Leber’s hereditary optic atrophy: further evidence for a defect of cyanide metabolism?

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SUMMARY We studied one patient with Leber’s optic atrophy (LOA) in the acute stage and 12 at the chronic stage of the disease, and measured the activity of rhodanese in white blood cells and the level of cyanide in whole blood. In the patient with acute disease the blood cyanide level was significantly increased at first. Treatment of this patient with cyanide antagonists reduced his cyanide level, but this was not accompanied by improvement in visual function. Rhodanese activity was normal in all patients, as were the blood cyanide levels in each of the 12 patients at the chronic stage of the disease. These findings suggest a temporary disturbance of cyanide metabolism during the acute phase of the disease in this family. The abnormal metabolic mechanism was not identified.

In 1965 Wilson1 observed that in most cases of Leber’s optic atrophy (LOA) the severity of the disease could be related to tobacco smoking. Because tobacco is the major exogenous source of cyanide in Europe, he suggested cyanide as a possible precipitant of visual loss in LOA. This impression was reinforced by the observation that optic neuritis occurs as a result of cyanide intoxication from ingestion of bitter cassava in man2 and in animals.3 These findings prompted a search for abnormal cyanide metabolism in LOA.

Wilson1 observed significantly increased urinary thiocyanate excretion in healthy smokers as compared with that in non-smokers. By contrast no difference between smokers and non-smokers was found in LOA patients, suggesting that patients with LOA are unable to respond to an increased intake of cyanide by increased detoxification. Rogers4 also observed relatively low levels of thiocyanate in patients with LOA.

Significantly reduced rhodanese (Ec: 2.8.1.1) activity, the key enzyme for cyanide detoxification, was found by Cagnianut et al.5 in the liver parenchymal cells and by Poole and Kind6 in the rectal mucosa of patients with LOA. However, Wilson1 reported normal rhodanese levels in the liver, Syme et al.7 in the red cells, and Nikoskelainen et al.8 in mitochondria of muscle samples of patients with Leber’s disease.

These observations prompted this further study of cyanide metabolism in patients with LOA.

Case report

In mid-January 1985 a 22-year-old male non-smoker noticed decreased vision in the left eye followed by a decrease in the right eye two weeks later. The visual acuity was 20/200 in the right eye and 20/300 in the left eye. The right optic disc of the patient was hyperaemic, with blurred margins. Circumpapillary telangiectatic microangiopathy, tortuosity of the retinal vessels, and swelling of the retinal nerve fibres were apparent. No abnormal vascular permeability was demonstrable by fluorescein angiography. In the left eye the optic disc was blurred only nasally.

At that time general examination (including urine tests), neurological examination, CT scan, and the flash electroretinogram (ERG) were normal. The mother and the grandmother of the patient had also suffered acute visual loss in the 4th decade of life.
and both were more severely affected than the propositus. Cyanide levels were determined according to the method described by Pranitis and Stolman, as modified by Wicht. Rhodanese (a mitochondrial enzyme) activity was measured in white blood cells by Sorbo's method.

A significantly increased blood cyanide level of 1-4 µg/ml (54 µmol/l) was found in the patient at the first examination (normal level <0-2 µg/ml (7-7 µmol/l)). In an attempt to lower the blood cyanide level the patient was treated for two days with 250 mg p-DMAP (which forms met-Hb), 600 mg CO₂-EDTA, 18 g vitamin B₁₂ (which both complex cyanide), and 375 ml of 8% sodium thiosulphate infusion (substrate for rhodanese). With this treatment the blood cyanide fell to 0-6 µg/ml (23 µmol/l). Because of cardiac side effects CO₂-EDTA was withdrawn. Treatment was continued with sodium thiosulphate, and the cyanide level returned to normal after four weeks (Fig. 1). Treatment was then stopped and the blood cyanide level rose to 0-8 µg/ml (30-8 µmol/l). The visual acuity with each eye gradually deteriorated to 20/600 over the next three months. Treatment with sodium thiosulphate was begun again in July 1985, and cyanide levels returned to normal values. The level remained normal after cessation of treatment (Fig. 1). By September 1985 the visual acuity with each eye remained at 20/600, and the optic discs were pale.

An additional 12 patients with chronic LOA (including the mother of the patient reported on above) were tested. White blood cell rhodanese activities and blood cyanide levels of all these patients were within the normal range.

**Discussion**

Our observation that the cyanide level was increased in a patient with acute LOA lends further support to the suggestion of Wilson that there is a defect in cyanide metabolism. We could not induce any improvement in visual acuity despite adequate reduction of the blood cyanide levels. The late onset of the treatment (two weeks after the second eye deteriorated) may explain the absence of visual recovery. It is remarkable that an increased cyanide level was observed only in the acute stage. In each of our chronic cases cyanide levels were normal, and in our patient with the acute disease it returned to normal after the attack. In contrast to the findings of Cagnianut et al. and Poole and Kind, none of our patients with LOA showed any reduction in rhodanese activity.

The finding in the patient with the acute disease implies a self-limiting derangement of cyanide metabolism. This would be in accordance with the behaviour of the disease in which acute dysfunction may or may not occur in patients with the abnormal inherited factor and funduscopic abnormalities. These observations also imply that a metabolic abnormality may be detected only during the acute stage, and that normal activity of enzymes involved in cyanide metabolism outside the acute stage of the disease would not necessarily exclude abnormal enzyme activity as contributing towards the pathogenesis of the disease.

We thank Professor O-E Lund for his encouragement and support. We are further deeply indebted to Professor A C Bird, Dr C Vesey, and Professor G W Weinstein for their help on the manuscript and
TA Berninger, L v Meyer, E Siess, O Schon, and F-D Goebel

Professor E Aulhorn, Professor A Blankenagel, Dr H Berdjis, and Professor K W Ruprecht for their support.

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Accepted for publication 9 June 1988.
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doi: 10.1136/bjo.73.4.314

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