Short report

Meige syndrome: double-blind crossover study of sodium valproate

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SUMMARY A double-blind crossover study of sodium valproate and placebo was conducted in five patients with Meige syndrome. CSF neurotransmitter studies were performed at the end of each treatment period. GABA levels were not influenced by the administration of sodium valproate. An increase in HVA levels was observed in every patient, which may reflect an increase in central dopaminergic activity. This finding may explain the trend towards clinical deterioration which was observed during treatment with sodium valproate. Sodium valproate appears to be ineffective in Meige syndrome.

Most pharmacological studies dealing with Meige syndrome (idiopathic or essential blepharospasm and oromandibular dystonia) have been based on the assumption of disruption of reciprocal dopaminergic-acetylcholinergic influences in the basal ganglia. Treatment based on this assumption has yielded inconsistent results. Decreased CSF gamma-amino-butyric acid (GABA) levels in adult onset dystonia have been reported, suggesting central GABA deficiency in this group of disorders. Several uncontrolled studies employing GABA mimetic or facilitating agents in Meige syndrome with a favourable effect have been reported. Sodium valproate is a GABA mimetic and an increase in CSF GABA levels in humans during administration of sodium valproate in therapeutic doses has been reported. We conducted a double-blind crossover study in five patients with Meige syndrome, in order to test the following hypothesis: sodium valproate has a beneficial effect in Meige syndrome by restoring decreased central GABA activity, which is demonstrable by an increase in CSF GABA levels during treatment.

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Methods

Five patients with Meige syndrome gave informed consent for the trial. Individual characteristics are shown in the table. In accordance with Meige's original description, only patients with both blepharospasm and oromandibular dystonia were included; in all patients blepharospasm, interfering with normal vision, was the major symptom. Treatment periods with sodium valproate and placebo lasted 6 weeks and the sequence in which the drugs were administered was randomised. All other medication known to exert an influence on the central nervous system was stopped at least 2 weeks prior to the study. Sodium valproate was given three times daily in increasing doses up to 20 mg/kg at the end of the first week; an identical scheme was used for the placebo tablets. After 6 weeks the medication was discontinued and after a drug free interval of 2 weeks duration the next phase started with the same build up of medication. The patients were seen at 2-week intervals for evaluation and for medication adjustment by an independent clinician. Lumbar puncture was performed under standard conditions at the end of both treatment phases; on the same day, sodium valproate serum levels were monitored at 8 hour intervals.

Patients were admitted at least 24 hours prior to the lumbar punctures and started on a vanillyl mandelic acid (VMA) restricted diet. After a fasting and bedrest period of 12 hours lumbar puncture was carried out between 8 and 10am with the patient in sitting position. CSF was acidified with 1 drop formic acid per ml and frozen at -80°C in small portions. The concentrations of 3-methoxy-4-hydroxyphenylglycol (MPHG), homovanillic acid (HVA), 5-hydroxy-indole-acetic acid (5-HIAA) and GABA were determined by high performance liquid chromatography (HPLC). MHPG,
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Table  Patient characteristics

<table>
<thead>
<tr>
<th>Patient</th>
<th>Sex</th>
<th>Age at onset (yr)</th>
<th>Duration (yr)</th>
<th>Blepharospasm</th>
<th>Oromandibular dystonia</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>M</td>
<td>52</td>
<td>2</td>
<td>++</td>
<td>+</td>
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<tr>
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<td>F</td>
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<td>M</td>
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<td>3</td>
<td>+ + +</td>
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<tr>
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<td>F</td>
<td>52</td>
<td>11</td>
<td>+ + +</td>
<td>+ + +</td>
</tr>
<tr>
<td>5</td>
<td>M</td>
<td>58</td>
<td>3</td>
<td>+ + +</td>
<td>+</td>
</tr>
</tbody>
</table>

+++ = severe, ++ = moderate, + = mild.

HVA and 5-HIAA were estimated by separation on a Nucleosil C18 column (dim 3.0 × 100 mm) and electrochemical detection (EC) at a potential of 700 mV. The injection volume was 50 μl CSF and the elution buffer had a composition of 0.07 M trichloroacetic acid, 0.03 M sodium acetate, 10⁻⁴ M EDTA, pH = 4.0, and 5% methanol. Flowrate was 0.40 ml/min and pressure about 7000 kPa. GABA was measured by separation on an ion exchange column (Dionex DC-4A; dim 5.0 × 75 mm) and fluorometric detection using the method of Böhlen et al⁴ with minor modifications. CSF samples were deproteinised with 12.5% 5-sulpho-salicylic acid (4:1 v/v) and 100 μl of the supernatant was injected on the column. Elution was carried out with 0.067 M sodium citrate at pH 4.50 and detection after reaction with o-phthalaldialdehyde. Flowrate was 0.50 ml/min and pressure about 200–250 kPa.

Assessment of the effect of treatment was based on a quadruple scoring system: (1) subjective scoring of abnormal movements, (2) clinical scoring of abnormal movements, (3) blind rating of randomised surface EMG recordings and (4) blind rating of randomised video tapes. All patients rated the severity of their symptoms at a range score of 0 to 10; only the scores of the 2-weekly examination dates were taken into account. Dystonic movements were assessed by the senior author separately in upper and lower face and rated for severity: 0 = absent, 1 = mild, 2 = moderate, 3 = severe. Surface EMG recordings and videotapes were made on each examination with the patients in standardised postures (that is, at rest, reading aloud and attempting to suppress the abnormal movements, for 1 minute each). All EMG recordings were made using surface electrodes placed on frontal, orbicularis oris, orbicularis oculi, and platysma muscles. Recordings were randomised and rated by a neurophysiologist blinded to the names of the patients and the medications administered. Likewise, the randomised videotapes were independently scored by two raters.

For each evaluation variable, scores were ranked from best to most severe (6–1) for each patient. The summed ranks within the placebo and sodium valproate groups were then compared.

Results

Treatment with sodium valproate at therapeutic serum levels (mean: 79–3 μg/ml, range: 44–128 μg/ml) did not result in clinical improvement (fig 1). In fact, in all comparisons sodium valproate was slightly inferior to placebo. Mean CSF GABA levels were not decreased below reference values in the placebo phase and no increase was observed during treatment with sodium valproate (fig 2a). However, CSF HVA levels increased in all patients during treatment with sodium valproate (p = 0.03 on the assumption of no effect on HVA level) (fig 2b). No significant effect was observed on both MHPG and 5-HIAA levels, but mean baseline (placebo) MHPG levels were elevated above reference control values (fig 2c).

Discussion

The cause of Meige syndrome, a disabling condition, has yet to be established.⁵ Recent research suggests an increased excitability of the interneurons of the blink and corneal reflexes, probably due to a hitherto undetected basal ganglia disorder.⁶ Symptomatic blepharospasm has been observed in patients with brain stem lesions,⁷ communicating hydrocephalus,⁸ and bilateral basal ganglia infarction.⁹ Drug treatment of blepharospasm and/or oromandibular dystonia remains unsatisfactory,⁴,¹⁵,¹⁰,¹¹ although effectiveness has been claimed of such
different compounds as cholinergics,\(^2\) and anticholinergics,\(^2\) benzodiazepines,\(^6\) \(^7\) \(^8\) \(^23\) \(^24\) dopamine agonists,\(^2\) \(^4\) \(^6\) \(^7\) \(^24\) \(^25\) antagonists,\(^1\) \(^2\) \(^3\) \(^4\) \(^7\) \(^8\) \(^22\) \(^23\) \(^24\) \(^27\) baclofen,\(^8\) \(^27\) lithium,\(^24\) choline,\(^28\) and cannabinoid.\(^29\) Most studies have dealt with attempts to manipulate either the dopamine or acetylcholine systems of the brain, but there appears to be no consistent pharmacologic response.\(^2\) \(^4\)

Recently, the role of GABA in a variety of dystonic and hyperkinetic movement disorders has been stressed.\(^30\) \(^32\) Neophytides \textit{et al} reported decreased CSF GABA concentrations in adult-onset dystonia,\(^2\) of which Meige syndrome is regarded a manifestation.\(^32\) In one uncontrolled study,\(^7\) treatment of a patient with Meige syndrome with a combination of sodium valproate and baclofen resulted in "substantial improvement". Sodium valproate appears to be effective in reducing symptoms in tardive dyskinesia,\(^30\) \(^34\) a disorder which may be clinically indistinguishable from Meige syndrome.\(^1\) \(^35\) \(^36\) Sodium valproate may exert its influence on GABAergic mechanism via direct GABA transaminase inhibition, but, more likely, by an effect on a postsynaptic element, probably the GABA receptor complex.\(^10\) There is, however, some doubt whether doses used in the treatment of patients are large enough to affect brain GABA.\(^37\) CSF GABA levels have also been reported.\(^5\) We were unable to confirm the finding of decreased (placebo) GABA levels in our patients, and no increase was observed during treatment with sodium valproate. One of our reasons for investigating the efficacy of sodium valproate in Meige syndrome was the theory that this disorder is due to a reduction in central GABA mediated inhibition, resulting in striatal dopaminergic predominance, secondary to disinhibition of dopaminergic nigral neurons.\(^1\) \(^7\) Accordingly, treatment with a GABA mimetic or facilitating agent would be expected to result in clinical improvement. Recently, however, evidence has been presented for the opposite phenomenon:\(^38\) as the threshold dose of GABA receptor agonists for inhibiting striatal acetylcholine neurons is substantially lower than that for depressing dopamine cells, low doses of these agents may selectively reduce striatal cholinergic activity in the absence of changes in dopaminergic transmission, thus resulting in a net dopaminergic preponderance. We did observe an increase in CSF HVA levels during treatment with sodium valproate. This also suggests an increase in the dopaminergic activity in the brain during treatment with sodium valproate\(^11\) and may explain the trend towards clinical deterioration we observed in our patients.

In conclusion, there appear to be no reasons to investigate further the use of sodium valproate in the treatment of Meige syndrome.
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References

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