BARTTER SYNDROME, A RARE CAUSE OF MALNUTRITION IN CHILDREN

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ABSTRACT
Three and half year old Haleema presented to OPD with complaints of failure to thrive polydipsia and polyuria. In past she visited different clinicians and hospitals. Scrutiny of the previous record showed alkalosis persistently, but other electrolytes values were different from different labs. We admitted her and investigations were repeated, which showed hypo-kalemia, hypo-chloremia, hypo-natremia, metabolic alkalosis, hyper-calciuria, and normal urine osmolality and specific gravity. Her plasma rennin and aldosterone level were also raised. She was discharged on treatment for Bartter syndrome. On follow up, 15 days later her weight and clinical symptoms were improved.

KEY WORDS:
Bartter Syndrome, Malnutrition, Children.

INTRODUCTION
Malnutrition is the most prevalent public health concern in most of under developed and developing countries. Poor household level of food security, access to health and sanitation services and child caring practices are the three main reason which results in malnutrition in our setup1. In south Asian countries, malnutrition is the most common public health issue in children under the age of five years. About half of the world’s malnourished children reside in India, Pakistan and Bangladesh. In developing countries, 16% of children are born with low birth weight. In south Asia, 27% of neonates are born with low birth weight. In Pakistan, 27% of our children <5 years are underweight and 32% are stunted. 42% of our pregnant women and 47% of our children<5 years are anemic. In last, nearly 50% of our population is malnourished1, 2.

Good nutrition in the very young age is very important because the effects of under nutrition in the early life are long lasted in terms of physical, social, mental and intellectual development of the child2. The most immediate and dangerous complication of malnutrition is premature death. The risk of death is increased even with mild malnutrition and thereby it increases exponentially as the severity of malnutrition is increased. If a child reaches his 1st birthday in malnourished state, he suffers irreversible physical and cognitive damage, and his future life, welfare and economic wellbeing are badly affected1. In south Asia Pakistan has the second highest infant and child mortality rates3. In 2005 hours under 5 years child mortality rate was 101/1000 live births and malnutrition underlies much of this child mortality4. There are so many causes of malnutrition, which are broadly divided into primary and secondary. Primary malnutrition is because deficient intake of calories/proteins, while secondary is because of excessive caloric expenditure that is not met, or food quantity and quality are good but it is not assimilated properly1. In this report we will present a case of secondary malnutrition which was extensively investigated at different hospital in Peshawar, but no cause was found. She had a changing picture of clinical features, and so she was investigated at different lines. She received different treatment regimens, but with very little improvement. And at last she was proved to be a case of Bartter syndrome.
CASE REPORT

Three and a half year old Haleema presented to us in OPD on 19th May 2015, with the main complaint of not gaining weight. She also had polyuria, polydipsia and vomiting since last two years. On examination, she was emaciated, irritable and having doll’s like faces. Her weight was 7.8kg, and height was 88cm. She always had a bottle of water in her hands. Both her height and weight were below the 5th centile on WHO growth charts. She was dehydrated. Her pulse rate was 110BPM. Her respiratory, abdominal, cardiovascular and central nervous system examination was unremarkable.

When she was one and a half years old she was admitted in a tertiary care hospital in Peshawar, with complaints of vomiting and not gaining weight. There her serum electrolytes were deranged with raised potassium and decreased sodium. For that reason her serum 17-OH progesterone level was also done that was normal. There her ABGs were also done that showed PH of 7.624, HCO3 of 20.5mmol/L, PCO2 of 20.3mmHg, and PO2 of 82.2mmHg. Also her ultrasound report was normal. But unfortunately they got discharge on will, before she was properly diagnosed. After that, in the last two years she visited different hospitals and private clinics, with different set of complaints, the common to all was not gaining weight, polydipsia and polyuria. Here with us, her urine examination was normal, with normal osmolality and specific gravity. Serum sodium was 132.6mmol/L, potassium was 2.04mmol/L and chloride was 91.3mmol/L. Serum alkaline phosphatase was normal but serum calcium was 8.2mg/dl. Serum magnesium was within normal range. Her serum creatinine and urea was also normal. Her Hb was 10g/dl. TLC, DLC and platelets count was also normal. MCV was 68.2fl. Her ABGs showed PH of 7.58 and HCO3 of 31.1mmol/L. Her ultrasound abdomen showed bilateral tiny renal concretions. Urinary chloride was 286mmol/24 hours (normal: 110-250), and urinary calcium was 380mmol/24 hours (normal: 100-300). Plasma rennin level was >500uIU/ml (normal: erect=4.4-46.1, supine=2.8-39.9) and aldosterone level was 35ng/dl (normal: standing=4-31, recumbent=1-16). So she was diagnosed to be a case of Bartter syndrome and treatment was started. On follow up, 15 days after discharge her weight was 8.6kg, and she was improved symptomatically.

DISCUSSION

Bartter syndrome is a rare disorder of renal tubules. It was 1st described in 1962 by Frederich Bartter and his colleagues. It is clinically and genetically hetero-genetic disorder characterized by renal salt wasting, hyper-calciiurea, hypo-kalemia, metabolic alkalosis, and normotensive hyper-reninemic hyperaldosteronism. It is transmitted in auto-somal recessive fashion; therefore it is more common in the off springs of consanguineous parents. Clinically Bartter syndrome is classified into two groups, depending upon the age at onset, antenatal/neonatal and classic. Genetically barter syndrome is classified into five types, depending on the underlying gene mutation. All of these mutant genes are expressed in the tubular epithelial cells of thick ascending limb of loop of henle. In type-1 there are loss-of-function mutations of SLC12A1 which encodes the apical sodium-potassium-chloride co transporter (NKCC2). In bartter syndrome type-2 there are loss-of-function mutations of KCNJ1, which encodes the apical inwardly-rectifying potassium channel (ROMK). BS type III is caused by loss-of-function mutations of CLCNKB which encodes the baso-lateral chloride channel (CiC-Kb). BS type IV is caused by loss-of-function mutations of BSND which encodes barttin and BS type V is caused by gain-of-function mutations of CASR which encodes the baso-lateral calcium sensing receptor (CASR). These defects results in defective re-absorption of sodium, potassium, chloride, hydrogen iron and calcium in the thick ascending limb of loop of henle, subsequently leading to hypo-kalemia, hypo-chloremic, metabolic alkalosis. Although hyper-calciiurea is an important feature, hypo-calcemia and hypo-natremia is usually not that much severe. The antenatal/neonatal bartter usually presents in neonatal or infantile life, with severe phenotypic
features like neonatal salt wasting and recurrent dehydration. These patients also usually have history of maternal poly-hydromnios. The classic bartter syndrome usually presents in childhood, with milder phenotype of recurrent episodes of mild dehydration, polydipsia, polyuria and failure to thrive. The dysmorphic features like triangular faces, large eyes and protruding ears can be present in both types\textsuperscript{5,7}. The diagnosis is mostly based on clinical history and characteristic bio-chemical picture of hypo-kalemia, hypo-chloremia, metabolic alkalosis, hyper-calciiuria and increased excretion of chloride in urine\textsuperscript{7}.

Treatment is supportive, by preventing dehydration, care of good and enough nutrition, and correcting the electrolytes derangement. These patients especially, needs very high amount of potassium daily. Potassium sparing diuretics are also helpful. Indomethacin is also given especially in antenatal/neonatal form. With accurate management the prognosis is usually good. Some patient may develop chronic interstitial nephritis and chronic renal disease\textsuperscript{7}. Our patient was a case of classic Bartter syndrome, leading to secondary malnutrition. Although Bartter syndrome is not a common cause of malnutrition, this case report will emphasize not to ignore the rare causes of malnutrition, if we cannot find any common cause. Putting all the malnutrition into the primary one, after investigating the common causes, and ignoring the important clues in history is not a realistic approach.

REFERENCES


CORRIGENDUM

In the issue Vol. 02 No.01 (Sep 2015-March 2016) of JGMDS in the CASE REPORT titled “Fibromatosis In Maxillofacial Region”, the name of the authors were aberrantly printed by omitting two authors. Henceforth the author list for the said article should be read as under:

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