Axonal sensorimotor neuropathy in patients with β-thalassaemia

E Stamboulis, N Vlachou, M Drossou-Servou, P Tsaftaridis, G Koutsis, N Katsaros, E Economou-Petersen, A Loutradi-Anagnostou

The purpose of this study was to investigate the prevalence of peripheral neuropathy in patients with β-thalassaemia.

Methods: Thirty-six patients with a mean age of 29.2 ± 8.2 years and 17 healthy controls with a mean age of 27.6 ± 9.1 years were included in this study. Measurements included the neuropathy symptoms score (NSS), the neuropathy disability score (NDS) as well as nerve conduction studies of two motor (ulnar and peroneal) and two sensory (ulnar and sural) nerves of the right limbs.

Results: A mainly sensory axonal polyneuropathy was present in 19 out of 36 patients (52.7%). Eight out of these 19 patients also had abnormal NDS values. The neuropathy correlated significantly with the age of the patients and the hematocrit. However, it did not correlate with the presence of antibodies against HCV, the ferritin levels, or with a history of transfusions, desferrioxamine treatment, or splenectomy.

Conclusions: This study showed a high prevalence of a predominantly sensory neuropathy in patients with β-thalassaemia. The electrophysiological data suggest that the underlying pathology is an axonopathy. Chronic hypoxia of the nerves resulting from severe anaemia may contribute to the pathogenesis of this neuropathy.

RESULTS

Nine out of 36 patients reported symptoms that could be attributed to neuropathy such as numbness, tingling, hyperpathia, cramps, muscular weakness, muscle tenderness, and restless legs. In eight patients neurological signs of neuropathy were demonstrated, such as reduction in vibration sense, hypoaesthesia of the feet, weakness of the extensor digitorum brevis, and reduced knee and achillaeal reflexes.

A total of 19 patients (52.7%) had pathological values in the nerve conduction studies (table 1). The most frequent abnormal finding was a reduction in the amplitude of the sensory action potential (SAP) of the sural nerve followed by a reduction in the amplitude of compound muscle action potential (CMAP) of the peroneal nerve. Further statistical analysis revealed a significant reduction in the amplitude of the SAP of the sural nerve (patients 17.25 ± 7.81, controls 24.40 ± 6.10, p<0.01), the amplitude of the CMAP (patients 7.81, controls 9.10, p<0.05).

Abbreviations: CMAP, compound muscle action potential; DFO, desferrioxamine; Ht, haematocrit; NSS, neuropathy disability score; NDS, neuropathy symptoms score; SAP, sensory action potential; SD, standard deviation.
Table 1  Patients with pathological nerve conduction studies

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<th>Patient</th>
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abn., number of patients with abnormal results; ampl, amplitude (in mV for CMAP and in mV for SAP); lat, latency (in ms); NSS, neuropathy symptoms score; NDS, neuropathy disability score; veloc, velocity (in m/s).
9.79±2.53, controls 7.92±1.88, p<0.01) and the motor velocity of the ulnar nerve (patients 59.29±3.61, controls 61.72±3.59, p<0.05), a prolonged distal motor latency of the ulnar nerve (patients 2.50±0.36, controls 2.15±0.33, p<0.01). A reduction in the amplitude of the SAP of the same nerve showed a tendency toward statistical significance (patients 36.47±13.39, controls 47.37±14.43, p<0.1).

Results of the NCS were further correlated with other clinical parameters. Statistical analysis showed that abnormal findings were more frequent in patients more than 20 years old (21–30: 12/21 patients with abnormal NCS, p<0.05, 31–40: 4/8 with abnormal NCS, p<0.1, 41–58: 3/3 with abnormal NCS, p<0.01) compared with patients less than 20 years old (0/4 with abnormal NCS); in patients with Ht less than 20 (Ht<30: 15/24 with abnormal NCS, Ht>30: 4/12 with abnormal NCS, p<0.01); and in patients with abnormal NDS (abnormal NDS: 8/8 with abnormal NCS, normal NDS: 11/28 with abnormal NCS, p<0.01). On the other hand, patients with a history of transfusions, DFO administration, or splenectomy were equally likely to have abnormal electrophysiological results as those with no such history (transfusion history: 14/28 with abnormal NCS, no transfusion history: 5/8 with abnormal NCS, χ² = 0.37 not significant (ns); receiving DFO: 15/28 with abnormal NCS, not receiving DFO: 4/8 with abnormal NCS, χ² = 0.02 ns; splenectomy: 12/9 with abnormal NCS, no splenectomy: 7/17 with abnormal NCS, χ² = 1.76 ns). The clinical severity of the disease (intermediate or major) did not appear to correlate with the frequency of abnormal findings (homozygous beta-thalassaemia: 17/32 with abnormal NCS, intermedia: 2/4 with abnormal NCS, χ² = 0.01 ns). Furthermore, the presence of antibodies against HCV or the presence of neuropathy symptoms was not associated with a significant increase in the frequency of pathological NCS (HCV+): 4/7 with abnormal NCS, HCV−: 15/29 with abnormal NCS, χ² = 0.05 ns; abnormal NSS: 5/9 with abnormal NCS, normal NSS: 14/27 with abnormal NCS, χ² = 0.03 ns). Patients with normal NCS had a mean ± SD value of ferritin of 2054.1±1876 and those with abnormal studies 2426.5±2355 (t-test = 0.52 ns).

DISCUSSION

The NCS in the present report demonstrated that more than half of the patients with beta-thalassaemia (52.7%) have evidence of a polyneuropathy. In eight out of the 19 patients with electrophysiologically established polyneuropathy, clinical signs of neuropathy were also present. The findings, as portrayed in table 1, suggest that the neuropathy is axonal and primarily sensory. The neuropathy first affected the lower limbs since pathological values were never limited to the upper limbs of any patient.

It has been reported in previous studies that the prevalence of polyneuropathy in patients with beta-thalassaemia is much lower than our results seem to suggest, ranging from 21% to 25%. This difference could be attributed to the fact that the patients in these studies were in the second decade of their life, whereas our patients were mostly in the third and fourth decades. The significant increase in neuropathy with age observed in our study has also been demonstrated in a similar Greek study in the past. In common with Papanastasiou et al, we found a significant increase in the frequency of neuropathy in patients with a low Ht and a lack of correlation with ferritin levels, administration of DFO, splenectomy, and transfusions. In addition, we found a lack of correlation of neuropathy with presence of antibodies against HCV. Our finding that severity of disease (homozygous vs intermedia) does not seem to correlate with the frequency of abnormal findings is probably due to the fact that the average age of patients with thalassaemia intermedia was greater than the average age of the rest of our patients (37.0±12.7 and 27.8±6.5, respectively).

A further interesting finding was that electrophysiologically established neuropathy correlated statistically with the presence of clinical signs of neuropathy, but not with the presence of clinical symptoms. It is possible that nerve hypoxia may lead to symptoms such as numbness, tingling, or cramps without causing any neural tissue damage.

Levine et al suggested that DFO can cause sensorimotor neurotoxicity in high doses. They reported two such cases. One patient had asymmetrical neuropathy of the upper limbs with electrophysiological examination showing a radiculopathy and the second patient who was diabetic developed a polyneuropathy. In both cases the neuropathy might not have been due to the administration of DFO.

Wong et al also attributed the subclinical neuropathy of their patients (seven out of 34 had pathological sensory conduction velocity of whom three had diabetes) to the long use of DFO but a statistically significant correlation with the dosage of DFO or the duration of its administration was not found. These findings cannot support the view that neuropathy in patients with beta-thalassaemia is due to toxicity of DFO.

The role of chronic ischaemia in the development of neuropathy has been demonstrated both in experimental studies and in clinical situations where ischaemia or hypoxia is present. The nerve ischaemia usually results in axonal damage. The large sensory myelinated fibres seem to be more severely affected. A high frequency of axonal neuropathy has been found in patients with chronic obstructive pulmonary disease, significantly higher in hypoxic groups. A mainly sensory axonal neuropathy probably due to the endoneural ischaemia, has been found in polycythaemia rubra vera, peripheral arterial disease and hereditary disorders of haemostasis. In accordance with the above, the predominantly sensory axonal polyneuropathy found in our patients could also be attributed to chronic hypoxia of the nerves resulting from longstanding anaemia. The higher prevalence of neuropathy in older patients and in patients with a low Ht is in agreement with this hypothesis.

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Competing interests: none declared

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Received 31 July 2003
In revised form 3 December 2003
Accepted 5 December 2003

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J Neurol Neurosurg Psychiatry 2004 75: 1483-1486
doi: 10.1136/jnnp.2003.024794

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