Myasthenic syndrome: effect of choline, plasmapheresis and tests for circulating factor

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SUMMARY In a patient with myasthenic syndrome neuro-muscular transmission was characterised by depression and facilitation. The relative extent of these two processes varied between muscles, and in the one muscle with time. Guanidine HCl treatment corrected the electrophysiological defect. Oral choline increased muscle action potential amplitude in response to single shocks. Intravenous choline produced features indicating cholinergic autonomic stimulation. Pimozide and plasmapheresis had no effect. Animal in-vivo and in-vitro studies performed to detect a circulating factor which interferes with neuro-muscular transmission were negative.

The myasthenic syndrome (MYS) is a well defined clinical entity. There remain a number of unresolved aspects, including a lack of knowledge of the cause and mechanism of the disorder. We have quantified certain parameters of neuro-muscular transmission in one patient with MYS in order to further define the pathophysiology of this condition. The transmission defect in MYS has two components, characterised by depression and facilitation. A variation in magnitude of these components with time, and at different neuro-muscular junctions, is demonstrated. This could explain the apparent concurrent presence of transmission features of myasthenia gravis (MG) and MYS which have been reported in some patients.

The effect of certain pharmacological agents, in particular choline, was assessed, and the influence of plasmapheresis noted. The patient’s serum was tested for its effect on neuro-muscular transmission in animal in-vivo and in-vitro preparations.

Patients and methods

A 56 year old woman presented with several months history of proximal weakness. Thyroid function tests were in the toxic range: T3 7-5 nmol/l (1-0-2-7), T4 250 nmol/l (60-145), FTI 360 (50-140), T3 resin uptake 144% (80-110), 123I thyroid uptake 20% at four hours. She was treated with carbimazole 60 mg/day and euthyroid status was reached after six weeks. Weakness persisted and three months later electrophysiological studies showed the characteristic features of MYS. Serum electrolytes including calcium, magnesium, manganese and cobalt were normal. Anti-acetylcholine receptor antibody test was negative. Repeated screening for an occult malignancy was negative.

Neuromuscular transmission was studied by recording compound action potential amplitude (CAPa) with superficial electrodes in response to supramaximal nerve stimulation. This was done in thenar, extensor digitorum brevis (EDB) and quadriceps muscles. Change in CAPa following nerve stimulation or exercise was expressed and plotted as a percentage change [(100×test CAPa/control CAPa)—100%]. Care was taken to have similar temperature at test sites.

In-vivo tests Edrophonium HCl was administered intravenously in a divided dose of 2 mg and 8 mg three minutes apart, preceded by atropine 0-6 mg intravenously. Choline was given intravenously as choline bitartrate. Calculated as the equivalent dose of choline Cl it was infused in normal saline at a rate of 27 mg/min over 70 min for a total dose of 38 mg/kg. Choline Cl was also given orally in a dose of 210 mg/kg/day for four weeks. Pimozide was given orally in a
dose of 6 mg/day for three weeks. Guanidine HCl was taken in a dose of 20 mg/kg/day orally over five months.

Continuous-flow plasma exchange was performed (Model 30 Haemonetics) with a total of six exchanges spaced 2, 5, 7, 9, and seven days apart. Mean exchange volume was 2520 ml, resulting in a drop in serum Ig A, G and M of 60%.

The effect of the patient's serum on the sciatic nerve-tibialis anterior muscle preparation of the cat was assessed. Twitch responses produced by supramaximal stimulation of the sciatic nerve were monitored. This was done under general anaesthesia with and without the addition of curare 50 mg/kg intravenously. The serum was injected intravenously or close intrarterially in single doses ranging from 0.1–1 ml.

**In-vitro tests** (I) Isolated rat phrenic nerve-diaphragm preparation. The effect of adding serum and plasma obtained at plasmopheresis was tested. In one series of experiments twitch tension elicited by supramaximal stimulation of the phrenic nerve at 0–1 Hz was monitored. In another series twitches were induced by direct muscle stimulation. (II) Isolated innervated chick biventer cervicis muscle preparation. The effect of serum on twitch responses elicited by supramaximal nerve stimulation was tested.

The preparations were suspended in tissue organ baths containing 25 ml of Krebs-Hanseleit solution for the diaphragm or McEwen's solution for the biventer cervicis muscle. The solutions were aerated with 5% CO₂ in oxygen and maintained at 37°C for the diaphragm or 39°C for the biventer cervicis muscle preparation. The serum was added in single doses of 0.1, 0.2, 0.4 ml with washout between, or as the same dose in cumulative mode with no washout. In either case each dose was left in the organ bath for five minutes. In the biventer cervicis preparation the test series was repeated after addition of curare 2 μg/ml to the solution to produce partial neuro-muscular block. Control preparations were run in parallel.

**Results**

Most of the results presented are for thenar muscles. The CAPa to single shocks measured at intervals over several months was low: 2.6±0.6 mV (mean±SD), n=15. The values ranged from 2.1–3.5 mV, with our normal range being 4.1–21.1 mV.

Stimulation at 1 Hz resulted in an exponential decay in CAPa (fig 1). At faster rates of stimulation up to 10 Hz, the response was characterised by depression followed by facilitation. The extent of depression and facilitation, and the transition point fluctuated in one muscle with time (fig 2A). It also varied between muscles tested on the same occasion (fig 2B). The dual characteristic of transmission was also defined by using twin stimuli (fig 3).

Stimulation at 20 Hz resulted in facilitation only. Faster rates of stimulation produced no additional potentiation. Fatigue of neuromuscular transmission was tested by prolonged stimulation at 20 Hz. The CAPa reached a plateau after 10 s, and was then maintained for 30 s, dropping thereafter 8% every 10 s. Normal subjects showed no change in CAPa over a 45 s period of nerve stimulation at this rate. Facilitation was also observed under physiological circumstances, such as maximal voluntary contraction. The potentiation and its time rate of decay are shown in fig 4.

Guanidine HCl treatment resulted in a three-fold increase in CAPa to 7.8±1.1 mV (mean±SD), range 6.7–9.2 mV. In association with this increase in CAPa, facilitation was no longer evident following exercise or during 20 Hz stimulation. An unexpected feature was noted after the first three days of guanidine treatment. The electrophysiological features of MYS were corrected in thenar muscles, but were still evident in the extensor digitorum brevis muscle (fig 5). After a further four days of treatment the defect was not apparent in either muscle group. The other drugs were administered when the patient was not receiving guanidine.

Edrophonium HCl infusion resulted in a 30% increase in CAPa. After one week of oral choline administration CAPa was 2.3 mV, and after a
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Further 2 1/2 weeks reached 4.5 mV. Choline was discontinued and three weeks later CAPa was 3.2 mV (fig 6A). 20 Hz stimulation at the corresponding times resulted in potentiation which was similar for the first second, and then diverged (fig 6B). Intravenous administration of choline produced no change in CAPa. The infusion was terminated because the patient complained of lightheadedness, and exhibited flushing of the cheeks, tremor and marked sweating. These features subsided within minutes of cessation of the infusion. For the next 24 hours, the patient experienced some abdominal cramps and diarrhoea. There was no subjective or objective change in strength as a result of guanidine or choline administration.

Pimozide produced no change in CAPa. Plasmapheresis did not result in clinical or electro-

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Fig 2 Change in CAPa. A1 and A2—for the thenar muscle at different times. B1—extensor digitorum brevis, B2—thenar muscle results measured on the same occasion.

Fig 3 Twin stimuli at 10 ms—10 s intervals. Amplitude of the second response relative to the first is plotted. Lines are fitted by eye.
physiological improvement over a five month period of observation. The cat preparation and the in-vitro tests showed no change in response as a result of adding the patient’s serum.

Fig 4  A and B: bar represents 15 s period of maximal voluntary contraction of thenar muscles. A—post-exercise potentiation and its decay, on four occasions, two being on the same day. B—results of A plotted on logarithmic scale. Circles and squares values obtained on same day, other values at different times. Time constant for faster decay was 23 s, for slower 51 s. Lines fitted by method of least squares.

Fig 5  Results after 3 days treatment with guanidine. 1—Thenar muscle results normal, 2—extensor digitorum brevis still shows features of MYS.

Fig 6  A—CAPa at different times in relation to treatment with oral choline Cl. B—change in CAPa following 20 Hz stimulation on the same occasions as readings in A were obtained.

Discussion

MYS is considered to be a pre-junction disorder, due to defect of transmitter release. The transmission defect has two characteristics—depression and facilitation, depending upon the rate of stimulation. Transmission characteristics fluctuate with time, and can vary with the muscle group examined. The selective effect of guanidine early in the course of treatment further points to the different severity of involvement of muscles in MYS. In-vitro studies have shown that the potentiating effect of guanidine is inversely related to initial quantal content at the neuro-muscular junction.

These findings may be relevant to the reports of patients who apparently have two neuromuscular junction disorders, MG and MYS, concurrently. These reports take two forms. The patient may exhibit electrophysiological features of MG and MYS in different muscle groups at the one time. Alternatively, the electrophysiological features may fluctuate between those of MG and MYS with time. These reports were made before estimation of anti-acetylcholine receptor antibody was available. There was also no information on force developed in the affected muscles. This may have been useful as the generation of force is different in MG and MYS. Our patient’s force records were typical of those
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seen in MYS (unpublished observations). An alternative explanation to the twin diagnosis of MG and MYS, may be that the observations represent fluctuation in the dual transmission characteristics of MYS.

The application of quantitative analysis to the electrophysiological changes in MYS has two aims. It defines in more rigorous terms the transmission characteristics, and the way these fluctuate in time. It could be used to compare the nature and extent of the defect between patients. The other potential benefit is that an analysis of the transmission defect may provide a clue to its mechanism. This relates to physiological studies aimed at determining the processes which act to increase transmitter release during and following repetitive stimulation. These have been separated to an extent by their decay time constant.\(^{10}\) Of the three processes arbitrarily subdivided on this criterion, our results fit the process of potentiation which decays with a time constant of tens of seconds to minutes. The variability in the decay constant shown in fig 4, also documents the fluctuation in transmission characteristics which occurs with time. The cause of this variability is uncertain. One possibility is that some change in the factor responsible for the pathophysiology of MYS is occurring. Another potential mechanism is change in local temperature. Animal studies have documented the sensitivity of facilitation of transmitter release to temperature, with the decay time constant of facilitation being inversely related to temperature.\(^{11}\) In MYS it has been found that cooling improves transmitter release\(^{12}\) and similarly in MG transmission is better at lower temperatures.\(^{13}\)

The potentiating effect of edrophonium HCl has been noted previously.\(^{14}\) Pimozide was used because of recent evidence indicating the presence of dopamine receptors with inhibitory influence on acetylcholine release at motor nerve terminals.\(^{15}\) Pimozide reversed this inhibition. The failure of pimozide to influence the electrophysiological features in this patient argues against the involvement of these receptors in the pathophysiology of MYS.

Choline has been used therapeutically in conditions such as tardive dyskinesia\(^{16}\) with the aim of increasing central acetylcholine (ACh) levels. Choline is a precursor of ACh, and its administration in animals has been shown to increase brain ACh concentration.\(^{17}\) Direct evidence of functional influence on ACh release is lacking in man, and only indirect evidence is available from animal studies.\(^{18}\) In MYS transmitter release at most neuro-muscular junctions is below the normal safety factor. A change in ACh release as a result of choline administration should therefore be observable. Our results suggest that prolonged oral choline administration caused at least a 40% increase in the release of an immediately available ACh pool. After 1s of 20 Hz stimulation, transmission tended to the same value as without treatment. The level of estimated increase in ACh release is in agreement with histochemical studies in animals showing raised ACh concentration after choline ingestion.\(^{17}\) The findings also suggest the presence of at least two ACh pools, only one of which is significantly influenced by choline intake. The existence of separate ACh pools in nerve terminals has been suggested by animal studies using labelled choline.\(^{19}\) It will be of interest to determine whether other patients with MYS show a similar response to oral choline.

Acute choline administration appeared to have no effect on transmitter release at the neuro-muscular junction. The clinical features exhibited by the patient were consistent with increased cholinergic autonomic activity. Transmission was therefore increased at some cholinergic sites. The mechanism underlying the difference in effect produced by acute and chronic choline administration, and in the acute setting the disparity in autonomic and neuro-muscular junction response remains uncertain. Animal studies indicate that the half-life of intravenously administered choline is less than one minute. The liver and kidneys remove 50%. Though conversion to ACh is rapid, it was concluded that this was a relatively minor pathway for choline metabolism.\(^{20}\) This coupled with the presence of specific high affinity transport systems for choline\(^{21}\) in parts of the nervous system suggests that in the acute setting available choline will be selectively distributed. The response to intravenous choline raises the possibility of acute manipulation of cholinergic function, in autonomic and possibly other parts of the nervous system.

Plasmapheresis was without effect. If the MYS was related to the presence of a circulating antibody, as in MG, then the schedule of plasmapheresis used should have produced a beneficial effect.\(^{22}\) If it was dependent on a circulating factor other than an antibody improvement may still have been expected. Alternatively the postulated factor, as the polypeptide botulism toxin, could be tightly bound to its site of action and produce long lasting effects. This is supported by the finding that prolonged washing of
biopsied inter-costal muscle preparation from
patients with MYS fails to alter the transmission
block.\textsuperscript{23} The lack of improvement in our patient
over a long follow-up period argues against this
possibility.

The animal and in-vitro studies showed no
evidence to suggest the presence of a circulating
factor which impairs neuromuscular transmis-
sion. Fractionation or concentration of the
serum was not performed, and it is possible that
the methods used were too insensitive or in some
other way inappropriate.

References

1 Lambert EH, Rooke ED. Myasthenic state and
lung cancer. In: Lord Brain, Norris FH, eds. The
remote effects of cancer on the nervous
system. New York: Grune and Stratton, 1965;

2 Moir M, Takamori M. Hyperthyroidism and
myasthenia gravis with features of Eaton-
Lambert Syndrome. Neurol (Minneap) 1976; 26:
882–7.

3 Schwartz MJ, Stalberg E. Myasthenia gravis
with features of the myasthenia syndrome.

4 Takamori M, Gutman L. Intermittent defect of
acetylcholine release in myasthenia gravis.

5 Brown GL. Close arterial injection in cat. J
Physiol 1938; 92:22.

6 Bulbring E. Isolated phrenic nerve diaphragm
preparation of the rat. Br J Pharmacol 1946; 1:
38–61.

7 Ginsborg BL, Warriner J. Isolated chick biventer
 cervicis nerve-muscle preparation. Br J Pharma-

8 Elmqvist D, Lambert EH. Detailed analysis of
neuromuscular transmission in a patient with
the myasthenic syndrome sometimes associated
with bronchogenic carcinoma. Mayo Clin Proc
1968; 43:689–713.

9 Kamenskaya MA, Elmqvist D, Thesleff S.
Guanidine and neuromuscular transmission.

10 Magleby KL, Zengel JE. Stimulation-induced
factors which affect augmentation and poten-
tiation of transmitter release of the neu-

11 Belnave RJ, Gage PW. Temperature sensitivity
of the time course of transmitter release. Brain

12 Ricker K, Hertel G, Stodieck S. The influence
of local cooling on neuromuscular transmission
in the myasthenic syndrome of Eaton and

13 Borenstein S, Desmedt JE. Treatment and
weather correlates of myasthenic fatigue. Lancet

14 Lambert EH. Defects of neuromuscular trans-
mission in syndromes other than myasthenia

15 Ganguly DK, Das M. Effects of oxotremorine
control by dietary choline. Science 1976; 191:
61–2.

18 Ulus IH, Wurtman RJ. Choline administration:
activation of tyrosine hydroxylase in dopa-
minergic neurons of rat brain. Science 1975; 194:
1060–1.

19 Aquilonius SM, Fleutge F, Schubert J, Sparf B,
Sundival A. Synthesis of acetylcholine in
different compartments of brain nerve terminals
in vivo as studied by the incorporation of choline
from plasma and the effect of pentobarbital on

20 Haubrich DR, Wang PGL, Wedeking PW. Dis-
tribution and metabolism of intravenously
administered chole (methyl-3H) and synthesis in
vivo of acetylcholine in various tissues of guinea

21 Kuhar MJ, Sathy VH, Roth RT, Aghaianian GK.
Choline: selective accumulation by central
cholinergic neurons. J Neurochem 1973; 20:
581–93.

22 Dau P, Lindstrom JM, Cassel CK, Denys EH,
Shev EE, Spitler LE. Plasmaphoresis and
immunosuppressive drug therapy in myasthenia

23 Lambert EH, Elmqvist D. Quantal components
of end-plate potentials in the myasthenic syn-
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H Kranz, D J Caddy, A M Williams and W Gay

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