CLINICAL PROFILE AND OUTCOME OF DIABETIC KETOACIDOSIS IN CHILDREN AT TERTIARY CARE HOSPITAL

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ABSTRACT: OBJECTIVE: To study the clinical profile of children admitted with diabetic ketoacidosis (DKA) in Sri Aurobindo Institute of Medical Sciences (SAIMS) Indore. **METHODS:** We retrospectively analysed the case records of 30 children admitted with DKA to our intensive care unit, from Jan 2011-Jan 2015. Information regarding personal details, chief complaints, clinical features, laboratory parameters, management and outcome was recorded using a predesigned pro forma. **RESULTS:** The median age at presentation was 8 years (range 9months- 13yrs); 12 boys and 18 girls were enrolled. Diabetes was newly diagnosed in 20 cases and known cases of type1 diabetes were 10. Commonest presenting complaints were fever (66%), rapid breathing (60%), vomiting (60%), and altered sensorium (26. 6%). Average length of PICU stay was 58. 6hrs. All of the cases had elevated HbA1c, except one. Co-morbities found were UTI in 12 (40%), dysentery in 2 (6. 6%), and viral hepatitis in1 (3. 33%). Cerebral edema was seen in 1 case. There was no mortality in our study. **CONCLUSION:** DKA in children if diagnosed early and managed timely has good outcome. Poor compliance is most important precipitating factor in known cases of T1DM.

KEYWORDS: Diabetic Ketoacidosis, Type I Dm, Cerebral Edema, Delayed Diagnosis.

INTRODUCTION: Diabetic ketoacidosis is a major complication of type1diabetes mellitus (T1DM) in children and is associated with increased risk of morbidity and mortality. Though type 1DM accounts for most cases of diabetes in childhood, approximately 50% of individuals present in adulthood. Around 20-40% of children with newly diagnosed type 1 DM present in DKA.^[1] Diabetic ketoacidosis accounts for nearly 0. 6 % of total PICU admissions.^[2] Though clinical features and management guidelines for DKA are well known, a high index of clinical suspicion is necessary for early diagnosis and timely intervention. There is lack of published studies on DKA from India. This study was done to study the clinical profile and outcome of DKA in central India.

MATERIALS AND METHODS: A retrospective study was done in PICU of tertiary level care hospital at Indore. Case records of children admitted with DKA from Jan 2011 to Jan 2015 were reviewed and data regarding personal details, presenting complaints, clinical features, family history of type1DM, laboratory parameters, management, duration of PICU stay and outcome was recorded using a predesigned pro forma.

DKA was diagnosed when blood sugar at admission was >200mg/dl with acidosis (pH <7.35, bicarbonates <15), and positive urine ketones. Severity was graded as mild, moderate and severe depending on blood gas and clinical features.^[1] Monitoring of heart rate, respiratory rate, blood pressure, level of consciousness and fluid charting was done hourly. Capillary blood glucose was measured every 2 hourly. Urine ketones, serum electrolytes, and arterial/venous blood gas was measured 4hrly.

J of Evolution of Med and Dent Sci/ eISSN- 2278-4802, pISSN- 2278-4748/ Vol. 4/ Issue 31/ Apr 16, 2015 Page 5329

ORIGINAL ARTICLE

Milwaukee protocol was followed for 24-36hr depending upon the severity of DKA. Insulin infusion was stopped when child was alert, tolerating oral feeds, metabolically stable. Initially rapid short acting regular insulin (human actrapid) was given subcutaneously. Then it was replaced by mixed insulin (30/70) once blood sugar stabilized. All children were given i/v ceftriaxone until blood/urine culture turned out sterile.

RESULTS: The median age at presentation was 8 years (range 9months- 13yrs) with A F: M ratio of 1. 5: 1. Eighteen females and 12 males were affected. Mean duration of symptoms before hospitalization was 5 days. Family history of type 1 DM was present in 2 cases (6.66%, in both cases sibling was affected.) Commonest presenting complaints were fever in 20cases (66%), rapid breathing in18 (60%), vomiting in 18 (60%), altered sensorium in 8 (26.6%), loose motions in 4 (13.33%), and seizures in 1 (3.33%). Classical h/o polyuria and polydipsia were seen only in 5 cases (16.6%). Among 30 cases, 20 were new cases and 10 were known cases of T1DM. Out of 10 known cases of T1DM incompliance was the cause of DKA in 6 (60%) and intercurrent illness in 4cases (40%). Average length of PICU stay was 58. 6hrs and for discharge from hospital was 7. 8days. The median time for the arterial blood gases to become normal was 26 hrs and changing over to subcutaneous insulin was 1.6 days. 29 cases were discharged and 1 got LAMA due to inaffordability after 48hrs of admission. She was 13year female was in ARF and required haemodialysis and ventilator. In associated co morbidities, UTI was seen in 12 cases, hypothyroidism in 1case, viral hepatitis in 1, acute renal failure in 1 and dysentery in 2 cases. Cerebral edema was seen in 1case.

DISCUSSION: Diagnosis of DKA is often missed, as classical history of polyuria and polydipsia is generally unavailable in children. The younger the child, the more difficult it is to diagnose DKA, especially with new-onset or previously undiagnosed diabetes mellitus (DM). In infants and young children symptoms may be nonspecific and this needs a high index of suspicion to diagnose DKA.

There exists difficulty in recognizing the symptoms of diabetes among parents and physicians. Studies have reported that unusual bed-wetting in a previously "dry" child occurred long before the diagnosis of T1DM; however, only in a few cases these parents were able to suspect the presence of hyperglycemia.^[3] Vigilance of parents and caretakers play an important role in early identification of T1DM. In a developing country like India, many a times due to social and economic reasons also first visit to the physician is delayed. Previous studies have shown that lack of parental and physician's awareness and delay in referral to appropriate centre as the major high risk factors for delayed diagnosis and mortality in DKA.^[4]

Ali et al^[5] had reported in 2011 that 30% of newly diagnosed children have had at least one related medical visit prior to diagnosis, suggesting the condition is being missed by doctors. Literature reports that DKA has been misdiagnosed as surgical emergencies with acute abdomen.^[6]

Lack of awareness about signs and symptoms of T1DM among general people and primary care physicians contributes to delay in diagnosis and more cases landing up in DKA.

The median age at presentation in our study was 8years where as two earlier studies reported it to be 9.1 and 8.2 years.^[7,8] Though literature suggests equal incidence in both genders, in our study the frequency of DKA was higher among girls similar to Neu etal.^[9] and Madoori etal.^[7] Most common presenting complaints was fever followed by vomiting and rapid breathing, in contrast to impaired level of consciousness as reported by jayshree etal.^[10]

ORIGINAL ARTICLE

History of symptoms of polyuria and polydipsia were given in only 5 (16.6%) cases as opposed to other studies^[3,7] which reported it to be 25% and 85% respectively. Most of our patients had new onset type 1DM. Major precipitating factors for DKA were infections (most commonly viral fever and UTI). Most common preventable cause of DKA among known cases of type1 DM was incompliance.

There was no mortality in our study group. Cerebral edema was seen in 1 case. Previous studies have reported life threatening cerebral edema in. 5-1% of DKA episodes.^[11,12] Symptomatic cerebral edema has a high mortality rate (21%–24%) with a substantial proportion of survivors (15%–26%) left with permanent neurologic sequelae.^[13] Mild, asymptomatic cerebral edema probably is present in most children who have DKA at the time of presentation and during therapy.^[14] Complications of DKA may extend beyond the acute period and may affect neurocognitive functions and quality of life in substantial number of children. Recent data suggest that DKA episodes without clinically overt cerebral edema have also been associated with permanent defects in memory function.^[15,16] DKA children should be followed regularly and assessed regarding neurocognitive impairments along with other complications of prolonged hyperglycemia.

Our study has certain limitations. Firstly, we studied DKA only in children less than 14 years of age, as the criteria for admission in our PICU was below 14yrs. Secondly, being a retrospective analysis, follow up of the cases could not be done.

Long term prospective trials are required for definitive rehydration therapy during first 24-48hrs. Proper management during initial hours lead to favorable outcome without life threatening complications like cerebral edema. Counseling and education of parents during hospitalization and discharge is very necessary, so that parents understand the importance of insulin injections. Regular follow up regarding long term complications of type 1DM is important.

REFERENCES:

- 1. Nelson textbook of paediatrics 19th edition, page no. 1976.
- 2. Burns MR, Bodansky HJ, Parslow RC. Paediatric intensive care admission for acute diabetes complication. Diabet Med 2010; 27: 705-8.
- 3. Maurizio vanelli, Giulia costi, Giovanni chiari, Lucia ghizzoni, Tommaso giacalone, Francesco chiarelli. Effectiveness of prevention programme for diabetic ketoacidosis in children. Diabetec care 1999; 22: 7-9.
- 4. Varadarajan Poovazhagi. Risk factors for mortality in children with diabetic ketoacidosis from developing countries. World J Diabetes. 2014; 5 (6): 932-938.
- 5. Ali K, Harnden A, Edge JA. Type 1 diabetes in children. BMJ. 2011; 342: d294 [PMID: 21325386 DOI: 10. 1136/bmj. d294].
- 6. Durai R, Hoque H, Ng P. The Acute Abdomen Commonly missed and mis-diagnosed Conditions: Review. Webmed Central Surgery 2010; 1: WMC001036 [DOI: 10. 9754/journal. wmc. 2010. 001036].
- 7. Madoori Srinivas, Kapil Challa., Kadiri Bhanu Varun, Shantappa Kallapalli and Srikanth. Clinical profile of diabetic ketoacidosis in children admitted at tertiary hospital karimnagar district. International journal of recent scientific research 2014; 5: 233-235.
- 8. Ramaswamy ganesh, R. arvind kumar, Thiruvengadam vasanthi. Clinical profile and outcome of diabetic ketoacidosis in children. The National medical Journal of India 2009; 22; 18-19.

J of Evolution of Med and Dent Sci/ eISSN- 2278-4802, pISSN- 2278-4748/ Vol. 4/ Issue 31/ Apr 16, 2015 Page 5331

- 9. Neu A, Willasch A, Ehehalt S, Hub R, Ranke MB. Ketoacidosis at onset of type1 diabetes mellitus in children- frequency and clinical presentation. Pediatr Diabetes 2003; 4: 77-81.
- 10. Jayashree M, Singhi S. Diabetic ketoacidosis: Predictors of outcome in a paediatric intensive care unit of a developing country. Pediatr Crit Care Med 2004; 5: 427–33.
- 11. Edge J, Hawkins M, Winter D, Dunger D. The risk and outcome of cerebral oedema developing during diabetic ketoacidosis. Arch Dis Child. 2001; 85: 16–22.
- 12. Glaser N, Barnett P, Mc Caslin I, et al. Risk Factors for Cerebral Edema in Children with Diabetic Ketoacidosis. N Engl J Med. 2001; 344: 264–9.
- 13. Wolfsdorf J, Glaser N, Sperling MA. Diabetic ketoacidosis in infants, children and adolescents: A consensus statement from American Diabetes Association. Diabetic Care. 2006; 29: 1150-9.
- 14. James P. Orlowski, Cheryl L. Cramer, Mariano R. Fiallos. Diabetic Ketoacidosis in the Paediatric ICU. Pediatr Clin N Am. 2008; 55: 577–587.
- 15. Ghetti S, Lee J, Holtpatrick C, DeMaster D, Glaser N. Diabetic ketoacidosis and memory impariment in children with Type 1 diabetes. J Pediatr. 2010; 156: 109–14.
- 16. Nicole S. Glaser, Simona Ghetti, T. Charles Casper, J. Michael Dean, Nathan Kuppermann. Pediatric Diabetic Ketoacidosis, Fluid Therapy and Cerebral Injury: The Design of a Factorial Randomized Controlled Trial. Pediatr Diabetes. 2013; 14 (6): 435–446.

Age group (years)	Girls	Boys	Known case of T1DM	New Case
<5	4	4	1	7
5-10	8	5	6	7
>10	6	3	3	6
Total	18	12	10	20

Table 1: Age and Sex wise distribution of children



ORIGINAL ARTICLE

Age group (in years)	Mild	Moderate	Severe		
<5	2	-	6		
5-10	3	3	7		
>10	2	2	5		
Total	7	5	18		
Table 2: Severity of DKA among various age groups					

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