Abstract

**Background:** Blood exchange transfusion (ECT) therapy is an essential part of the modern health care for high-risk icteric neonates.

**Objectives:** Therefore, this study sought to examine the effectiveness of monitoring sodium disorders and the required interventions indicating the overall state of health of hospitalized jaundiced neonates.

**Methods:** This prospective analytic study was performed on 49 neonates diagnosed with severe jaundice undergoing screened exchange blood transfusion from November 2018 to May 2019. Serum sodium ion concentrations testing was performed before the exchange, then 1 - 3 hours later, and again 24 hours after receiving ECT, using a sensitive ELISA kit in a laboratory. Using the newest version of SPSS 24 software program, the association between different numerical variables was calculated by a repeated-measure ANOVA test. P < 0.05 criterion was set as the threshold of significance.

**Results:** Out of the total 49 neonates, 24 (48.5%) were girls and 25 (51.5%) were boys, of which 93.9% and 6.1% were term and preterm-born neonates, respectively. The average birth weight was 2843 ± 284/510 g and the mean period of hospital stay was 4.55 ± 2.399 days. The various causes of severe neonatal jaundice of all the hospitalized cases were 45% unknown, 43% combined ABO and Rh incompatibility, 10% breastfeeding pattern, and 2% G6PD deficiency. A continual fall in serum sodium was observed following banked blood exchange (P < 0.05). However, these reversible changes were outside the expected normal range. Despite a diminished sodium concentration, the mean sodium levels were within the laboratory reference range of serum (137 - 142 mEq/L) in all the three measured duration times. Moreover, variation in the amount of serum sodium was associated with an unknown underlying cause that led to neonatal jaundice (P = 0.04) and this difference did not reach statistical significance in terms of birth weight, gestational age, and other causes accounting for high bilirubin (P > 0.05).

**Conclusions:** Given these facts, it was concluded that neonatal jaundice was significantly associated with post-exchange serum sodium changes within the laboratory reference range of serum.

**Keywords:** Jaundice, Exchange Transfusion, Neonatate, Sodium

1. Background

Neonatal hyperbilirubinemia is a very abundant, multifactorial, and quite challenging disorder resulting from elevated total serum bilirubin (TSB) level which is experienced within the neonatal period, particularly in the first week of life (1). This extremely prevalent medical problem affects approximately 8% - 11% of symptomatic infants who have the yellow coloration of the skin and sclera of the eyes caused by bilirubin (1). Currently, two clinically safe and effective treatment approaches are used in the most benign neonatal jaundice cases for reduction of the acute high bilirubin blood level, which are phototherapy and the acceptable rapid decrement method of exchange transfusion (2-4). According to the neonatal jaundice care guideline, both of these methods must be timely applied at total bilirubin serum level or when exceeding it, depending on the threshold of phototherapy and ECT treatments to avoid causal over-treatment complications (5, 6). Furthermore, all healthy term newborns with severe jaundice should be screened for simultaneous presence of the recurrence and exacerbation of persistent hyperbilirubinemia and potential high-risk presence of significant short-term and long-term side effects expressed with exchange transfusion therapy (2, 7). ECT therapy is the most common med-
clinical procedure for conditions that require immediate clinical intervention such as preterm infants, breast milk and feeding jaundice, ABO isoimmunization, and glucose-6-phosphate dehydrogenase (G6PD) inconsistencies (7). According to the pooled data from hospital-based case control studies, almost in two-thirds of these cases, ECT was not in a good condition and even newborn death could occur within seven days after the transfusion in up to 10 percent of the severe cases (8). In other words, because of the increasing adverse effects of significant complications and severe risk factors accompanied by blood exchange, this procedure is considered as the most frequent documented cause of infants’ fatality, with the reported incidence rate of about 10 to 74 percent (9). Some of the most often clinical events submitted to ECT were thrombocytopenia (44%), hypocalcemia (29%), and metabolic acidosis (24%), of which 69%, 74%, and 44%, respectively needed rescue treatment (10). Besides, serum sodium metabolic abnormalities together with calcium homeostasis disturbances (hypocalcemia) and thrombocytopenia are frequently observed in routine clinical practices (11). In other words, ECT is the primary identified contributing factor for the increased risk of encountering harmful hyperbilirubinemia recurrence in 18% of term newborns and it increases mortality in preterm infants and infants with other comorbidities (11).

Many previous studies have reported that the most important reason of the apparent sodium ion concentration change near the upper limit of standard value in most cases, during and after exchange blood transfusion, is poor nutrition in the first several days of life that leads to physiological loss of extracellular fluid’s volume, body water imbalance, and dehydration (5, 12-14). Additionally, serum sodium ion concentration disturbances following administration of an extensive quantity of stored blood products, which are typically sodium bicarbonate-rich solutions, could be deduced as poor prognosis of renal failure (15, 16). Shared clinical symptoms of many primary diseases in infancy such as meningitis, cardiac arrhythmias, sepsis, and congenital metabolic disorders, as well as hypoglycemia and hypocalcemia, can be mimicked by medical signs associated with sodium alterations (17). Therefore, preventing more severe abnormalities due to unnatural serum sodium ratio changes, especially neurological dysfunction and precipitation of severe heart failure and arrhythmia, could equally be supported by timely professional medical care interventions through ensuring well-organized treatment of transfusion-dependent patients (18,19). There are a few quantitative and qualitative surveys conducted for determining and modifying the increased risk of detrimental clinical outcomes correlated with possible fatal sodium abrupt changes during and after the exchange transfusion in newborns.

2. Objectives

Therefore, this study aimed to investigate the effectiveness of monitoring sodium disorders and the required interventions indicating the overall state of health of hospitalized jaundiced infants.

3. Methods

3.1. Experimental Pattern and Subjecting Technique

This prospective analytic study was performed on 49 neonates diagnosed with features of neonatal jaundice who were admitted to the neonatal ward of Abouzar Hospital in Ahvaz, Iran in order to undergo screened exchange blood transfusion according to the protocol authorized by the transfusion medicine unit of the hospital in a period from November 2018 to May 2019.

More specialized primary practices regarding pediatric healthcare were performed by physician assistants, including the critical need for blood exchange. Serum sodium ion concentration test of each of the desired samples was performed in a laboratory at three recommended points in time: Before, 1 - 3 hours after, and then 24 hours following blood exchange using the most sensitive and specific ELISA kits. In the next step, the collected data related to measured parameters was used to compare the considerable differences in these three times alongside other etiological risk factors. In this study, necessary hospital interventions and functional follow-up visits to patients who suffered from a severe disorder were done based on their presentation.

The ethics committee of Jundishapur University of Medical Sciences approved this clinical research (code: IR.AJUMS.REC.1398.541), and written informed consent was taken from all the parents or guardians of the subjects.

3.2. Statistical Analysis

The Kolmogorov-Smirnov test was used to examine the normal distribution of information. The statistics related to collected data with quantitative values were reported as mean, median, and standard deviation. In contrast, descriptive statistics involving frequency, the relative frequency, and percentage distribution were computed as appropriate summary measures for categorical or qualitative data. In the parametric multivariate analysis using the newest version of SPSS 24 software program, a
probability-calculating technique for determining statistical strength of the correlation between different numerical quantitative variables was assessed using a repeated-measures ANOVA test. The $P < 0.05$ criterion was set as the threshold of significance.

4. Results

4.1. Fundamental Individual Dataset and Specific Clinical Evidence of the Research Output Affiliated to the Experimental Trial

Sub-groups of the general clinical features of this study’s patients are demonstrated in Table 1. As shown there, in a total of 49 infants, 24 (48.5%) were girls and 25 (51.5%) were boys. The gestational ages for the statistical recording of neonatal jaundice varied from eight (17%) for more than 40 weeks, 28 (58%) for 37 to 40 weeks, 7 (16%) for 34 to 37 weeks, to four (9%) for 30 to 34 weeks. Birth weights of newborns were classified as 38 (77%) for babies that weighed between 2,500 to 4,000 g, 7 (15%) for those who weighed 1,500 to 2,000 g, and 1 (2%) for those weighing between 1,000 and 1,500 g, while, three (6%) was attributed to the newborns that weighed more than 4 kg (Table 1). The mean length of hospital stay of newborns of all the hospital deliveries was $4.55 \pm 2.399$ days. All neonates were clinically followed up for a maximum period of four days and exchange transfusion treatment and related therapeutic modalities were completed in four (8%) hospitalized patients within a maximum length of 10 days.

Only a 3-day-old 3,100 g male infant born at 37 weeks’ gestation had G6PD deficiency - leading cause of jaundice without manifestation of severe bacterial sepsis or other pathological detectable drivers of neonatal jaundice. Jaundice was observed a few days after delivery in four male breastfed term newborns with birth weights between 2,500 and 4,000 g, and gestational ages of 39 to 40 weeks. No dissimilarities were identified in the neurological investigation of these newborns. A 3,100 g 38-week gestational age girl was identified with breast milk jaundice in the first week after birth (the seventh day of life), while her medical examinations were normal. The observational trial of this study suggested that the fourth case that had ABO incompatibility and received double exchange transfusion due to acute hyperbilirubinemia, did not have comparatively lower bilirubin serum level, and one exchanged case notably had the highest bilirubin level only one day after, which was more than 25 mg/dL. Further exchange transfusion courses were performed mainly in 10 infants aged four days and at raised total serum bilirubin level of 28 mg/dL.

| Table 1. Demographic Baseline Characteristics Allocated to the Research Patients |
|-----------------------------------------------|---------------------------------------------|
| Characteristics                        | Patient (N = 49), No. (%)                        |
| Gestational age, wk                           |                                              |
| < 30                                         | 0 (0)                                        |
| 30 - 34                                      | 4 (9)                                        |
| 34 - 37                                      | 7 (16)                                       |
| 37 - 40                                      | 28 (58)                                      |
| > 40                                         | 8 (17)                                       |
| Gender                                       |                                              |
| Boy                                          | 25 (51.5)                                    |
| Girl                                         | 24 (48.5)                                    |
| Delivery weight, g                           |                                              |
| < 1000                                       | 0 (0)                                        |
| 1000 - 1500                                  | 1 (2)                                        |
| 1500 - 2000                                  | 7 (15)                                       |
| 2500 - 4000                                  | 38 (77)                                      |
| > 4 kg                                       | 3 (6)                                        |
| Causes of jaundice                           |                                              |
| Idiopathic                                   | 22 (45)                                      |
| ABO isoimmunization                          | 11 (23)                                      |
| Rh isoimmunization                           | 10 (20)                                      |
| Breast milk                                  | 1 (2)                                        |
| Breast feeding                               | 4 (8)                                        |
| G6PD inconsistencies                         | 1 (2)                                        |

4.2. The Account of Various Factors, Particularly Numerous Causes Leading to Jaundice, Different Birth Weights, and Actual Gestational Age Associated With Post-Exchange Serum Sodium Levels Disturbances

All the newborn infants with jaundice in this study that suffered from a broad range of clinical issues and had different birth weights and distinct gestational ages demonstrated significant mean sodium serum level declines at 1 to 3 hours and 24 hours after exchange transfusion in comparison to the time before the exchange. Despite a diminished sodium concentration, the mean sodium levels stayed within the laboratory reference range of the serum (137 - 142 mEq/L) at the three points in time mentioned above. In addition, variation in the amount of serum sodium was associated with an unknown underlying cause that led to neonatal jaundice ($P = 0.04$, Table 2) and this difference did not reach statistical significance in various trends in birth weight, gestational ages, and other causes accounting for high bilirubin ($P > 0.05$). The most common changes of the mean value in blood sodium level were noticeably found in infants aged six days. The high-
in the level of serum sodium was not observed in the study and after ECT in icteric neonates. Furthermore, alterations in sodium levels of about 2.8% to 3.2% were observed during and after blood exchange, in contrast to the time before the exchange transfusion. In addition, Jurfest-Ceccon et al. (24) showed that an increasing frequency in serum sodium levels was found that the mean sodium levels remained within the normal range directly after the ECT process in neonatal hyperbilirubinemia. This is consistent with the results of this original research on patients who had ECT, demonstrated a declining trend in serum sodium concentration in neonates, but these changes were outside an abnormal range of expected values. The same observations were reported in the prospective study of Esmaeilivand et al. (21), which delineated decreased sodium levels after ECT in 8% of neonates with serious jaundice. Despite their interventions, hyponatremia was detected in some of the jaundiced neonates. Consequently, sodium concentrations and serum sodium levels did not elevate (22). Similarly, the most recent literature has proposed that this method does not carry much risk for serum sodium imbalance in the situation of jaundice in newborn patients. For example, Calladine et al. (23) and Ballot and Rugamba (5) found that the mean sodium levels remained within the normal range directly after the ECT process in neonatal hyperbilirubinemia, which is consistent with the results of the present study. In comparison with these pilot feasibility studies and the current study, a large number of clinical transfusion trials have provided evidence that a considerable surge in serum sodium average in neonates and higher incidence of hyponatremia have happened during and after blood exchange, in contrast to the time before the exchange transfusion. In addition, Jurfest-Ceccon et al. (24) showed that an increasing frequency in serum sodium levels of about 2.8% to 3.2% was observed during and after ECT in icteric neonates. Furthermore, alteration in the level of serum sodium was not observed in the study of Wani et al. (8), in which they concluded that a heedful exchange transfusion strategy did not support this negative impact. Such dramatically different data across clinical practice fields related to neonatal jaundice and various researches that were mentioned above can be attributed to the differences in the clinical status of term neonates and survey periods. Given these facts, it is concluded that neonatal jaundice is significantly associated with changes in serum sodium level after receiving ECT, based largely on a baby's age and mean total serum bilirubin levels.

5. Discussion

A less aggressive exchange transfusion is capable of getting rid of extreme neonatal hyperbilirubinemia. This potentially emergency practice is regularly utilized in term infants for suppressing the occurrence of acute bilirubin encephalopathy (ABE), kernicterus features, and death (20). Reversible electrolyte imbalances, especially natural serum sodium and potassium disturbances, are the two most increasingly frequent abnormalities that are referred for administration of ECT therapy (14). To the best of the authors’ knowledge, there are a few conducted quantitative and qualitative surveys for determining and improving the increased risk of detrimental clinical outcomes correlated with possible fatal abrupt changes in sodium during and after the exchange transfusion in newborns. Therefore, this study was aimed to examine the effectiveness of monitoring sodium disorders and the required interventions regarding the overall state of health of hospitalized jaundiced neonates.

At first, primary stratified findings based on the results of this original research on patients who had ECT, demonstrated a declining trend in serum sodium concentration in neonates, but these changes were outside an abnormal range of expected values. The same observations were reported in the prospective study of Esmaeilivand et al. (21), which delineated decreased sodium levels after ECT in 8% of neonates with serious jaundice. Despite their interventions, hyponatremia was detected in some of the jaundiced neonates and serum sodium levels did not elevate (22). Similarly, the most recent literature has proposed that this method does not carry much risk for serum sodium imbalance in the situation of jaundice in newborn patients. For example, Calladine et al. (23) and Ballot and Rugamba (5) found that the mean sodium levels remained within the normal range directly after the ECT process in neonatal hyperbilirubinemia, which is consistent with the results of the present study. In comparison with these pilot feasibility studies and the current study, a large number of clinical transfusion trials have provided evidence that a considerable surge in serum sodium average in neonates and higher incidence of hyponatremia have happened during and after blood exchange, in contrast to the time before the exchange transfusion. In addition, Jurfest-Ceccon et al. (24) showed that an increasing frequency in serum sodium levels of about 2.8% to 3.2% was observed during and after ECT in icteric neonates. Furthermore, alteration in the level of serum sodium was not observed in the study of Wani et al. (8), in which they concluded that a heedful exchange transfusion strategy did not support this negative impact. Such dramatically different data across clinical practice fields related to neonatal jaundice and various researches that were mentioned above can be attributed to the differences in the clinical status of term neonates and survey periods. Given these facts, it is concluded that neonatal jaundice is significantly associated with changes in serum sodium level after receiving ECT, based largely on a baby’s age and mean total serum bilirubin levels.

5.1. Conclusions

In summary, notwithstanding the highly therapeutic effectiveness of ECT modality as a suitable alternative treatment option for neonatal jaundice (NJ), this approach has its dire consequences. Jaundiced neonates were most susceptible to electrolyte imbalance panel, which is the tendency for rapid decline in serum sodium levels. Certainly, future studies with larger populations and longer follow-up periods are needed that can directly lead to further examination of significant negative electrolyte changes coincided with ECT therapy and long-term follow-up treatment of infant patients showing the characteristics of jaundice. Therefore, the task of differential diagnosis of direct ongoing health side-effects produced by ECT medication in icteric neonates could be effective in early diagnosis of relative clinical manifestations and indications and inhibiting several principal causes of elevated morbidity or mortality rates that occur due to sodium disturbances and other serious complications.

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Footnotes

Authors’ Contribution: AKH, and MZ contributed to the design and implementation of the research. AS did the analysis of the results and verified the analytical methods. SRM carried out the experiment, manuscript writing and editing.

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**References**

1. Ullah S, Rahman K, Hedayati M. Hyperbilirubinemia in Neonates: Types, Causes, Clinical Examinations, Preventive Measures and Treatments: A Narrative Review Article. *Iran J Public Health*. 2016;45(5):558-68. [PubMed: 27398128]. [PubMed Central: PMC4936999].

2. Bujandric N, Grujic J. Exchange Transfusion for Severe Neonatal Hyperbilirubinemia: 17 Years’ Experience from Vojvodina, Serbia. *Indian J Hematol Blood Transfus*. 2016;32(2):208-14. doi: 10.1007/s12288-015-05344. [PubMed: 27065585]. [PubMed Central: PMC4790007].

3. Alizadeh Taheri P, Sadeghi M, Sajjadian N. Severe neonatal hyperbilirubinemia leading to exchange transfusion. *Med J Islam Repub Iran*. 2014;28(4). [PubMed: 25405299]. [PubMed Central: PMC4219984].

4. Edris AA, Ghany EA, Razek AR, Zahran AM. The role of intensive phototherapy in the management of severe neonatal hyperbilirubinemia. *Indian J Med Res*. 2014;140. [PubMed: 24605703].

5. Ballot DE, Rugamba G. Exchange Transfusion for Neonatal Hyperbilirubinemia in Johannesburg, South Africa, from 2006 to 2011. *Int Sch Res Notices*. 2015;2016:1268149. doi: 10.1155/2016/1268149. [PubMed: 27382636]. [PubMed Central: PMC4937011].

6. Zardosht R, Shah Farhat A, Saedii R, Parvin F. Assesment Efficacy and Complication of the Distance between Phototherapy Lamps and Neonate’s Body Level on Serum Bilirubin Decrease and Phototherapy Complications in Neonatal Hyperbilirubinemia. *Iran J Neonatol IJN*. 2010;10(1):47-52.

7. Sabzehei MK, Basiri B, Shokouhi M, Torabian S. Complications of Exchange Transfusion in Hospitalized Neonates in Two Neonatal Centers in Hamadan, A Five-Year Experience. *J Compr Pediatr*. 2015;6(2). doi: 10.17795/compreped-20547.

8. Wani MI, Mudasir N, Roumisa L, Mohd R, Syed Wajid A, Bashir Ahmad C. Impact of Double Volume Exchange Transfusion on Biochemical Parameters in Neonatal Hyperbilirubinemia. *Int J Pediatr Res*. 2018;4(4). doi: 10.23937/2469-5769/1500038.

9. Yu C, Li H, Zhang Q, He H, Chen X, Hua Z. Report about term infants with severe hyperbilirubinemia undergoing exchange transfusion in Southwestern China during an 11-year period, from 2001 to 2011. *PLoS One*. 2017;12(6). e0179550. doi: 10.1371/journal.pone.0179550. [PubMed: 28662081]. [PubMed Central: PMC5491324].

10. Ibehwe RC, Ibehwe MU, Muoneke VU. Outcome of exchange blood transfusions done for neonatal jaundice in abakaliki, South eastern Nigeria. *J Clin Neonatol*. 2012;1(1):34-7. doi: 10.4103/2249-4847.92239. [PubMed: 24027883]. [PubMed Central: PMC768882].

11. Rajagopal R, Chakraborti S, Grover S, Khehra N. Knowledge, experience & attitudes concerning electroconvulsive therapy among patients & their relatives. *Indian J Med Res*. 2012;135:201-10. [PubMed: 22444682]. [PubMed Central: PMC338851].

12. Kumar P, Chacham S, Kumar J, Dutta S. Adverse events following blood exchange transfusion for neonatal hyperbilirubinemia: A prospective study. *J Clin Neonatol*. 2019;8(2). doi: 10.4103/jcn.JCN_96_18.

13. Opoku-Okrah C, Acquah BK, Dogbe EE. Changes in potassium and sodium concentrations in stored blood. *Pan Afr Med J*. 2015;20:236. doi: 10.11604/pamj.2015.20.236.5801. [PubMed: 27386012]. [PubMed Central: PMC4919675].

14. Antwi-Baffour S, Adjei JK, Tryawo F, Kyerehene R, Botchway FA, Seidu MA. A Study of the Change in Sodium and Potassium Ion Concentrations in Stored Donor Blood and Their Effect on Electrolyte Balance of Recipients. *Biomed Res Int*. 2019;2019:862897. doi: 10.1155/2019/862897. [PubMed: 31662997]. [PubMed Central: PMC679281].

15. Sá CAM, Santos MCP, Carvalho MD, Moreira MEL. Eventos adversos associados à exsanguineotransfusão na doença hemolitica perinatal: experiência de dez anos. *Rev Paul Pediatr*. 2014;32(2):168-72. doi: 10.1590/S0104-74012014005000006.

16. Chen W, Abramowitz MK. Treatment of metabolic acidosis in patients with CKD. *Am J Kidney Dis*. 2014;63(4):473-80. doi: 10.1053/j.ajkd.2013.06.017. [PubMed: 23932089]. [PubMed Central: PMC3949691].

17. Kovesdy CP. Pathogenesis, consequences and treatment of metabolic acidosis in chronic kidney disease. 2014. Available from: http://UpToDate.com.

18. Satar M, Arisoy AE, Celik IH. Turkish Neonatal Society guideline on neonatal infections-diagnosis and treatment. *Turk Pediatr Ars*. 2018;53(3):558-560. doi: 10.3752/turkpediarsars.2018.018099. [PubMed: 31236022]. [PubMed Central: PMC6568293].

19. Espay AJ. Neurologic complications of electrolyte disturbances and acid-base balance. *Handb Clin Neurol*. 2014;119:365-82. doi:10.1016/B978-0-7020-4086-3.00023-0. [PubMed: 25280346]. [PubMed Central: PMC4464645].

20. Keshavarzkhani I, Asadian S, Siavash V, Mohammadi M, Siavashi A. [Comparison of the transfusion complications in term and preterm neonates with jaundice]. *J Babol Uni Med Sci*. 2016;18(9):49-55. Persian.
22. Mabogunje CA, Olaifa SM, Olusanya BO. Facility-based constraints to exchange transfusions for neonatal hyperbilirubinemia in resource-limited settings. World J Clin Pediatr. 2016;5(2):182-90. doi: 10.5409/wjcp.v5.i2.182. [PubMed: 27170928]. [PubMed Central: PMC4857231].

23. Calladine M, Gairdner D, Naidoo BT, Orrell DH. Acid-base changes following exchange transfusion with citrated blood. Arch Dis Child. 1965;40(214):626-31. doi: 10.1136/adc.40.214.626. [PubMed: 5891758]. [PubMed Central: PMC2019484].

24. Jurfest-Cecon ME, de Albuquerque Diniz EM, de Araujo-Ramos JL, Costa-Vaz FA. Biochemical and hematometric effects of exchange transfusion in isoimmunized neonates. Bol Med Hosp Infant Mex. 1993;50(3):367-76. Spanish. [PubMed: 8442881].