Hypernatraemic dehydration in infants with acute gastroenteritis at King Edward VIII Hospital, Durban, South Africa

T Hariram, MB ChB, DCH (SA), FC Paed (SA); K L Naidoo, MB ChB, DCH (SA), FC Paed; S Ramji, MB ChB

Department of Paediatrics and Child Health, University of KwaZulu-Natal, Durban, South Africa

Corresponding author: T Hariram (terishiapillay@gmail.com)

Background. Acute gastroenteritis (AGE) is a leading cause of infant mortality, with hypernatraemic dehydration contributing to increased morbidity and mortality.

Objectives. To determine the prevalence of hypernatraemia secondary to AGE in admitted infants in Durban, South Africa. To describe the feeding choices, nutritional status and outcomes of these patients. To determine the association between admission sodium (Na) level, the rate of Na correction and clinical outcomes.

Methods. A retrospective chart review was conducted on cases of hypernatraemic dehydration admitted in 2014 to a South African hospital. Serum Na levels were corroborated with National Health Laboratory Services results. Descriptive and analytical statistics were done using Statistical Package for Social Sciences version 22.

Results. A 12.3% prevalence of hypernatraemia (n=41/334) was found. The majority of infants were formula-fed (76%) with a 21% incidence of malnutrition and 66% HIV exposure rate in this cohort. A high rate of neurological abnormalities (63%), and a 4.9% case fatality rate was found. Shock on admission was present in 92% of patients who developed severe neurological complications. The mean admission Na was higher in those with severe neurological complications (164.2 ± 158.4 mmol/L, p=0.08). The mean rate of Na change was not faster in those with severe neurological morbidity (0.61 ± 0.91 mmol/L/hr; p=0.1).

Conclusion. Hypernatraemic dehydration remains a significant problem in South Africa. High rates of formula feeding may be a contributory factor and the correlation with HIV infection needs investigation. Poor neurological outcomes were noted particularly in those presenting with hypernatraemia and shock. Although the mean admission Na level was higher in patients with severe neurological complications, this was not statistically significant in this sample. This study supports the hypotheses that neurological complications in diarrhoea-related hypernatraemia are largely associated with the severity of the dehydration that occurs prior to presentation rather than following rehydration.

Approximately 76% of deaths in children under 5 years of age occur within the first 12 months of life, with diarrhoeal disease ranked as one of the leading causes of infant mortality in South Africa (SA). Hypernatraemic dehydration as a complication of diarrhoeal disease, contributes to the morbidity and mortality of affected patients, with a prevalence of 6.4% to 13.7%, respectively, reported in developing countries.

Diarrhoea-related hypernatraemic dehydration occurs as a result of a negative water balance, with water loss exceeding sodium (Na) loss. Salt poisoning – commonly due to incorrect constitution of homemade oral rehydration therapy (ORT) and over-concentrated formula feeds – is also an important factor in the aetiology of hypernatraemia.

Acute onset hypernatraemia increases serum osmolality, causing fluid to shift from the intracelluar to the extracellular space, leading to cerebral cell dehydration and shrinkage. This can perpetuate structural changes resulting in intracranial haemorrhages. As a means of adaptation to gradual onset hypernatraemia, the brain increases its intracellular content of idogenic osmoles, thus regaining its cell volume. In this event, rapid correction of the Na with hypertonic intravenous fluids can lead to cerebral oedema.

Thus, neurological abnormalities are common, ranging from irritability and tone disturbances to seizures and coma. Mortality rates vary between 3% and 13%, and hypernatraemic dehydration has been implicated as a predictor of death in children younger than 5 years old admitted with acute gastroenteritis (AGE).

Despite high morbidity and mortality rates, the management of hypernatraemic dehydration is still not standardised in national guidelines. Institutions and teaching hospitals have developed individually tailored protocols developed by clinicians over many years. In KwaZulu-Natal (KZN), where AGE rates remain high, such a protocol has been in place for the past decade in Durban hospitals, including King Edward VIII Hospital (KEH), which is the major teaching hospital in the province.

There are no published randomised controlled trials that clearly document the ideal choice of intravenous fluid, or the optimal rate of Na correction that can be undertaken without developing cerebral oedema, and there are conflicting results as to the correlation between the rate of fall of Na and clinical outcomes.

To reduce its impact on the neurological morbidity of infants in SA, there is a need to further understand this condition and the factors that may contribute to such adverse outcomes. The aim of this study was to describe the extent of hypernatraemic dehydration in an HIV-endemic province in SA and to clarify if the major contributor to these outcomes is the severity of the hypernatraemia itself, or complications arising from rapid fluid correction.

The objectives of the study were: (i) to determine the prevalence of hypernatraemia (Na ≥150 mmol/L) in all infants admitted with AGE at KEH from January until December 2014; (ii) to describe the feeding patterns, nutritional status and clinical outcomes of these infants; and (iii) to determine the association between admission Na level, rate of Na correction and clinical outcomes.

Methods
This study was a retrospective chart review conducted at KEH, a regional hospital within the Durban Metropolitan area in KZN. Purposive sampling was used to identify cases of hypernatraemic dehydration in infants younger than 12 months old. Utilising a physician-verified institution database, all cases of AGE in infants admitted between January and December 2014 were identified. The admission serum Na levels of these cases were traced via the National Health Laboratory Service (NHLS) Trakcare Web Results. Those with an admission Na level ≥150 mmol/L were included in the study. This two-step process aimed to prevent any omission of cases of hypernatraemia that might have not been documented in the admission database. The inpatient charts of identified cases were retrieved from the KEH medical records department for detailed clinical review. Demographic characteristics of each patient, nutritional status, feeding choice, as well as clinical data, including documented neurological symptoms, signs and outcomes (length of stay or mortality) were captured. Nutritional status was based on the weight-for-height Z-score, the presence of oedema and visible severe wasting. A 10% adjustment in admission weight was made to correct for the degree of dehydration. The serum Na on admission and subsequent Na levels during the course of inpatient management, together with their sampling times, was used to calculate the average rate of Na reduction until correction (<150 mmol/L) within the first 24 hours of admission.

To improve accuracy and minimise inter-observer variability, the clinical examination findings of the attending specialist paediatrician served as the primary source of data related to clinical symptoms and signs. For the purposes of this study, we recorded and classified the presence of irritability and/or abnormal muscle tone during the period of admission as a mild neurological abnormality. The occurrence of seizures and/or encephalopathy classified the patient as having a severe neurological abnormality.

It is plausible for a single patient to have more than one documented neurological abnormality during their admission; therefore, each patient was classified based on the most severe complication they developed, with encephalopathy considered most severe on the spectrum. This eliminated any duplication of results during the process of data analysis.

Data were captured on an Excel spreadsheet and subsequently analysed using the Statistical Package for Social Sciences version 22 (IBM Corp., USA). Descriptive statistics, including frequencies, percentages, means, medians and standard deviations (SD) were used to summarise results. Independent Samples t-test or the Mann-Whitney test was used to test for association between admission serum Na level, the rate of Na correction and outcomes. The level of significance was set at 0.05.

Ethics
Ethical approval was obtained from the Biomedical Research Ethics Committee of the University of KwaZulu-Natal (ref. no. BE266/15).

Table 1. HIV status of infants with AGE admitted to KEH in 2014

<table>
<thead>
<tr>
<th>HIV status</th>
<th>AGE with hypernatraemia (N=41), n (%)</th>
<th>AGE without hypernatraemia (N=293), n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unexposed</td>
<td>14 (34)</td>
<td>151 (52)</td>
</tr>
<tr>
<td>Exposed, HIV-uninfected</td>
<td>20 (49)</td>
<td>91 (31)</td>
</tr>
<tr>
<td>Exposed, HIV-unknown</td>
<td>4 (10)</td>
<td>33 (11)</td>
</tr>
<tr>
<td>Exposed, HIV-positive</td>
<td>3 (7)</td>
<td>16 (5)</td>
</tr>
<tr>
<td>Unknown</td>
<td>0</td>
<td>2 (1)</td>
</tr>
</tbody>
</table>

*The HIV status of the three patients with missing files was obtained from the hospital admission database.

Results
There were 1 422 infants admitted to KEH in 2014, of whom 334 (23.5%) had an admission diagnosis of AGE. The prevalence of hypernatraemic dehydration secondary to AGE in infants was found to be 12.3% (n=41/334 cases). Three of the 41 patient files could not be traced, leaving 38 patients eligible for chart review. The sample population consisted of 84% infants under the age of 6 months (n=32/38) and 16% between 6 to 12 months of age (n=6/38); 55% were male (n=21/38) and 45% female (n=17/38). ORT had been used in 55% of patients (n=21) prior to presentation, 8% of patients (n=3) had not used ORT and 37% of the patient files (n=14) did not document whether ORT had been used or not. An initial clinical assessment of shock was present in 45% of the patients (n=17), of whom 65% also experienced a severe neurological complication. The HIV status of all infants with AGE is described in Table 1.

The hypernatraemic cohort was noted to have a 66% HIV exposure rate in comparison with 47% in those infants admitted for AGE without hypernatraemia, and this was a statistically significant difference (p=0.03). Table 2 lists the nutritional status and documented feeding choices and practices of the sampled cohort.

With regards to nutrition and feeding, 21% of hypernatraemic patients were classified as either moderate or severe acute malnutrition. The nutritional classification of patients admitted for AGE without hypernatraemia was recorded as 64% normal, 17% underweight and 19% severely underweight for age.

Seventy-six percent of the hypernatraemic cohort were exclusively formula fed with only 16% exclusively breastfed. Of those receiving...
formula feeds, 18% of patient caregivers were questioned about their formula dilution practices. Table 3 describes the clinical outcomes of the sampled cohort.

The majority (90%) of patients were discharged from KEH and 2 patients (4.9%) were transferred to an intensive care unit. The mortality rate of infants with hypernatraemic dehydration at KEH was 4.9%. Both patients who demised had an initial clinical assessment of shock, as well as co-morbid conditions, one of which was severe sepsis with disseminated intravascular coagulation and the other was a chromosomal anomaly.

Neurological abnormalities were documented in 63% of reviewed infants, with 32% developing seizures and/or encephalopathy. Of those who developed seizures, 55% were preceded by irritability and/or abnormal muscle tone. Of those who developed encephalopathy, 88% had preceding seizures. An initial clinical assessment of shock was present in 92% of patients who developed severe neurological complications. The association between admission serum Na level and neurological outcome is shown in Table 4.

The mean (SD) admission Na level for the sample population was 160.2 (9.49) mmol/L (range 150 to 181). A higher admission serum Na level was seen in those who developed severe neurological complications. The association between admission serum Na level and neurological outcome is shown in Table 4.

The mean (SD) admission Na level for the sample population was 160.2 (9.49) mmol/L (range 150 to 181). A higher admission serum Na level was seen in those who developed severe neurological abnormalities; however, this association was not statistically significant.

There was no statistically significant difference between the median admission Na of those who died (n=2) and those who survived (n=36) (160.5 v. 157.5 mmol/L, p=1 using Mann-Whitney test).

The association between rate of Na correction and neurological outcome is shown in Table 5. The rate of Na correction ranged from −0.1 to 2.66 mmol/L/hr, with a mean (SD) of 0.81 (0.53) mmol/L/hr (n=37, as one patient had only an admission serum Na record with no subsequent measurements).

Severe neurological complications were not associated with a faster mean rate of Na correction. The sample population was also grouped into three categories according to rate of Na correction: <1 mmol/L/hr; 1 - 2 mmol/L/hr; and >2 mmol/L/hr. The categories were compared with outcomes. This showed that 70% of all patients had a rate of Na correction <1 mmol/L/hr. Of those with a rate of Na correction ≥1 mmol/L/hr, 18% developed a severe neurological complication. There was no statistically significant difference in the development of neurological complications between the groups (p=0.489 using Fisher’s exact test). Both patients who demised had severe neurological morbidity, and both had a rate of Na correction <1 mmol/L/hr.

**Discussion**

This study highlights the persistent burden of hypernatraemic dehydration secondary to AGE in SA. While the developed world now reports a decline in cases of hypernatraemic dehydration,[11] the prevalence rate of 12.3% in our study remains similar to that of earlier studies conducted in developing countries.[12]

The incidence of hypernatraemia is said to increase with decreasing age,[12,13] as was the case in this study, with 84% of infants being under the age of 6 months. The decreased thirst response, as well as the inability to access adequate amounts of free water despite ongoing losses, contributes to this particular age group being identified as high-risk.[14]

The hypernatraemic cohort in this study had a 66% HIV exposure prevalence, compared with 47% in those infants admitted for AGE without hypernatraemia, and this was a statistically significant difference. Both these results were also noted to be higher than the provincial HIV exposure prevalence of 40.1% in KZN.[12] It has been shown that HIV exposed uninfected infants have an increased risk of infectious morbidity, particularly when born to mothers with advanced disease,[12,13] and this could explain the increased prevalence among these infants with diarrhoeal disease. This study, however, describes a higher HIV exposure prevalence in relation to hypernatraemic dehydration, warranting further studies in this regard.

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**Table 3. Clinical outcomes**

<table>
<thead>
<tr>
<th>Outcomes (N=41)*</th>
<th>n (%)</th>
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<tbody>
<tr>
<td>Discharged</td>
<td>77 (18.7)</td>
</tr>
<tr>
<td>≤7 days</td>
<td>24 (58.5)</td>
</tr>
<tr>
<td>&gt;7 days</td>
<td>13 (31.7)</td>
</tr>
<tr>
<td>Transferred to PICU</td>
<td>2 (4.9)</td>
</tr>
<tr>
<td>Demised</td>
<td>2 (4.9)</td>
</tr>
<tr>
<td>Spectrum of neurological abnormalities (N=38)</td>
<td></td>
</tr>
<tr>
<td>Nil</td>
<td>14 (37)</td>
</tr>
<tr>
<td>Irritability and/or abnormal muscle tone only</td>
<td>12 (31.5)</td>
</tr>
<tr>
<td>Seizures</td>
<td>4 (10.5)</td>
</tr>
<tr>
<td>Seizures progressing to encephalopathy</td>
<td>7 (18)</td>
</tr>
<tr>
<td>Encephalopathy</td>
<td>1 (3)</td>
</tr>
<tr>
<td>PICU = paediatric intensive care unit.</td>
<td></td>
</tr>
<tr>
<td>*The outcomes of the three patients with missing files was obtained from the hospital admission database.</td>
<td></td>
</tr>
</tbody>
</table>

**Table 4. Association between admission serum Na level (mmol/L) and severe neurological outcomes***

<table>
<thead>
<tr>
<th>Clinical outcome (N=38)</th>
<th>n (%)</th>
<th>Admission serum Na (mmol/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>No severe neurological abnormality noted</td>
<td>26 (68.4)</td>
<td>158.4</td>
</tr>
<tr>
<td>Seizures and/or encephalopathy present</td>
<td>12 (31.6)</td>
<td>164.2</td>
</tr>
<tr>
<td>SD = standard deviation, CI = confidence interval.</td>
<td></td>
<td></td>
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<tr>
<td>*p-value (two sample t-test) = 0.08.</td>
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**Table 5. Association between rate of Na correction (mmol/L/hr) and severe neurological outcomes**

<table>
<thead>
<tr>
<th>Clinical outcome (N=37)</th>
<th>n (%)</th>
<th>Rate of Na correction (mmol/L/hr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>No severe neurological abnormality noted</td>
<td>25 (67.6)</td>
<td>0.91</td>
</tr>
<tr>
<td>Seizures and/or encephalopathy present</td>
<td>12 (32.4)</td>
<td>0.61</td>
</tr>
<tr>
<td>SD = standard deviation, CI = confidence interval.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>p-value (two sample t-test) = 0.1.</td>
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</table>
Due to 31% of mothers using formula feeds indicating that their HIV status was the reason for their feeding choice, and as formula feeds are a recognised risk factor for the development of hypernatraemic dehydration, we considered the effect of these feeding choices on HIV exposure prevalence and hypernatraemia; however, when we compared the HIV-exposed with the HIV-unexposed patients within the hypernatraemic cohort, we found an equally high number of patients who were exclusively formula-fed (76% v. 77%).

The use of formula feeds in developing countries is increasing, with only 8.3% of SA infants under the age of 6 months reported to be exclusively breastfed. KEH advocates the current national infant feeding strategy, which recommends exclusive breastfeeding for all infants until 6 months of age, including those born to HIV-positive mothers. In keeping with this, the distribution of free formula feeds to HIV-positive mothers was discontinued in 2011. Despite this, the low rate of exclusive breastfeeding was also apparent in our study at KEH, with 76% of infants being formula-fed and 8% mixed-fed with both formula and breastmilk.

Although the majority of patients were formula-fed, we noted that 82% of caregivers were not questioned about formula dilution practices. Over-concentration of formula feeds leads to a high osmotic load resulting in hypernatraemia, so obtaining this information from caregivers is vital to identify such errors in feed dilution.

ORT serves as the first line management for the treatment of dehydration at the primary healthcare level; however, Dippenaar et al. showed that 14% of caregivers added too much salt to the homemade sugar-salt solution, which is a known risk factor for the development of hypernatraemia. We noted that 37% of caregivers were not asked about the use of ORT prior to admission. This should be reinforced as an important component of the history-taking process for any patient admitted with AGE, to identify errors in ORT constitution and prevent recurrences.

Approximately 21% of our study population was found to have moderate and severe acute malnutrition. This was an unexpected finding, as hypernatraemic dehydration is classically described in the well-nourished infant and its incidence is said to be inversely related to the degree of malnutrition. A study by Chisti et al. in Bangladesh reported that 28% of their hypernatraemic patients had severe wasting. This finding could be attributed to the higher rates of malnutrition in developing countries overall, as more than one-third of the infants admitted to KEH for AGE without hypernatraemia were also found to be underweight or severely underweight for age. Approximately two-thirds of patients in this study experienced abnormal neurological manifestations, of which 50% were seizures and/or encephalopathy. Robertson et al. also reported a high incidence of seizures within their sample of hypernatraemic patients in Cape Town.

The mortality rate of 4.9% for hypernatraemic dehydration was similar to that of other studies. However, it was higher in comparison to the age-matched mortality rate of 1% for infants admitted to KEH for AGE without hypernatraemia. This highlights the increased risk of mortality in patients with hypernatraemic dehydration, but is it a higher admission Na level or a rapid rate of Na correction that contributes to this increased risk?

Admission Na level and rate of Na correction

The management of hypernatraemic dehydration is based on providing adequate free water to correct the serum Na level, however, the rate at which it is corrected remains a point of contention. A study conducted in Cape Town found that rate of Na correction was not significantly associated with adverse outcomes. In contrast, a Chinese study concluded that a faster rate of Na correction was a significant risk factor for the development of cerebral oedema.

In the face of these conflicting study results, no definitively safe rate of correction has yet been advocated. Moritz and Ayus recommend that, unless symptoms of hypernatraemic encephalopathy are already present, a rate of correction not exceeding 1 mmol/L/hr is reasonable. Another recommendation by Duggal et al. is to factor in the rate of development of the hypernatraemia. A gradual onset requires slower correction at 0.5 mmol/L/hr, while an acute onset can tolerate faster correction, with a rate of 1 mmol/L/hr deemed acceptable. The experience at KEH has been to aim for a rate of Na correction between 0.5 and 1 mmol/L/hr using a standardised hospital protocol that guides the fluid management of hypernatraemic dehydration. This protocol was developed by senior clinicians using their collective experience in the management of such cases over a number of years.

Standard ORT is used in cases of mild hypernatraemic dehydration. Where intravenous rehydration is required, fluids containing a Na concentration of approximately two-thirds that of the patient’s Na level is used, at an initial rate of 10 mL/kg/hr. This accounts for maintenance requirements, deficit and ongoing losses, and is continued until the level of hydration improves and the patient is able to tolerate adequate amounts of ORT.

In this study, we found that 70% of hypernatraemic patients had a rate of Na correction <1 mmol/L/hr, in keeping with recommendations in the literature.

Our study showed no statistically significant correlation between a faster rate of Na correction within the first 24 hours and adverse neurological outcomes and mortality. Of those with a rate of Na correction ≥1 mmol/L/hr, only 18% developed a severe neurological complication.

In the light of these findings, it appears that the hypernatremia itself would be the more likely cause for adverse outcomes in these patients. This study showed that the mean admission Na level was higher in those with severe neurological morbidity and mortality; however, this association did not reach statistical significance.

We postulate that the development of neurological complications could be related to an interaction of factors prior to presentation, including the severity of the hypernatremia together with the clinical presence of shock. In this study, 92% of patients with severe neurological complications were in shock on presentation, highlighting it as an important factor in those with adverse outcomes. It is known that the degree of dehydration is often underestimated in hypernatraemic patients due to the relatively well-preserved intravascular volume. Thus, the signs of severe dehydration and shock occur late, and may initially be missed by caregivers. This leads to inadequate provision of ORT and delayed presentation, while intracerebral cerebral dehydration prevails. Hypernatremia and increasing plasma arginine vasopressin concentrations have also been shown to affect haemostatic function, thus promoting a hypercoagulable state, although the mechanism of this effect remains unclear. This association with an increased risk of thrombosis, together with the haemo-concentrated state induced by hypovolaemic shock, are risk factors for the development of cerebral sinus venous thrombosis, which can present with seizures and neurological abnormalities. The use of neuroimaging would ideally differentiate between intracranial haemorrhages, thrombosis and cerebral oedema in patients who experience severe neurological complications during admission; however, this is not always feasible in resource-limited settings.

Study limitations

As this was a retrospective study, data were obtained from information documented in patient files, which were often poorly recorded with regard to ORT use and formula dilution.

There was a small sample size. There was a lack of definite time-based data for all neurological signs. There was no case control group in this study with which to make clear comparisons with the hypernatraemic group.
Conclusion and recommendations
This study confirms that hypernatraemic dehydration related to AGE remains a significant challenge in South Africa, with a prevalence of 12.3% in a high HIV-exposed infant cohort. The predominant use of formula feeds, with low breastfeeding rates noted in these patients, may be a contributing factor. We also reaffirm the high rate of neurological morbidity and increased risk of mortality in hypernatraemic patients in this study. Mean admission Na was higher in cases with severe neurological complications and mortality, although this finding did not reach statistical significance. A faster rate of Na correction within the first 24 hours was not significantly associated with adverse outcome. This study suggests that the development of neurological complications could be related to an interaction of factors prior to presentation, including the severity of the hypernatraemia together with the clinical presence of shock. Larger multicentre trials are needed to further delineate these associations, to assist with management guidelines and reduce complications.

We recommend that greater emphasis should be placed on the pre-hospital management of patients with AGE, including educating caregivers on the correct use and constitution of ORT, appropriate practices and quick referral for help, to prevent the development of hypernatraemia and shock. Formula-fed infants should be identified as at risk for hypernatraemia. This study highlighted the poor documentation of ORT use and constitution, as well as formula dilution practices in patient records at the institution where the study took place. We suggest that existing admission templates be altered to remind staff to enquire about the relevant history in AGE cases.

In the management of infants with hypernatraemia following AGE, healthcare workers should aim for a reduction in serum Na between 0.5 and 1 mmol/L/hr, as per recommendations in the literature[1,2] to prevent any added risk of cerebral oedema.

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Author contributions. TH was responsible for study design, data collection, data analysis and drafting of the manuscript. KN and SR supervised the study and reviewed the manuscript.

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Conflicts of interest. None.

22. Duggal AK, Yadav P, Agarwal AK, Rewari BB. Clinical approach to altered serum sodium levels. JACM 2006;7(2):91-103.

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