestations of defects in catabolism or anabolism of nutrients or energy-producing molecules. Although rare individually, collectively IEM cause substantial morbidity and mortality (1). They can be classified as disorders involving either large molecules (complexes or organelles) or small molecules. Organelle diseases are characterized by a gradual, often insidious, onset of symptoms, a relatively slowly progressive clinical course and specific clinical signs, which may be characteristic enough to make a diagnosis. Response to supportive therapy is generally only fair or poor. In contrast, IEM involving small molecules tend to be characterized by a rapid onset of symptoms and a clinical course featuring remissions and relapses. Physical findings are generally non-specific, which makes their diagnosis challenging at first presentation. However, small molecule disorders tend to respond well to aggressive supportive therapy (2).

Diagnosis of small molecule disorders is challenging due to the episodic nature of the metabolic illness, wide range of non-specific clinical symptoms also associated with common clinical conditions, general lack of experience among paediatric subspecialists and the need for expensive investigations (3).

No statistics on small molecule disorders of IEM are available in Egypt. The aim of this retrospective study was to analyse the epidemiological and clinical data, final diagnoses and clinical outcomes of patients admitted to the paediatric intensive care unit (PICU) at Alexandria University Children’s Hospital with a suspected or confirmed diagnosis of small molecule
disorders.

**Methods**

In this 5-year retrospective study, we collected medical records of all patients admitted to the PICU at El-Shatby Children’s Hospital, a tertiary care teaching hospital affiliated to Alexandria University (serving 4 governorates comprising 14 million people), from 1 January 2010 to 31 December 2014 (n = 1417). Ages ranged from 1 month to 6 years. The policy of our hospital is that all cases of small molecule disorders presented at the emergency room are admitted to PICU. Inclusion criteria were: confirmed diagnosis of small molecule disorders and suspected diagnosis of small molecule disorders at the time of PICU admission or during PICU stay, with no subsequent confirmation of diagnosis (according to PICU protocol). The exclusion criterion was previous diagnosis of small molecule disorders in patients admitted to the PICU during the study period. Out of 1417 PICU admissions, 34 patients (2.4%) had a suspected or confirmed small molecule disorder.

To identify patients with suspected small molecule disorders and determine their management, according to our PICU policy, paediatric intensivists were required to enter specific information on patient symptoms and signs as well as laboratory investigations undertaken into a computer-based protocol. The symptoms and signs of clinical suspicion of small molecule disorders comprise a history of consanguinity, siblings with a history of similar illness or previous unexplained death, respiratory distress, apnoea, unexplained neurological symptoms, unexplained acute liver failure, feeding difficulties, failure to thrive, hepatomegaly and splenomegaly. Initial simple laboratory investigations performed immediately on admission to the PICU included random blood glucose, ketone bodies, arterial blood gases, anion gap, blood lactate, pyruvate and lactate/pyruvate ratio, blood ammonia and uric acid. Results from the laboratory investigations helped to classify small molecule disorders into 5 categories: aminoacidopathies, organic acidopathies, fatty acid oxidation defects, primary lactic acidosis and urea cycle defects.

From the medical records, we gathered demographic data, clinical data and laboratory findings suggesting small molecule disorders. Other data that were recorded by the residents were the Paediatric Index of Mortality 2 (PIM2) at admission, a scoring system for rating the severity of medical illness for children encompassing 10 physiological variables collected from the time of first physical contact between the patient and the PICU team up until 1 hour after physical PICU admission (4,5), and paediatric logistic organ dysfunction (PELOD) score on day 2 in the PICU, a scoring system covering of physical and laboratory variables representing 6 organ systems: nervous, cardiovascular, renal, respiratory, haematologic and hepatic systems. Each variable is assigned points (0, 1, 10, or 20) based on the level of severity, each organ dysfunction receives points for the variable associated with the highest number of points (6). The final diagnosis is documented as either a case of small molecule disorder (i.e. confirmed) or a non-small
molecule disorder (i.e. not confirmed or proven to be an alternative diagnosis) and patient clinical outcome (i.e. PICU discharge, death or death within ≤ 24 h of PICU admission).

All data were analysed using SPSS, version 20.0 (7). The Kolmogorov–Smirnov test of normality revealed significance in the distribution of some variables, so the non-parametric statistics were adopted. Qualitative data are presented as numbers and percentages, and quantitative data as minimum, maximum, median and interquartile range (IQR). Comparison between different groups of category variables was made using the chi-squared test. When > 20% of the cells had an expected count < 5, correction for chi-squared was made using Fisher’s exact test or the Monte Carlo correction.

Multivariate logistic regression analysis was done for prediction of mortality with PIM2 and PELOD as independent predictors. Area under the ROC Youden index (the vertical distance between the 45 degree line and the point on the ROC curve) was performed to determine the best cut-off value for the variable tested. Pair-wise comparison of areas under the ROC was carried out using MedCalc, version 14, for PIM2 and PELOD for discrimination for death (8,9).

The calibration was assessed by directly comparing the observed and customized predicted mortality. We employed the Hosmer–Lemeshow goodness-of-fit test, where a P-value > 0.1 indicates acceptable calibration (10).

Results

Out of 1417 PICU admissions from January 2010 to December 2014, 34 patients (2.4%) had a suspected or confirmed small molecule disorder. The age of onset ranged from 1.17 to 72.00 [median (IQR) 9.00 (2.88–24.00)] months. The male to female ratio was 1:1.27 (15:19 cases). Patient weight ranged from 1.55 to 20 [median (IQR) 7.00 (4.78–10.25)] kg. Patient length of stay in PICU ranged from 1 to 26 [median (IQR) 3.50 (2.00–10.00)] days. PIM2 ranged from 20.00 to 90.80 [median (IQR) 44.95 (27.60–85.00)].The mortality rate in the study group (18/34, 52.9%) was more than twice the general mortality in the PICU at Alexandria University Children’s Hospital (24.1%). There were 7 deaths within 24 h, and the PELOD score on day 2 for the remaining 27 cases ranged from 4 to 31 [median (IQR) 10.00 (5.00–20.00)] (Table 1).

Eight of the 34 cases (23.5%) were confirmed as small molecule disorders, whether suspected on admission or not suspected on admission and subsequently confirmed during PICU stay. The diagnosis of small molecule disorders was confirmed in 22.7% of cases (5/22) suspected on PICU admission, and in 25.0% of cases (3/12) not suspected on PICU admission (Table 2).
Of the 8 confirmed cases, 3 were diagnosed as organic acidaemia, 2 as urea cycle defects, 2 as aminoacidopathies and 1 as primary lactic acidosis. The best clinical outcome was obtained with cases of urea cycle defects and organic acidaemia where all patients, except 1 who died, were discharged.

A significantly higher rate of consanguinity was evident among cases confirmed as small molecule disorders compared to unconfirmed cases [4 cases (50.0%) vs 3 (11.5%) respectively, \( P = 0.037 \)]. No statistically significant difference were found in terms of sex, age on presentation to PICU, history of sibling death in infancy (unknown cause or stillbirth) and place of residence (Table 3).

There was a significantly higher rate of jaundice in the confirmed compared with the non-confirmed small molecule disorders groups [3 cases (37.0%) vs 1 case (3.8%) respectively, \( P = 0.033 \)] (Table 4). In addition, from the laboratory results obtained from initial investigations on PICU admission, there was a significantly higher rate of hyperammonaemia and hyperuricaemia in the confirmed compared with the non-confirmed cases of small molecule disorders group [5 cases (62.5%) vs 3 cases (11.5%), \( P = 0.009 \), respectively; 7 cases (87.5%) vs 8 cases 30.8%, \( P = 0.011 \), respectively] (Table 5). No other clinical presentations or laboratory results showed any statistically significant difference between the groups.

Using the predictive model of mortality, examining the PIM2 and PELOD score as independent covariates, the classification accuracy of the model (block 0, i.e. before implementation of the statistical model) was 59.25%, with variations in the PELOD and PIM2 scores accounting for 50.1% of the variation of occurrence of death in the study sample (–2 Log likelihood = 23.975, Nagelkerke R2 = 0.501). The regression model was well calibrated. The Hosmer–Lemeshow \( \chi^2 \) value was 4.954 (\( P = 0.666 \)). The overall model was statistically significant (\( \chi^2 = 12.524, P = 0.002 \)) (independent covariates had predictive capacity). The predictive capacity of the model increased from 59.2% (of the basic “null” model) to 81.5%. Only the PIM2 score was a statistically significant predictor of mortality [odds ratio (OR) = 1.050, 95% CI: 1.008–1.094] (\( P = 0.019 \)). The PELOD score was not a statistically significant predictor of mortality (OR = 1.100, 95% CI: 0.976–1.240) (\( P = 0.118 \)) (Table 6).

We found that PIM2 was a statistically significant discriminator of death: area under the ROC curve = 0.849 (95% CI: 0.685–0.948) (\( Z = 5.411, P < 0.0001 \)). The diagnostic criterion using the Youden index is the level of > 71.6, with sensitivity 64.71% (95% CI: 38.3–85.8), specificity 94.12% (95% CI: 71.3–99.9), positive predictive value 91.7% and negative predictive value
72.7%. The PELOD score was a statistically significant discriminator of death: area under the ROC curve = 0.759 (95% CI: 0.556–0.901) (Z = 2.600, P = 0.0093). The diagnostic criterion using the Youden index is the level of > 10.0, with sensitivity 72.73% (95% CI: 39.0–94.0), specificity 68.75% (95% CI: 41.3–89.0%), positive predictive value 61.5% and negative predictive value 78.6%. Pair-wise comparison for the 2 ROC curves showed no statistically significant difference (Figure 1).

Discussion

Over 500 known IEMs, over 100 involving neonates, have been described (11). The frequencies for each individual IEM vary; although most are very rare, collectively they are common (12). In Egypt, 1 in every 32 individuals is reported to harbour a gene for an IEM (13). An extended metabolic screen carried out in 2004 on 231 suspected paediatric cases of IEM (44 neonates and 187 children) in Egypt revealed that abnormal results were detected in 8.56% (14). The majority of IEMs from various countries have been reported to be small molecule disorders. A study in Thailand showed that 74.3% of diagnosed IEM cases were small molecule disorders (15), and a 25-year retrospective study in Saudi Arabia described a cumulative incidence of 150 IEM cases per 100 000 live births (0.15%), of which 54% were small molecule disorders (16). The incidence of IEMs is reportedly lower in Western countries, e.g. 1 in 2500 live births (0.04%) in British Columbia, Canada (17) and 1 in 2555 live births (0.04%) in Italy (18). The difference may be related to consanguineous marriage and intermarriage. In Egypt, the rate of consanguineous marriage is very high (35.3%), especially among first cousins (86%), and in rural areas, Upper Egypt and Cairo (19).

In our 5-year study, the incidence of suspected or confirmed cases of small molecule disorders was 2.4% (34/1417) of all PICU admissions, with only 8 confirmed cases of small molecule disorders of the 34 cases (0.56% of all 1417 PICU admissions). This low incidence could be explained by a high pre-admission mortality rate of patients with small molecule disorders. This is comparable with the results from a 5-year study of PICU admissions in a French teaching hospital (incidence 2.2% of all PICU admissions) (20) and a 1-year PICU study in India (incidence 2.6% of all PICU admissions) of suspected or diagnosed IEM, including small molecule disorders (21).

In a study from Brazil, only 6.7% (4/59) of suspected cases admitted to the PICU were diagnosed as IEM (22). In contrast, in a 3-year study in Pakistan 26% of suspected cases were diagnosed as IEM, including small molecule disorders (23). In our study, 1.5% (22/1417) of all PICU admissions were suspected cases of IEM, and only 22.7% (5/22) of these were confirmed to be small molecule disorders.
In developed countries, small molecule disorders are typically diagnosed using high-performance liquid chromatography (HPLC) and tandem mass spectrophotometry (3). Gas chromatography/mass spectroscopy is an advanced technique for diagnosing and confirming metabolic disorders and also for mass neonatal screening (18). Enzyme assays with leukocytes or erythrocytes are used to confirm a specific enzyme deficiency (3). These diagnostic techniques are expensive and beyond the financial capacity of many developing countries. Therefore, in developing countries, diagnosis of IEM relies on simple clinical and laboratory tests for the selective screening of high-risk individuals suspected of IEM, in order to identify the small molecule disorders so that early empirical supportive treatment can be initiated. This approach is often supported by other approaches to confirm the final diagnosis. For example, in Thailand, clinicians work in collaboration with the Chulabhorn Research Institute in Bangkok for the use of high-performance liquid chromatography (15), and in Pakistan (23) and Saudi Arabia (16) samples are sent to specialized laboratories in Japan and the United States, respectively, to confirm the final diagnosis. Thus, screening for IEM without primary selection is a costly process for developing countries with limited resources. Patients with small molecule disorders are usually diagnosed following PICU admission and are considered acute emergencies amenable to lifesaving therapy (20). However, their prognosis is very poor due to the delay in diagnosis and management (15). Therefore, small molecule disorders warrant heightened attention, as early diagnosis is essential for early treatment and improved clinical outcome.

A French study reported a mortality rate of 28.6% (20/70) of confirmed IEM cases among PICU admissions, which was twice that observed for all PICU admissions and 4 times that observed in European PICUs (20). A retrospective study in Italy reported a PICU mortality rate of 25.3% of confirmed IEM cases (18). A study from India found a mortality rate of 36% among PICU admissions of confirmed IEM cases (21). Consistent with these findings from previous studies, in our study, the mortality rate among the 34 patients we studied was 52.9%, whereas the mortality rate among confirmed cases of small molecule disorders was 37.5% (3/8), which was higher than our total PICU mortality rate of 24.1%. We should, however, take into consideration the very small number in the sample. The high mortality rate may be due to delayed diagnosis, encephalopathy, respiratory distress, heart failure or dehydration.

The rate of various clinical presentations did not show any statistically significant differences between confirmed and non-confirmed small molecule disorders groups, except for the presence of jaundice, which was significantly higher in the confirmed groups. This may be explained by the presence of severe illness in patients with small molecule disorders, who usually present with multiple organ dysfunction syndromes, including pre-hepatic failure, which manifests as jaundice. Our finding is in contrast to those of a prior report in which about half of the IEM patients, including those with small molecule disorders, had acute encephalopathy as the most common clinical presentation (15). A 3-year study in Pakistan showed that respiratory distress and developmental delay were the most common clinical presentations in cases
diagnosed with IEM, including small molecule disorders (23). This difference could be explained by the variability of symptoms and severity and the non-specificity of small molecule disorders.

Comparison of laboratory results between confirmed small molecule disorders and not confirmed cases revealed a significantly higher rate of hyperammonaemia and hyperuricemia in the confirmed group. This may be because of urea cycle defects and organic acidaemia, which represented 62.5% (5/8) of cases, usually present with hyperammonaemia. This finding is in agreement with prior findings of a significantly higher rate of hyperammonaemia in the IEM group, including small molecule disorders, compared to the non-IEM group, although, contrary to our findings, metabolic acidosis and ketosis were also found to be frequent (15).

We evaluated the performance of the PELOD score compared with PIM2 for predicting mortality in small molecule disorders in PICU; most research uses these scores generally for survival in PICU (24,25), however in our study, only the PIM2 score was a statistically significant predictor of mortality. Similarly, a 2013 study in India reported the under-prediction of mortality by PELOD-2 compared to PIM2 (24); it is possible that the predictive mortality models could be population sensitive, so validation studies are necessary before application in other settings and populations.

A limitation of our study is the small sample size; this ensues from the rarity of small molecule disorders.

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References

4. Slater A, Shann F, Pearson G; Paediatric Index of Mortality (PIM) Study Group. PIM2: a


