Sjögren syndrome and lupus erythematosus nephritis

A similar case of Sjögren syndrome and subsequent development of systemic lupus erythematosus has been reported in a 12 year old boy. In our patient, two points remain to be clarified. whether the seizures were the first symptom of systemic lupus erythematosus central nervous system involvement and whether any relation can be found between the phenobarbitone treatment and the onset of systemic lupus erythematosus.

Interpretation of sequential renal biopsies. The first renal biopsy showed notable interstitial lymphoplasmocytic infiltrate without glomerular alteration. Similar features have been reported in Sjögren syndrome, but in some cases of systemic lupus erythematosus, where interstitial lymphocytic infiltrate has been found on renal biopsy, the initial nephropathy worsened and rapidly progressed to renal failure. On the second renal biopsy a mild diffuse proliferative nephritis typical of systemic lupus erythematosus was found, but the interstitial lymphocytic infiltrate persisted. Thus, the sequential renal changes in our patient may have been those of systemic lupus erythematosus, with lymphocytic interstitial infiltrate at the beginning, progressing to the typical glomerular lesions.

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References


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Systemic carnitine deficiency exacerbated by a strict vegetarian diet

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SUMMARY A 12 year old boy suffered episodes of vomiting, lethargy, and hypoglycaemia from the age of 1 year. Adhering to a vegetarian diet caused an increase in frequency and severity of the attacks. It was found that he was suffering from systemic carnitine deficiency that responded promptly to treatment with L-carnitine.

Carnitine deficiency, first described in 1973, can manifest itself in two forms—myopathic and systemic. In myopathic carnitine deficiency, carnitine concentrations are reduced only in muscle, but in the systemic form, carnitine concentrations are reduced in several tissues, and plasma concentrations are also reduced. The patients often present in infancy with features resembling Reye’s syndrome, (encephalopathy, fatty liver, raised hepatic enzymes, and hypoglycaemia). Recently, dietary dependent carnitine deficiency has been described. We report a child with systemic carnitine deficiency that was aggravated by a strict vegetarian diet. treatment with L-carnitine resulted in a dramatic improvement.

Case report

The patient, who is now 12 years old, was first evaluated at the age of 3 years. He was born after a normal pregnancy and delivery to healthy parents. His birthweight was 3.2 kg. At the age of 1 year he suffered his first episode of vomiting and lethargy, and low serum glucose values were noted. He was treated with intravenous glucose and was discharged after three days. Since then similar episodes occurred every few months. When first evaluated at our hospital a blood glucose of 20 mg/dl (1-1 mmol/l)
and raised hepatic enzymes were noted during an acute attack. The urine was positive for ketone bodies. He was stuporous on admission but responded to glucose administration and after two days stopped vomiting. His weight and height were at the 25th centile, and his psychomotor development was normal. Investigations for hypoglycaemia were negative, and he was discharged home with the presumptive diagnosis of ketotic hypoglycaemia of infancy.

Over the subsequent years he was admitted to hospital every four to five months with similar episodes that clinically resembled Reye’s syndrome. On no occasion were ketone bodies found in the urine, despite the hypoglycaemia. His development proceeded normally and at that time muscle weakness was not noted. At the age of 10 years he experienced only one episode during the year, but became a vegetarian and refused to eat meat. Two months later he had a further episode of vomiting, lethargy, liver enlargement, and hypoglycaemia, and from that time his condition deteriorated. The frequency of attacks increased to one every three weeks and he started to complain of muscle weakness.

In October 1982 he was again admitted to hospital, and when his condition improved, plasma carnitine concentrations were measured and found to be low. Total carnitine was 12 nmol/ml (normal values mean SD, 65 (15) nmol/ml total carnitine and 40 (10) nmol/ml free carnitine). At the beginning of November 1982 he was admitted with his worst attack so far. Vomiting persisted for a week and the muscle weakness progressed to the extent that he could not raise his head. Diplopia and dysphagia subsequently developed. The hepatomegaly became more pronounced and liver enzyme values rose sharply.

Total parenteral nutrition was begun and L-carnitine (100 mg/kg/day) was added. The boy’s condition improved dramatically over the course of a few days. He was placed on a high carbohydrate diet with mid-chain triglycerides, gained weight, and the muscle weakness improved gradually. While on treatment the total carnitine plasma concentration was 73 nmol/ml with 69 nmol/ml free carnitine. A 12 hour fasting blood glucose value was still 60 mg/dl (3-3 mmol/l), but blood ketone-body concentration was lower than an age matched control after a 12 hour fast. His acetoacetate and β-OH-butyrates values were 0-349 mmol/ml and 0-256 mmol/ml respectively, while the control had corresponding values of 0-704 mmol/ml and 0-613 mmol/ml. The fasting test was not prolonged beyond 12 hours because of a recent report of sudden death after fasting in a patient with carnitine deficiency.4

The child has now been on carnitine treatment for 10 months and has not suffered any attacks; he has gained 7 kg in weight and feels very well. Muscle strength has now returned to normal. It should be noted that he still adheres to the meat-free diet.

Discussion

Since the first report of systemic carnitine deficiency in 1976,5 more than 30 cases have been reported. Deficiency of carnitine blocks the mitochondrial oxidation of fatty acids to carbon dioxide in all tissues and to ketone bodies in the liver. Thus, patients who lack carnitine will become hypoglycaemic after a short period of starvation. Despite constant low carnitine concentrations, acute attacks of weakness and encephalopathy occur only intermittently. Whatever the exact exacerbating mechanism, it seems that most acute episodes are associated with fasting or semi-starvation. It is conceivable that mobilisation and uptake of free fatty acids in the face of a limited capacity for oxidation somehow have an initiating role.2

Exogenous carnitine is not usually needed for maintaining normal carnitine concentrations, as biosynthesis from lysine takes place. Carnitine is synthesised in the hepatocytes and released into the plasma for transfer to peripheral tissues.6 In human beings, carnitine may also be derived from the diet, and beer is a primary source. Vegetables contain a negligible amount of carnitine, and thus vegetarians have only endogenous carnitine. Newborns and infants are incapable of synthesising the necessary amount of carnitine and without exogenous carnitine, derived from milk products, symptoms of carnitine deficiency can occur.3

We believe that in our patient exogenous carnitine played an important role. Notwithstanding the impaired endogenous carnitine production, our patient, while on a regular diet, had sufficient carnitine to prevent recurring episodes of encephalopathy and the other signs of systemic carnitine deficiency. With the change to a vegetarian diet without meat and milk products, the amount of exogenous carnitine ingested was negligible, and frequent attacks occurring every two to three weeks resulted. The introduction of L-carnitine (100 mg/kg/day) increased the blood carnitine concentration to normal and despite the cohesiveness of the vegetarian diet, no further attacks have occurred over the past 10 months.

References

Fifty five years ago

A syndrome in the rat resembling pink disease in man

Ruby O Stern (National Hospital, Queen Square, London)

‘Investigation of the pathological changes occurring in young rats fed on dried egg-white shows that very characteristic changes occur in the central nervous system. These pathological changes still further emphasize the similarities of the condition in the rat to Swift’s disease in man. These similarities may be thus briefly summarised:—

(i). Both diseases occur in young animals. Swift’s disease has never been recorded in adults, while in the adult rat it is only after from three to five months that cutaneous lesions can be produced.
(ii). The diseases may occur on a diet of mothers’ milk or a ration containing ample supplies of the known vitamins.
(iii). The clinical symptoms of both diseases are nutritional, nervous and cutaneous.
(iv). In rats there is a characteristic ‘kangaroo’ position, in children a knee-elbow position.
(v). In rats and children there is a curious mousy odour.
(vi). Death is often due to an intercurrent broncho-pneumonia.
(vii). There is no evidence that the diseases are due to an infection with bacteria or ultramicroscopic viruses.
(viii). The pathological changes in the skin and nervous system are similar.’

Archives of Disease in Childhood 1929;4:1-11.
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