DISTAL RENAL TUBULAR (TYPE 1) ACIDOSIS- A CASE REPORT

Shyamali Datta¹, Jasninder Singh²

¹Assistant Professor, Department of Paediatrics, Mata Gujri Memorial Medical College, Kishanganj, Bihar.
²Postgraduate Student, Department of Paediatrics, Mata Gujri Memorial Medical College, Kishanganj, Bihar.

ABSTRACT

BACKGROUND

A 7-year-old female child presented with growth failure, features of rickets and history of multiple attacks of paralysis. On examination she was retarded in height and weight, had flaccid paralysis of limbs and features of rickets which was not responding to Vitamin D therapy previously. On investigation she showed metabolic acidosis, hypokalaemia, high urinary pH nephrolithiasis and increased renal cortical echogenicity in ultrasonography. Treatment with alkali and potassium was satisfactory, vitamin D and calcium were also given.

KEYWORDS

Distal Renal Tubular Acidosis, Rickets (Refractory), Hypokalaemic Periodic Paralysis, Growth Failure.


BACKGROUND

Case Report

A seven years old female child of non-consanguineous marriage from a distant Bihar village, presented with Growth failure, Bone deformities, Polyuria and episodes of Gastroenteritis with vomiting followed by weakness of limbs. For bony deformities, she was treated with Injectable Vitamin D at least on three occasions during her lifetime in last three - four years (as seen in her treatment records), but with no results.

On examination, she showed retardation of weight and height (weight for age= 22 kgs, whereas her body weight was 16.5 kgs i.e. - 1 SD;² Height for age= 119 cm, whereas her height was 100.5 cm i.e. - 2 SD). Widen ing of bone ends at elbow, wrist, knee and ankle seen,³,⁴ which the mother told to be better after present hospital treatment. Muscle power and tendon jerks are diminished,⁵ sensory system unaffected. No cardiorespiratory problem seen, only tachycardia noted on several occasion (92 - 96/minute), ECG normal, Ear and Eye examination showed no abnormality. No sibling history of similar disease. No H/O exposure to any industrial waste⁶ or any H/O of any tumour or prolonged drug therapy.⁷ As she was admitted with Gastroenteritis and Dehydration, she was given IV Ringer Lactate.

On investigation Blood Count was normal, no significant pallor, blood urea 32.1 mg/dL and creatinine was 0.81 showing good renal function. Liver function tests were normal. X-ray bones showed widening and fraying of epiphysis- suggestive of Rickets (which did not respond to multiple Vitamin D doses at past.³ Serum Sodium 132 mEq/L and Potassium 2.6 mEq/L were low in more than one instances. Serum Calcium was low normal (7.5; 7.9 mEq/L). Serum Bicarbonate was

12.5 mEq/L- low. Urine pH was high at 7.25 with +1 Albumin. Blood paO2 82.6 mmHg and pCo2 22.8 mmHg, pH 7.265. Ultrasonography showed increased cortical echogenicity of left-sided kidney and 3.8 mm single stone in left renal mid-pole. The girl was given oral Sodium Bicarbonate 2 mEq/kg in three divided doses⁸ and oral Potassium Chloride 2 mEq/kg daily. Injectable Vitamin D and Calcium orally was also given. Followup after three months showed no further paralytic incidence, no progression of bony lesions, growth also improved as evident from gain in body weight.

Financial or Other, Competing Interest: None.
Corresponding Author:
Dr. Shyamali Datta,
Ashirbad Apartment, Flat 2A,
Khudiram Sarani,
Cooch Behar-736101,
West Bengal.
E-mail: data.shyamali@gmail.com
DOI: 10.14260/Jemds/2017/538
DISCUSSION
Defective secretion of Hydrogen ion (H+) in distal renal tubule or reduced proximal renal tubular reabsorption of Bicarbonate (HCO3-) leads to metabolic acidosis. Urine pH remains high. Renal function is usually normal. It may be sporadic or secondary to Wilms' Tumour Drugs ifosfamide
Heavy metal Lithium or Genetic (cystinosis).

Our case was sporadic with Hypokalaemic muscle weakness (seen in Type I variety) due to urinary potassium loss. Refractory Rickets is due to bone demineralisation in an attempt to buffer acidosis is seen in Type I variety. Renal calculi is also seen in distal renal tubular acidosis (Type I variety). Growth retardation is due to mobilisation of organic materials from bone in an attempt to buffer metabolic acidosis. Bowing of lower extremities is related to weight bearing at the age of walking. Waddling gait in such children, Genu valgum; Genu varum; short stature is also seen.
Radiographic findings include metaphyses widening, fraying and coarse appearing of trabecular bones, Cupping of metaphyses occurs at proximal and distal tibia and at distal femur, radius and ulna bone.

CONCLUSION
So the case according to our clinical examination, investigation, treatment pattern and response tally very well to be a patient of Distal Renal Tubular Acidosis Type 1 variety.

REFERENCES