Controversies about the treatment of myasthenia gravis

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SUMMARY Clinicians treating patients with myasthenia gravis must choose cholinergic drugs, corticosteroids, immunosuppressive drugs, thymectomy, or plasmapheresis. Clinicians must decide the sequence or combination of these therapies and when to deem lack of improvement a sign for a different therapeutic approach. Because controlled trials have not been done to evaluate therapies that may require months or years before benefit is evident, controversy abounds.

It is a privilege to participate in this Festschrift to honour Ian Simpson who has shown us the power of clinical thought in analysing a medical disorder as a biological problem. Among his many contributions, he has been widely applauded for his studies of myasthenia gravis, including an early analysis of thymectomy, his introduction of the concept of myasthenia as an autoimmune disease, and the critical notion that pharmacological deductions about the physiological abnormality in myasthenia presumed normal morphological geometry at the endplate. Simultaneously with Simpson's closely reasoned arguments, Nastuk and Strauss and their colleagues provided the first evidence of humoral abnormality and AG Engel and his colleagues provided evidence that the morphology of the endplate is not normal in human myasthenia.

It is my assignment in this symposium to review controversies about the therapy of myasthenia. Controversy is the unwanted child of ignorance, and we may be ignorant because we do not have the means to find the answers to questions, or because we have the means but cannot apply them. Too often in medicine, our ignorance seems to be due to asthenia of the will. We know how to find the answers, but we do not. We know that we could find out whether this or that proposed treatment is effective by setting up a controlled trial. But patients are not rabbits. Patients want to be treated and physicians have their own emotional problems; they want to treat. We adopt therapeutic programmes because they seem logical or useful, and these therapies become so embedded in custom that it is ultimately deemed unethical to even consider a controlled trial.

We are not alone in this dilemma. When I was a medical student, standard treatments included vagotomy for peptic ulcer and sympathectomy for hypertension; current controversies include radical mastectomy, tonsillectomy, carotid endarterectomy, and coronary by-pass surgery. Nevertheless, when physicians are entrusted with the care of patients with myasthenia gravis, they must choose between cholinergic drugs, steroids, more specific immunosuppressive drugs, plasmapheresis, and thymectomy. There is uncertainty about the value of all of these and there is difference of opinion about specific details of each therapy, or the sequence to be chosen. Lacking appropriate evidence, the physician must nevertheless make a choice.

That is the dilemma. Recognising that there are differences of opinion, I will try to make the questions clear, and to provide fair statements of divergent views. I have my own views, of course, but will try to separate assertion from fact.

CHolinergic DRUGS

Little attention is being paid to cholinergic drug therapy for at least two reasons: First, these drugs do not ordinarily restore normal life to patients with myasthenia. Second, the apparent clinical benefit of immunotherapies and evidence
of abnormal immune mechanisms in the disease have shifted therapeutic attention to measures that might alter the course of the disease, rather than merely reversing symptoms. Reviving archaic debates may not seem appropriate but almost all clinicians use anticholinesterase inhibitors to start treatment, and it may be useful to indicate that there are still unanswered questions, even about this primeval form of drug therapy. **Which anticholinesterase medication is most effective?** The immediate effects of cholinergic drugs are so dramatic that the therapeutic response can be considered part of the definition of the disease.8 There is no need for a controlled trial when ptosis disappears before your very eyes or ophthalmoplegia melts into normal motion. The therapeutic benefit of oral medication, however, is usually much less dramatic. Some disabled patients may be returned to normal activity, but symptoms are rarely relieved completely. Of the three drugs now available (neostigmine, pyridostigmine, ambenonium), pyridostigmine has proven the most popular, but this choice was never based on any kind of formal assessment. Rather, patients were asked which drug they preferred, and most chose pyridostigmine because it caused the fewest gastrointestinal effects. There has never been convincing evidence that one drug is better than another for specific symptoms, or that one drug can be effective when another is not.

*How is the optimal dosage of a cholinergic drug determined?* There is still no way to determine the optimal dosage of pyridostigmine therapy, except by a trial that depends upon subjective responses of patient and physician. Osserman8 introduced the edrophonium test to determine optimal dosage. A patient already taking oral medication is given 2-0 mg edrophonium intravenously. If symptoms improve, the patient is deemed undertreated and the oral dosage is increased. If symptoms worsen, overtreatment is presumed and the oral dosage is decreased. If edrophonium has no effect, oral treatment is deemed optimal.

However, the edrophonium test has never been assessed formally, to determine whether these conclusions are justified. It was never demonstrated why 2-0 mg should be the decisive dose, rather than 3-0 or 5-0 or some other quantity, or why it should be the same in every patient. In untreated patients the same dose of edrophonium may have a diagnostic effect one day, no effect the next, or may improve function of some muscles but worsen others; how then can the test be reliable in treated patients? We never assessed this test formally either, but we have not found it a reliable guide to oral therapy, and the test does not seem to be used very often nowadays.

If there is no objective way to determine optimal dosage, we have to rely upon statements of the patient about the response of specific symptoms. It is my impression that patients and physicians tend to increase the dosage progressively, so long as the patient is not too distressed by side-effects. Often, the dosage can be sharply reduced without ill effect. Although there is no direct evidence to support the statement, few patients seem to show objective benefit from doses larger than 120 mg every two or three hours. That is why “pyridostigmine requirement”, at least in terms of daily dosage, should not be used as a measure of therapeutic effect for thymectomy other measures designed to alter the course of myasthenia.

*How often does cholinergic crisis occur?* Years ago, when irreversible inhibitors of cholinesterase were evaluated in the treatment of myasthenia, it became apparent that overdosage could cause weakness and pyridine-aldoxine methi-dide (PAM) was introduced as a specific antidote for this kind of organophosphate intoxication. That cholinergic crisis may occur is indisputable, and it can be demonstrated experimentally,9 but it is not clear whether or how often it occurs in patients treated with the standard drugs now in use. Indeed, as crisis itself has become less frequent, the question has disappeared from the literature.

There are still, however, occasional references to the use of edrophonium as a method to determine whether weakness is “myasthenic” or “cholinergic.” As originally described,8 the test again required an arbitrary dosage of edrophonium (1-0 or 2-0 mg), but interpretation now depended on only two choices: if weakness improved, it was “myasthenic”; if there was no improvement, or if weakness worsened, the weakness was “cholinergic.” Again, however, this interpretation was never assessed formally, and it is not clear why “no response” should be deemed optimal in one situation but not in another. We have been sceptical about this interpretation because it has been standard practice in our institution for 30 years to discontinue cholinergic medication whenever a patient requires respiratory support. Never have we seen a patient improve immediately after discontinuing cholinergic medication; either there is no immediate change, or, more often, the weakness becomes perceptibly worse. If cholinergic crisis
occurred, patients should improve promptly when the drug is withdrawn. There has been no convincing documentation of acute cholinergic crisis in any patient taking only oral medication. **Should administration of cholinergic drugs be discontinued during crisis?** We define crisis as an exacerbation of myasthenic symptoms that requires mechanical ventilation. In 1949, Randt and Korey gave one patient 18-0 mg neostigmine intravenously in one hour; the equivalent oral dose would be 440 mg or almost 30 standard tablets of 15 mg each. Within 60 hours, that patient received 50 mg neostigmine intravenously and 95 mg intramuscularly.10 There were neither beneficial nor deleterious effects but they concluded that cholinergic drug therapy could not terminate crisis, and continued use could complicate management by causing diarrhoea or increasing pulmonary secretions. For these reasons, it has become standard practice in our institution and others to discontinue cholinergic drug therapy as soon as mechanical ventilation is started.

One practical effect of this approach is that the distinction between “cholinergic” and “myasthenic” crisis becomes inconsequential, since the possibly offending drug is withdrawn anyway. It does, however, leave the question of when to resume drug therapy, and this is important after thymectomy as after crisis of other cause. There are no clear answers to this question but we view crisis as a temporary exacerbation; it is our task to keep the patient alive until the cause of the exacerbation subsides, assuming that it is due to transient effects of respiratory infection, aspiration pneumonitis, or surgery. Therefore, pyridostigmine does not seem useful and we cannot “get the patient off the respirator with pyridostigmine” while the patient is febrile, or while other complications are overtly active. Once these complications subside, respiration begins to improve and pyridostigmine can be started again. **Could treatment be improved by monitoring blood levels of cholinergic drugs?** Little is known about the pharmacokinetics of pyridostigmine. After intravenous injection of 14C-labelled drug, 44-47% of the administered label appeared in the urine within one hour.11 Because excretion of the drug exceeded creatinine excretion, there seemed to be tubular excretion as well as glomerular filtration.12 13 Interactions with other drugs and hepatic metabolism have not been defined, and much of orally administered drug appears in the faeces.13

There have been few studies of blood levels since the introduction of gas-liquid chromatography. In one study,14 serum levels were lower in patients deemed “poorly controlled” than in those who seemed to be responding to therapy. Because parenteral administration of drug raised the low blood levels to the range of those in good control, the originally low drug levels were attributed to malabsorption. However, in another study,15 plasma pyridostigmine content was restricted to a narrow range (20-60 mg/ml) despite wide variations in daily dosage (60-660 mg) among different patients. In still another study, peak levels and area under the curve were higher in poorly controlled patients.16 Further work seems to be necessary to resolve these differences, and to indicate whether therapeutic drug monitoring has a role in myasthenia. **Does chronic administration of cholinergic drugs have deleterious effects?** From time to time, clinicians wonder whether patients may become “refractory” to the beneficial effects of cholinergic drugs, perhaps because of desensitisation of receptors or because of drug-induced changes in morphology. Experimentally, chronic administration of these drugs can induce both physiological and morphological changes.17-20 However, there is virtually no evidence that this is true in humans. The dosages used in the animals were much larger than patients take, and there is no clinical evidence of declining effect of cholinergic drugs that might be dependent upon dosage or duration of treatment (but that has not been assessed formally either). Mayer et al21 presented evidence that myasthenic crisis might be due to drug-induced changes, but questions were raised about their data.21 The possibly beneficial effects of a “drug holiday” in treatment of myasthenic crisis have not been evaluated.

**THYMECTOMY**

**Is thymectomy effective in the treatment of myasthenia?** The world over, thymectomy is standard therapy for myasthenia. There is no debate about that statement. Yet the beneficial effects of thymectomy were discovered by accident when Blalock22 removed a cystic thymoma from a young woman with myasthenia, and the myasthenic symptoms subsequently improved. On the basis of that experience, the operation was gradually adopted, in large part due to the influential work of John Simpson.1 But in those days shortly after World War II, no one knew the function of the thymus, so there was not much rationale for thymectomy. Even today, it is not known whether benefit is bestowed because a source of antibodies is removed,
because the source of antigen is removed, or for some other reason.

In the early days, there was controversy about the therapeutic value of thymectomy, and the considerable operative mortality was a restraint. Even at the Massachusetts General Hospital, six of the first ten thymectomies ended in death. But, Keynes soon pointed out that results were better for patients who lacked thymoma, and gradually improvements in surgery, anaesthesia, and respiratory care reduced the mortality rate to virtually nil. For instance, at the Columbia-Presbyterian Medical Center, there were four deaths in 21 operations between, 1942 and 1955, but there were no deaths in 33 operations between 1955 and 1966, and there have been no deaths in 101 trans-sternal thymectomies from 1963 to 1979. Other centres have experienced the same changes; as the risks declined, more operations were done.

Results of the modern operation have been summarised in many reports since 1970 and the generally held view can be stated succinctly. When there is no thymoma, thymectomy is followed by improvement in 66% to 86% of the patients. There has not really been much debate about this; McQuillen is a brave but solitary voice, questioning the nature of the Emperor Thymectomy's robes, and McQuillen's doubts were expressed by comparing only the incidence of complete remission in operated and unoperated cases. It may not be appropriate to disregard improvement short of complete remission, and some have questioned McQuillen's acceptance of possibly brief and early remissions in unoperated cases, in contrast to lasting remissions after thymectomy.

Whether McQuillen is absolutely correct or not, other students have been disquieted by the lack of any prospectively controlled study of thymectomy. Although several authors compared results in operated and unoperated cases, only one study was concerned with matched cases, comparing patients of the same age, sex, duration of symptoms, and severity of symptoms. Even in that study, operated and unoperated patients were not contemporary; operations were probably done more often in recent years, after improvements in respiratory care (and also after the introduction of steroid therapy).

Therefore, despite the almost universal belief in the efficacy of thymectomy, its value has not really been proven. Also, there have been some inexplicable cases, such as improvement of myasthenia after mediastinal exploration without thymectomy, or after removal of an echinococcus cyst of the lung. Some have even claimed that Blalock's original case was not really a thymectomy, because the tumor was entirely cystic. It is possible, though doubted by most, that improvement in respiratory care now permits patients to live after crisis that would have been fatal years ago, and that the natural history of the disease is then one of improvement. Simpson, Genkins et al., and others believe that the disease is "active" for a finite time, although there is no way to measure "activity" of the disease except by symptoms. These possible doubts come rather late in the game and it is now difficult to conceive of a prospectively controlled trial of thymectomy, but some investigators in the USA are making the attempt.

Is cervical thymectomy preferable to trans-sternal thymectomy? Barring the nagging but fundamental doubts of McQuillen, the major controversy about thymectomy concerns the operative technique. The standard operation involves splitting the sternum and although operative mortality has been essentially obliterated, it is still a major operation; respiratory assistance is often required in the postoperative period and patients may have to remain in an intensive care unit for several days. It was therefore of great interest, in 1967, when Kark and Papa- testas and their associates began to write about transcervical thymectomy. The morbidity of this procedure is much less than the trans-sternal operation and it spares the patients from a disfiguring scar on the chest. The morbidity is so slight that the indications can be extended to children or the elderly, or to patients with only slight functional disability, groups not ordinarily selected for the trans-sternal operation.

Since the results of the transcervical operation are claimed to be equivalent to the standard procedure, it is rather strange that the operation has not been adopted by more than one centre in the US and one in Europe. The major argument against the cervical approach has been stated by Jaretzki; it is not possible to do a total thymectomy through the neck in every patient; complete removal of the thymus requires thorough exploration of the mediastinum. Several other arguments have been adduced: (1) There has been no prospective comparison of the two operations and more patients with mild myasthenia (who might be expected to have a better prognosis anyhow) are selected for the transcervical operation; outcomes of mild and severe cases operated in different epochs cannot be compared. (2) The cervical operation...
was introduced in 1967 and insufficient numbers of patients have been followed long enough for evaluation. (3) Takahashi et al.\textsuperscript{66} reported remissions or improvement in 54\% of 13 patients four years after cervical thymectomy performed between 1969 and 1973. For reasons similar to Jaretzki\textsuperscript{56} they changed to a trans-sternal operation in 1973 and among 43 patients who had this “radical” operation, the rate of remission and improvement was 88\%-4\% at four years, and improvement was seen sooner. (4) Even the proponents of the cervical operation believe the sternum should be split to remove thymomas, but not all thymic tumors are detected before surgery by standard radiograms (and computed tomography of the mediastinum may be falsely “positive”).\textsuperscript{57} (5) If the sternal operation is “thorough”, it is known that all recognisable glandular tissue has been removed. If the patient does not improve after cervical thymectomy, however, physicians may wonder about residual thymus. For instance, Stump et al.\textsuperscript{58} reported that residual thymic tissue was found by a trans-sternal operation in all of five patients who had not improved after an earlier cervical thymectomy. Even the trans-sternal operation may leave tissue behind.\textsuperscript{28 59}

This dispute will only be resolved by the results. What is needed is some evaluation of truly comparable patients subjected to the two procedures and followed for a prolonged period of observation; this is not available. If the cervical operation ultimately proves to be beneficial, it would mean that “total” thymectomy is not necessary for therapeutic effect. This conclusion is not impossible, since no one knows what deleterious influence is removed by thymectomy, nor why improvement may be delayed as long as seven years; merely reducing the burden may suffice. Is it possible to predict which patients will benefit from thymectomy? Other controversies pale by comparison to the questions about thymectomy itself, or the surgical approach, but other differences of opinion should be noted. Originally, it seemed that young women were the best candidates for thymectomy, but this could have been an artefact of selection since young adult patients would be most likely to be selected for a major surgical procedure, and most young adult patients with myasthenia (and no thymoma) are women; some men and some older patients have improved in virtually every report. Short duration of symptoms has been deemed important, especially in recent years,\textsuperscript{29 33} but in the single largest series, improvement rates were the same for patients with symptoms for more or less than five years.\textsuperscript{44} The histology of the thymus has been considered important, but some authors regarded presence of germinal centres as favourable;\textsuperscript{66} others considered lack of germinal centres favourable,\textsuperscript{73 36 46 61} or found that thymic histology was not related to outcome.\textsuperscript{29 62 64} As McQuillen and Leone\textsuperscript{48} pointed out, these discrepancies may not be surprising because it has not been proven that germinal centres are actually abnormally numerous in glands of patients with myasthenia.\textsuperscript{65 66}

There has been concern about doing thymectomies in pre-adolescent children\textsuperscript{47}, but the rates of improvement seem no different from those in adults and no long-range harm has become evident, even when the operation was performed in children as young as two years.\textsuperscript{45 65-74}

It is probably fair to state that there are no reliable predictors of favourable outcome. The most common indication for thymectomy is disabling myasthenia, but “disabling” permits considerable room for subjective choice by the physician. As the operation becomes safer, and as more physicians become convinced of the value of the operation, more patients are selected for thymectomy and, increasingly, during the first year or two of symptoms. Can “early” thymectomy prevent “progression” of myasthenia? Although it is difficult to find evidence in the literature, it has been stated that early thymectomy prevents “progression” of mild myasthenia to a severe state.\textsuperscript{53} Because of this view, the Mount Sinai group has attempted to identify which patients with solely ocular myasthenia are destined to have generalised disease.\textsuperscript{73} If a patient with solely ocular symptoms shows either a decremental response to repetitive stimulation of a limb nerve or abnormal sensitivity to curare to a local challenge of limb muscles with this drug, the disease is considered “generalised” and some of these patients have been subjected to thymectomy. Bever et al.\textsuperscript{76} however, found no correlation between the results of curare test or repetitive stimulation and subsequent-course of patients with ocular myasthenia.

Does steroid therapy enhance the results of thymectomy? It is probably true that, in most centres, steroid therapy is started only after thymectomy for most patients, at some time when physicians worry that improvement is not evident soon enough. This time depends upon the severity of symptoms; the more severe the disease, the sooner steroid therapy will be instituted. However, Johns and his associates\textsuperscript{57} have urged treatment with steroids before thymectomy because
steroid-induced improvement may make the postoperative course less onerous. For other clinicians, this policy introduces new problems. The outcome of steroid therapy is itself uncertain in time and degree; if patients are chosen on grounds of disability, why operate on a patient in remission, steroid-induced or spontaneous? To avoid uncertainty about the appropriate timing of thymectomy after steroid therapy, why not do the thymectomy first and hope that this will avoid the risks of steroid therapy? These are some of the reasons why most centres do thymectomy first.

A new aspect of steroid therapy was introduced by Bolooki and Schwartzman78 who advocate administration of methylprednisone, given intravenously in divided doses just before, during and after the operation for a total of 1000 mg in 24 hours. They believe that the regimen greatly reduced the morbidity of the trans-sternal approach; all 32 patients so treated left the intensive care unit within 24 hours. There have been no controlled studies of this approach, nor any reports from other centres.

Why should thymomas be excised? There is no debate about the advisability of surgical excision of thymoma in patients with myasthenia; it is done routinely. However, when a thymoma is present, thymectomy is less likely to be followed by improvement than in patients without thymoma. The indication for surgery of thymoma is therefore said to be the possibility that the tumour may ultimately invade neighbouring structures. In fact, symptoms are only rarely due to invasion of pericardium, lung, or superior vena cava79 and the myastenia sometimes does improve. Possible improvement of myasthenia is the “secret indication” in these patients, as in those without tumour.

One of the mysteries of thymectomy is how it exerts its beneficial effects. This is illustrated by some patients who do not have symptoms of myasthenia until some time (often years) after a thymoma has been removed. At first, it seemed that some thymic tissue was probably left behind in these cases,80 but Namba and Grob81 provided evidence that there had been apparently complete thymectomy in several patients with this kind of delayed myasthenia. Does that mean thymic cells had seeded other lymphoid organs before the tumour and thymic gland were removed? In contrast, however, myasthenia and tumour may both reappear after initial remission82; some patients experience remissions even if the thymoma is not removed83; and sometimes there is no autopsy evidence of thymic tissue even in fatal myasthenia.10 84 There is still much to be learned about the relation of the thymus to myasthenia.85

STEROIDS

Is steroid therapy effective in myasthenia gravis? Even to raise this question seems almost sacrilegious, because prednisone is now a staple of current therapy. It was not always so; there was a lag of almost 20 years from the first use of ACTH to some kind of general acceptance. But, as with other forms of myasthenia therapy, “acceptance” is not to be equated with rigorous proof of efficacy, safety, or clear indications for use. A brief review of steroid therapy provides a commentary on the sociology of academic neurology, especially in the United States.

ACTH first became available for therapeutic trials in 1950; its effects in the treatment of rheumatoid arthritis led to the award of a Nobel Prize and to use in other diseases of unknown cause. At that time, myasthenia was not regarded as an autoimmune disease. The first neurologists to be entrusted with limited supplies were Houston Merritt and Harold Wolff. Merritt, with Gilbert Glaser, used the drug in many different disorders, including one patient with severe myasthenia gravis who had “a partial remission” but died a few weeks later.86 87 Wolff, on the other hand, concentrated on myasthenia, reporting some improvement in ten of 15 cases; the improvement, however, was preceded by exacerbation of weakness and one patient died on therapy.88 89 Therefore two of the first 16 patients treated with ACTH died. Attention was directed to the possibly adverse effects of ACTH even though severe myasthenia was indeed grave in those days; the reported benefits were overshadowed.

Negative views of the effects of corticosteroids were reinforced by two other early reports, one from the Mayo Clinic80 and the other from Johns Hopkins81; both papers were influential because the writers were such highly respected authorities on myasthenia. The Hopkins paper was especially powerful and the story was told in the title of the paper: a patient was being treated with cortisone for rheumatoid arthritis when symptoms of myasthenia first appeared. If a drug cannot prevent the first symptoms of a disease, how can it be effective in therapy?

As a result of these experiences, there were no reports of steroid therapy from any of the major myasthenia centres for more than a decade. There were still, however, occasional reports of benefit92 98 and this provided the background
for a peculiar episode in the early 60s. At that
time, several clinicians were concerned about the
lack of controlled trials to evaluate thymectomy
and they met to design a suitable protocol. The
question of appropriate controls, however,
seemed insurmountable; even then, the notion of
sham operation was theoretically desirable but
was deemed ethically impossible. Also, the co-
operation of individualistic centres was difficult
to achieve. To prepare for some kind of
cooperative trial of thymectomy, therefore, the
participants agreed to study the effects of ACTH
on ocular myasthenia.

For several reasons, this seemed to be an ideal
way to devise methods of cooperative inves-
tigation. Eye movements before and after therapy
could be photographed and measured, objectively
and quantitatively, without recourse to subjective
evaluation and difficult-to-define words such as
"improved". Evaluation could be completely
"blind"; photographs were sent to another city
and measurements were made without knowl-
dge of therapy, whether it was placebo or
ACTH, or even whether the photographs were
taken before, during, or after therapy. Finally,
because the patients had solely ocular symptoms
and because the duration of therapy was limited,
the patients were not being subject to any serious
risk.

So the trial was initiated. Cooperation among
the participating centres was excellent. The re-
sults were unequivocal. ACTH therapy, 580
units given in eight days, had no effect.94

The issue could have ended there. But the very
next year, at an international conference on
myasthenia, von Reis et al95 described and pre-
sented motion pictures to prove, dramatically,
how beneficial ACTH therapy could be in
patients with severe generalised myasthenia.
Within a year, Grob and Namba96 and Osserman
and Genkins97 confirmed these reports and
steroid therapy was established throughout the
world. The next logical step was taken when oral
therapy with prednisone replaced injections of
ACTH in 1971.98 The wave of enthusiasm sank
the cooperative trial, which disappeared from the
literature without leaving a ripple. How could it
have been so wrong? Perhaps because gener-
alised and ocular myasthenia are not the same.
Perhaps the dosage or duration of therapy was
inadequate, but similar dosages and duration
were said to be successful in generalised myas-
thenia. Perhaps the controlled trial was not
wrong, because a critical review of the literature
of prednisone indicates persisting uncertainties
despite uniform endorsement of this form of
therapy.

To evaluate the effects of prednisone therapy,
we have analysed the 15 reports (from 10
centres) since 1971.99-112 Of these, only one at-
ttempted a controlled study; Howard et al107 found
that seven of ten treated patients improved but
so did three of ten receiving placebo, and the
difference was not statistically significant. The
number of patients was small, and the patients
all had mild myasthenia (because, ethically, the
investigators did not believe they could withhold
standard therapy from patients with severe symp-
toms). Therefore, there has been no adequately
controlled study, and no controlled study of any
kind for patients with severe disease (who are
most likely to be treated with prednisone).

Fischer and Schwartzman102 described uni-
formly beneficial results in eight patients with
solely ocular symptoms. The other reports con-
cerned 216 patients with generalised myasthenia,
but, taking the most recent reports, only three of
them described series of more than 20 patients
each.101 109 110 The results of each report, bar one,
were fantastic.

Some reported improvement in every single
patient treated99-101 104 108 and the rate dropped
only slightly in later reports from three of these
centres, to 84%105 89%106 and 92%.110

The only dissent from these reports of almost
universal improvement came from Kornfeld
et al109 who found benefit before thymectomy in
only 17% of patients with thymoma and only
30% of those without thymoma. After thryme-
tomy, however, these authors found 100% improvement in patients with thymoma but still
only 50% of patients without thymoma. Since
negative therapeutic results are less likely to be
reported the experience in some other centres
may be in line with the results of Kornfeld et al,
but that, of course, cannot be proven until the
data are published.

There are other discrepancies in these published
reports. Transient exacerbation of weakness was
reported by some,99 101 104 107 110 denied by
others,98 100 102 103 and not mentioned by Kornfeld
et al.109 Signs of improvement were noted as early
as the first day or within one week100 102; within
two weeks, but later extended to one month106;
one month103; 40 days104; or 50 days.110

It is important to know the upper limit of time-
to-improvement because, if there is no improve-
ment, the clinician must know when it is
appropriate to discontinue therapy as a failure.
This question was not specifically addressed in
any of these reports but it would seem that two
or three months should be adequate. Time-to-
maximum-benefit is similarly important, also not specifically considered, but given as less than one month, 8 months, 15 months and even 55 months.

Almost all of these investigators first used a daily dosage of at least 50 mg (or 100 mg on alternate days). There is virtually no information about the long-term use of prednisone; how long full dosage should be given, when or to what levels dosage may be reduced; in how many patients drug-induced remission is permanent and prednisone therapy can be discontinued; or how often it must be continued because patients relapse when dosage is reduced. Sanders et al. routinely used prednisone before thymectomy so that the postoperative course will be more benign, but the data of Kornfeld et al. suggest that this policy may not be appropriate. It is not clear whether prednisone is more or less beneficial in children than in adults, because it has been used in only a few children, and similar questions may be raised about the elderly.

Is prednisone therapy safe? Do the benefits justify the risks? Some authors reported no complications or nothing more than the cosmetic effects of Cushing syndrome or glycosuria. It is not possible to ascertain the incidence of more serious complications that have included vertebral collapse; congestive heart failure; gastric haemorrhage; ruptured diverticulum; tracheal abscess; psychosis; cataracts; aseptic necrosis of bone; and sepsis.

Two deaths were attributed to steroid therapy. No one has described the signed release used to explain these potential side effects in attempts to obtain informed consent for chronic prednisone therapy.

Does steroid therapy have pharmacological effects at the neuromuscular junction? The rationale for use of steroid therapy depends upon immunological theories of the disease, although it is not known precisely how steroids alter the pathological immune state. Several investigators have evaluated the possibility that steroids might repair the physiological abnormality of myasthenia but the balance of evidence suggests that this cannot be the basis for the beneficial effect.

How can we summarise the present state of prednisone therapy? It is now standard and customary, but has not been subjected to controlled trial and the incidence of documented risk has not been ascertained. It is therefore uncertain whether it should be used at all in patients who are not seriously disabled, in children, or in the elderly. It is not clear whether it should be used before or after thymectomy. It is uncertain how long full dosage should be continued before deeming any trial a failure. It is not known how long full dosage should be continued when there is apparent benefit, or how often treatment can be terminated without adverse effect. The reports of almost universal benefit cannot be taken at face value because of a single published dissent and because, if the results were as good as described, there would be little interest in plasmapheresis or immunosuppressive drugs.

**IMMUNOSUPPRESSIVE DRUGS**

At the 1970 international meeting on myasthenia, I was asked to review experience with immunosuppressive drugs. There had been reports of immunosuppressive drug therapy in only 46 patients and 38 of them had improved, but 38 of these patients (including 32 who improved) had been reported from one centre by Mertens et al. By means of a poll of centres in the US, we could obtain information about 14 cases treated with immunosuppressive drugs, but without detailed data; improvement was reported in four of these cases and the drug was discontinued in three. In sum, at that time, 60 patients were known to have been treated and 36 had improved.

A decade later, Hertel et al. held my paper responsible for a chilling effect on the use of these drugs, but that view exaggerates any influence I might have (or anyone else, for that matter). What stopped research on the use of immunosuppressive drugs in the US was the litigious nature of American society. At the 1970 meeting, I described the tragic case of a woman with life-threatening myasthenia who had an exaggerated bone marrow response to 6-mercaptopurine, became infected, suffered a myasthenic crisis, and died. That case resulted in a law-suit, euphemistically called "professional liability" here; the patient died in 1963 and legal action continued until the case was settled in 1973. The case was widely known among American investigators and probably did more to inhibit the use of these drugs than anything else; no physician wants to be accused of malpractice. For that reason, now, as in 1970, we have to look to European experience to evaluate immunosuppressive drug therapy.

Since 1971, there have been reports of only four series of patients who were treated with azathioprine without simultaneous prednisone therapy. Matell et al. found that 78% of 26 patients improved and three patients were in
complete remission; the dosage was 2 mg/kg body weight. Improvement was not seen until after six to 12 weeks and the maximal effect appeared in six to 15 months. Because the "gradual effect makes it difficult to evaluate the effect properly", azathioprine was discontinued every year or two to determine whether there had been a spontaneous remission, but the results of this test were not stated. The drug had to be withdrawn because of agranulocytosis in only one patient. Sepsis and pneumonia affected two patients, one fatally.

Hertel et al\textsuperscript{122} used 150-200 mg azathioprine without steroids, to treat 64 patients. Thirty-three patients were given the drug after thymectomy, so that the prolonged improvement included the effects of both treatments but they implied that azathioprine hastened or enhanced the results of thymectomy (although specific data were not given); four years after thymectomy, they were still improving. Very severe myasthenia was treated with both corticosteroids and azathioprine in 15 cases and all but one improved. Only one patient had severe bone marrow depression that lasted for several months, and therapy did not have to be discontinued in any patient because of hepatic or gastrointestinal toxicity. One patient died of sepsis and another had orchitis.

Reuther et al\textsuperscript{124} studied AChR antibody responses in nine of the patients mentioned by Hertel et al\textsuperscript{122}; three had thymectomies. Titres declined to about 70\% of the initial values by three years and to 40\% after five years. Although all patients improved and seven were in remission, abnormal antibody titres persisted in all of them.

Newsom-Davis et al\textsuperscript{125} gave azathioprine in a daily dosage of 2.5 mg/kg, to six patients; five had thymectomy and five were receiving prednisone. They were compared to seven patients who received the drug (concomitant with prednisone in four and who were also treated by plasma exchange. Six of the 13 patients improved in the observation period of four to 12 weeks, and all patients showed a decline in antibody titres, but there were no significant differences between the two groups. No serious side-effects were reported.

There have been reports of treatment with other immunosuppressive drugs that are not generally available\textsuperscript{126} or with antithymocyte globulin\textsuperscript{127, 128} but the numbers of patients were too small to evaluate critically.

Since 1970, there has been one major change; there is now a clear rationale for the use of immunosuppressive drugs. Because of a single

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**PLASMAPHERESIS**

That myasthenia might be an autoimmune disorder was first suggested by Smithers in 1959\textsuperscript{129}; the theory captured attention after the seminal paper of Simpson\textsuperscript{2} and the demonstration of antibodies to muscle striations by Strauss et al\textsuperscript{1} in 1960. By the end of the decade, however, there was much uncertainty because the striation-binding antibodies were lacking in most cases, could not explain the disorder of neuromuscular transmission, and could not be demonstrated consistently in neonatal myasthenia. Attention was therefore being directed to alterations of cell-mediated immunity, when the first attempts were made in 1969 to remove immunopathogens from the myasthenic patient. Since cell-mediated mechanisms were suspected, it was therefore surprising that the benefits of thoracic duct drainage seemed to be due to something in plasma rather than the lymphocytes.\textsuperscript{131, 132} In retrospect, another observation of that time could have been due to removal of plasma, for an infant with neonatal myasthenia improved promptly after an exchange transfusion for Rh-incompatibility.\textsuperscript{133}

These early observations made more sense a few years later, after the elucidation of experimental autoimmune myasthenia and the demonstration of antibodies to AChR in human patients. The technology of human blood separation had also advanced and it was therefore logical to attempt to remove the pathogenic antibodies by plasma exchange, or plasmapheresis. The first report, by Pinching et al\textsuperscript{134} had a major impact; there is now published information about the use of this technique in 94 patients\textsuperscript{135-138} and 11 patients have been studied at the Colum-
Controversies about the treatment of myasthenia gravis 653

At the Presbyterian Medical Center.\textsuperscript{143}

As usual, the first reports were most enthusiastic, reporting universal improvement in patients who received prednisone and either azathioprine or cyclophosphamide in addition to plasma exchange.\textsuperscript{134, 135} But there have been some disappointments. Howard \textit{et al}\textsuperscript{137} reported improvement in all eight patients who received prednisone as well as "not adequately" in five. Behan \textit{et al}\textsuperscript{142} reported six failures in 21 patients who were receiving both prednisone and azathioprine at the time of plasma exchange.

With a procedure so new, it is not appropriate to speak of "controversy" but there are certainly unanswered questions, and we shall pose some of these.

\textbf{What are the risks of plasma exchange?} Apparently not many. The worst on record were one fatality attributed to azathioprine\textsuperscript{142} and one case of subacute bacterial endocarditis\textsuperscript{134} in which a Scribner shunt had been used; venous catheterisation usually provides adequate access. Susceptibility to infection may increase and influenza-like syndromes may be more frequent after plasma exchange.\textsuperscript{137, 147} Mild transient worsening of myasthenia was reported only by Lisak \textit{et al}\textsuperscript{128} and there is no evidence of adverse effect from loss of pyridostigmine or prednisone in the discarded plasma.

\textbf{How expensive is plasma exchange?} In New York, it costs $350 for the disposable tubing used in each exchange. To this must be added the cost of purchasing and maintaining the cell separator, professional costs for physicians and nurses who perform the exchange, and hospital costs. The total could be at least $800 for each treatment. The procedure could be performed on some ambulatory patients, but laboratory tests are necessary so hospital costs cannot be completely eliminated.

\textbf{Can we predict, on clinical grounds, which patients are most likely to improve?} The answer seems to be "No". Improvement has been reported in patients with mild or severe symptoms, in the young and the elderly (although not yet in many children); before or after thymectomy; with or without prior or concomitant administration of prednisone or immunosuppressive drugs; and with or without thymoma. Congenital myasthenia may or may not be an exception; the procedure failed in two children\textsuperscript{141} and one adult with lifelong symptoms\textsuperscript{143} but this need not imply universal failure in these cases.

\textbf{Can we predict improvement on the basis of antibody titres?} Because plasma exchange was introduced to decrease the titre of circulating antibodies to AChR, it would seem logical to reserve the procedure for patients with high titres of these antibodies, but there are several theoretical reasons why this might not be so: (1) Antibodies are not detected in all patients with myasthenia gravis; depending upon the laboratory and the method, abnormal titres were present in as many as 85%\textsuperscript{144} of patients or as few as 50%.\textsuperscript{145} In either case, a substantial number lack the antibodies. In six patients with congenital myasthenia antibodies were lacking.\textsuperscript{147} (2) When antibodies are present, there is no strict relation between the titre and severity of symptoms. (3) If antibodies to AChR cause the symptoms of myasthenia, these observations suggest that the amount of antibody bound to neuromuscular junctions may not be proportionate to the titre of circulating antibody. (4) Alternatively, the beneficial effects of plasma exchange could be due to removal of a substance other than antibody to AChR. For instance, plasma content of thymic hormone decreases after thymectomy\textsuperscript{147} and is probably also removed by plasma exchange, or there may be changes in immune complexes\textsuperscript{148} or other plasma constituents that have not yet been identified.

However, efficacy of a treatment can force us to revise theory, and it is therefore necessary to evaluate the data. In general terms, and in most patients, AChR antibody titres have declined as patients improved after plasmapheresis,\textsuperscript{136, 141} and two patients with congenital myasthenia who lacked antibody both failed to improve.\textsuperscript{145} But there have been some notable exceptions. Newsom-Davis \textit{et al}\textsuperscript{141} reported a significant drop in antibody titre in one clinical failure of plasma exchange. Dau \textit{et al}\textsuperscript{136} found comparable changes in antibody titres of patients who showed little or much improvement. Howard \textit{et al}\textsuperscript{137} reported improvement in two patients who lacked detectable antibody, and we have seen one such patient.\textsuperscript{148} It is therefore necessary to gather more data about the relation of antibody responses to clinical improvement after plasma exchange.

\textbf{Is it necessary to administer prednisone or immunosuppressive drugs for therapeutic benefit of plasma exchange?} This question cannot be answered from available data and it is of major practical importance because of the risks of the drugs. The question cannot be answered because almost all of the published cases were receiving steroids or immunosuppressive drugs. Both Newsom-Davis \textit{et al}\textsuperscript{141} and Dau \textit{et al}\textsuperscript{136} were impressed by the theory (that drugs are necessary to delay reappearance of the pathogenic
antibodies) and by a few failures in patients who did not respond without these adjuvants. Additionally, the two groups noted “rebound” in four patients who were not receiving azathioprine; that is, antibody titres after plasma exchange gradually increased to levels higher than the original as symptoms returned. However, one of the patients of Dau et al.\textsuperscript{136} was apparently better clinically at the time of antibody “rebound” and symptoms returned only later. Lisak and Schotland\textsuperscript{138} reported that seven of nine patients “have done well without additional treatment”.

Therefore, the role of prednisone and immunosuppressive drugs is not yet defined. 

**Does plasmapheresis add to the benefit of chronic azathioprine therapy alone?** While documenting the acute benefits of plasma exchange, Newsom-Davis et al.\textsuperscript{141} found that the long-term effects of azathioprine alone were not improved by intermittent plasma exchange.

**What are the indications for plasmapheresis?** Considering the costs and still unknown risks, almost all investigators have reserved plasmapheresis for patients who are seriously disabled, usually after other forms of treatment have failed. However, the procedure could be used before other treatments in some circumstances. For instance, it may be effective in the treatment of myasthenic crisis (defined as the need for assisted respiration) if patients do not recover independent ventilation in a few days. Or it could be used to improve the clinical condition of a patient who is being prepared for thymectomy, hoping to make the postoperative course less arduous. For patients who have failed to respond to thymectomy, prednisone or immunosuppressive drugs, plasma exchange has already been of value in providing sustained improvement for weeks or months. For some patients, intermittent plasma exchange may be a suitable alternative to the risks of chronic therapy with steroids or immunosuppressive drugs.

**Are controlled trials necessary to evaluate the therapeutic effects of plasmapheresis?** If a patient has stable and disabling symptoms of myasthenia for several months, and if these symptoms improve objectively within a few days of a series of plasma exchanges, controlled trials seem superfluous. Indeed, it would probably be ethically impossible to design a controlled trial that would include sham plasmapheresis. In this regard plasmapheresis differs from the problems generated by interventions that may take weeks, months or years to have an effect, as in the use of thymectomy, steroids, or immunosuppressive drugs. Another aspect of plasma exchange, however, poses a different problem. The procedure is already being used to treat patients in respiratory crisis, and this may be one of the most important indications. However, it will be difficult to ascertain that treatment with plasmapheresis actually shortens the duration of these episodes of crisis. Because crisis seems to be becoming less frequent, it is not likely that controlled trials can be done in any one institution and we will probably have to compare the duration of crisis before and after the time plasma exchange was introduced.

**Conclusion**

This essay may seem nihilistic to some; it is not meant to be. In our centre, we tend to recommend thymectomy more and more. If the patient does not then improve within a reasonable period of time (and “reasonable” permits considerable latitude, depending upon the severity of symptoms and the patient’s desires as well as the judgement of one or more physicians), then we use prednisone. If there is still no improvement after some uncertain period of time, or if there have been serious side-effects, we then use azathioprine. We have begun to use plasmapheresis, but everyone recognises the uncertainty of the proper application of this technique. Sometimes, if a patient has a crisis before thymectomy, steroids or immunosuppressive drugs are used before the operation.

In practice, therefore, we behave like most other physicians who care for patients with myasthenia. Therapeutic nihilism can go just so far. But I am repeatedly concerned about what seems to be uncritical acceptance of new therapies. Each of the major approaches to alter the course of the disease (thymectomy, prednisone, immunosuppressive drugs) has been endorsed enthusiastically without controlled trials. Yet if any were the panacea that original reports claimed, would plasmapheresis be greeted so warmly?

Indeed, where do all the subjects for plasmapheresis come from, if thymectomy, prednisone and azathioprine are uniformly beneficial?

These techniques are now all enshrined as “accepted practice” and they bear the sacred imprimatur of “the literature”. It becomes increasingly difficult to justify, on ethical grounds, a controlled trial. Yet for treatments that require months or years to bestow benefit, and for a
The author's views have been tempered by association with other clinical investigators, including Paul FA Hoefer, Henry Aranow Jr, Audrey S Penn, Robert E Lovelace, MR Olarte, and Richard Schoenfeldt. Alfred Jaretzki III has provided thoughtful leadership in questions about thymectomy. George Zito gave generous assistance in the analysis of data about steroid therapy. To all of them I am indebted, but none of them is responsible for any errors, intemperate comments, or apparent lapses of judgement in this essay.

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14. Cohan SL, Dretchen KL, Neal A. Malabsorption disease whose “natural” history is not now known, controls would certainly seem desirable. If controlled studies had been done early, we would not be left with so many unanswered questions (table). Clinicians must decide whether to use thymectomy, plasmapheresis, steroids or immunosuppressive drugs, in what sequence or combinations, and when to deem lack of improvement a sign for a different therapeutic approach. Even proponents of specific regimens must admit that the guidelines are not clear. That is the problem, in general terms. It may all sort out in time. In the meantime, each centre will proceed according to the preferences of local physicians. Certainly, myasthenia is not the grave disease it was only 20 years ago; we must be doing something right.

Table Thirty unanswered questions about the therapy of myasthenia gravis

1. How should we determine the optimal dose of a cholinergic drug?
2. Is one cholinergic drug more effective than another?
3. Does cholinergic crisis occur in patients taking, by mouth, conventional reversible inhibitors of anticholinesterase?
4. Is the edrophonium test of value in determining optimal dosage or in defining cholinergic crisis?
5. In the management of myasthenic crisis, should cholinergic drug therapy be discontinued? If so, when should it be re-instituted?
6. Would management be improved by monitoring blood levels of cholinergic drugs?
7. Is thymectomy of proven value in the treatment of myasthenia?
8. If so, how does it work?
9. Which patients are most likely to improve after thymectomy?
10. Does thymectomy prevent progression of myasthenia to a more serious disease? Should every patient with generalised myasthenia have thymectomy? As soon as the diagnosis is made?
11. Should prednisone be given routinely before thymectomy?
12. Should thymectomy be done before adolescence or after some arbitrary older age (for example after 65 years)?
13. Is cervical thymectomy as effective as trans-liquid thymectomy?
14. Is prednisone of proven value in treating myasthenia?
15. Is one corticosteroid preferable to another?
16. What dosage should be used? For how long?
17. If there is no improvement, how long should steroid therapy be continued before deeming the trial a failure?
18. Are steroids safer or more hazardous than azathioprine?
19. How should properly informed consent be obtained before a patient embarks upon prolonged steroid therapy?
20. Are immunosuppressive drugs of proven value in treating myasthenia?
21. Is any one immunosuppressive drug preferable to another?
22. Should immunosuppressive drugs be given before, after, or with steroid drugs?
23. What dosage should be used? Should it be arbitrary, according to body weight, or should mild leukopenia be induced?
24. Which patients will benefit most from plasmapheresis?
25. Does plasmapheresis shorten the duration of myasthenic crisis?
26. Is plasmapheresis of benefit in preparing patients for thymectomy?
27. Is there a role for plasmapheresis in long-term management of myasthenia?
28. Is plasmapheresis beneficial because it removes antibody to AChR or because it removes something else?
29. Can plasmapheresis be beneficial if antibodies to AChR are not detected in an individual patient?
30. Is it necessary to administer prednisone or azathioprine for either acute or long-lasting benefit of plasmapheresis?


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