Treatment of erectile dysfunction with sildenafil citrate (Viagra) in parkinsonism due to Parkinson’s disease or multiple system atrophy with observations on orthostatic hypotension

I F Hussain, C M Brady, M J Swinn, C J Mathias, C J Fowler

Abstract
Objectives—To assess the efficacy and safety of sildenafil citrate (Viagra) in men with erectile dysfunction and parkinsonism due either to Parkinson’s disease or multiple system atrophy.

Methods—Twenty-four patients with erectile disease were recruited, 12 with Parkinson’s disease and 12 with multiple system atrophy, into a randomised, double blind, placebo controlled, crossover study of sildenafil citrate. The starting dose was 50 mg active or placebo medication with the opportunity for dose adjustment depending on efficacy and tolerability. The international index of erectile function questionnaire (IIEF) was used to assess treatment efficacy and a quality of life questionnaire to assess the effect of treatment on sex life and whole life. Criteria for entry included a definite neurological diagnosis and a standing systolic blood pressure of 90–180 mm Hg and diastolic blood pressure of 50–110 mm Hg, on treatment if necessary. Blood pressure was taken at randomisation (visit 2) and crossover (visit 5) lying, sitting, and standing, before and 1 hour after taking the study medication in hospital.

Results—Sildenafil citrate was efficacious in men with parkinsonism with a significant improvement, as demonstrated in questionnaire responses, in ability to achieve and maintain an erection and improvement in quality of sex life. In Parkinson’s disease there was minimal change in blood pressure between active and placebo medication. In multiple system atrophy, six patients were studied before recruitment was stopped because three men showed a severe drop in blood pressure 1 hour after taking the active medication. Two were already known to have orthostatic hypotension and were receiving treatment with ephedrine and midodrine but the third had asymptomatic hypotension. However, the blood pressures in all three had been within the inclusion criterion for the study protocol. Despite a significant postural fall in blood pressure after sildenafil, all patients with multiple system atrophy reported a good erectile response and were reluctant to discontinue the medication.

Conclusions—Sildenafil citrate (50 mg) is efficacious in the treatment of erectile dysfunction in parkinsonism due to Parkinson’s disease or multiple system atrophy; however, it may unmask or exacerbate hypotension in multiple system atrophy. As Parkinson’s disease may be diagnostically difficult to distinguish from multiple system atrophy, especially in the early stages, we recommend measurement of lying and standing blood pressure before prescribing sildenafil to men with parkinsonism. Furthermore, such patients should be made aware of seeking medical advice if they develop symptoms on treatment suggestive of orthostatic hypotension.

Keywords: erectile dysfunction; Parkinson’s disease; multiple system atrophy; viagra; orthostatic hypotension

Estimates of the prevalence of erectile dysfunction in men with Parkinson’s disease show that it is a significant problem, affecting 60% of a group of men compared with an age matched healthy non-parkinsonian group, in whom the prevalence was 37.5%.

Typically the problem only affects men some years after the neurological disease has been established. A survey of young patients with Parkinson’s disease (mean age 49.6 years) that included their partners also, showed a high level of sexual dysfunction with the most severely affected couples being those in which the patient was male. Erectile dysfunction and premature ejaculation were complaints in a significant proportion although in general terms sexual dysfunction seemed to be multifactorial with no single cause identified. There is experimental evidence that dopaminergic mechanisms are involved in determining libido and causing penile erection and the possible role of a central deficiency of dopamine in this disease remains to be examined.

In many cases of multiple system atrophy, urinary incontinence and other autonomic disturbances may precede the onset of major neurological symptoms but urogenital symptoms are usually manifest before symptoms of orthostatic hypotension. In a study of the clinical features and treatment of genitourinary dysfunction in 62 patients with multiple system atrophy, 96% of the men had erectile dysfunction, with this as the first symptom in 37%. In another study of 100 patients with multiple system...
system atrophy, autonomic symptoms, including erectile dysfunction, were the initial features of the disease in 41% and had subsequently developed in 97% of male patients; symptomatic orthostatic hypotension, although present in 68%, was severe and caused fainting in only 15% of patients. The symptoms caused by orthostatic hypotension have been studied in 40 patients with multiple system atrophy, and despite an orthostatic blood pressure fall of >30 mm Hg less than 50% have syncope (unlike pure autonomic failure), whereas others only have symptoms such as fatigue, lethargy, and weakness.

The extensive clinical database that already exists for sildenafil citrate demonstrates that it is efficacious in the management of erectile dysfunction, and safe when appropriately used. The contraindication with any form of nitrate therapy is absolute because a synergistic response can result in severe hypotension. In normal volunteer studies sildenafil reduced mean maximum supine systolic and diastolic blood pressure by about 8 mm Hg and 5 mm Hg without an effect on the heart rate.

The safety and efficacy of sildenafil for the treatment of erectile dysfunction in parkinsonism due to Parkinson's disease and multiple system atrophy had not been studied before. Although multiple system atrophy is less common than Parkinson's disease, it is a disease in which most men have erectile dysfunction and because orthostatic hypotension may be a complication, such a study is of particular relevance.

Methods

PATIENTS

Twenty four male patients with a well documented history of erectile dysfunction, 12 with a diagnosis of Parkinson's disease, and 12 with multiple system atrophy had not been studied before. Although multiple system atrophy is less common than Parkinson's disease, it is a disease in which most men have erectile dysfunction and because orthostatic hypotension may be a complication, such a study is of particular relevance.

Twenty four male patients with a well documented history of erectile dysfunction, 12 with a diagnosis of Parkinson's disease, and 12 with multiple system atrophy had not been studied before. Although multiple system atrophy had previously been made by a consultant neurologist after investigation and according to accepted inclusion criteria.

Patients were excluded if they had no stable sexual partner, penile deformity, other sexual or psychological disorders, or a known history of alcohol or drug dependence. Patients with diabetes, retinitis pigmentosa, a history of stroke or myocardial infarction, or significant cardiac history were also excluded. Treated resting and standing systolic blood pressure was required to be between 90 and 180 mm Hg and diastolic BP between 50 and 110 mm Hg.

Written documentation from the patient's family doctor was required to ensure that they were not on any nitrate medication, and lipid abnormality and thyroid, renal, hepatic, and haematological disease were excluded by screening blood tests.

The median age of the men with Parkinson's disease was 61 years (range 48–68) and for the men with multiple system atrophy, 54 years (range 46–61). The median duration of erectile dysfunction in the men with Parkinson's disease was 54 months (range 12–72) and 57 months (range 24–90) in the men with multiple system atrophy.

Efficacy

Efficacy was judged by the international index of erectile function (IIEF), a psychometrically validated 15 item self administered questionnaire covering the domains of erection, orgasm, desire, intercourse satisfaction, and overall satisfaction with sex life. This includes the ability to obtain an erection (question 3) and the ability to maintain an erection adequate for intercourse on a scale of 0–5 (question 4), where “almost always” or “always” scores 5. In normal healthy men the mean responses for these questions are 4.0 and 4.2.

A quality of life (QoL) questionnaire was also used. This assessed eight factors including whole life, vocational and financial situation, and sex life. Responses range from 0 to 5 (very dissatisfying to very satisfying).

The patient's partner was also given a brief questionnaire to complete independently.

The primary efficacy end points were the responses to questions 3 and 4 of the IIEF and responses to the QoL questionnaire.

Trial Design

The trial was designed as a randomised, double blind, placebo controlled, crossover study with flexible dosage starting at 50 mg active medication or placebo and was conducted over a 24 week period in each patient. The dose could be titrated up to 100 mg or down to 25 mg depending on efficacy and tolerability. Medication was on an “as needed” basis 1 hour before sexual activity, and patients were provided with three tablets a week but advised not to take more than one each day.

At visit 1, consent was obtained before demographic data and screening blood samples were taken. Patients were advised to stop all therapy for erectile dysfunction and sexual activity was encouraged.

After a 4 week run in period (visit 2), patients filled in a baseline IIEF questionnaire and were randomised to receive either 50 mg sildenafil citrate or placebo medication. The first dose was taken in the department and the heart rate with lying, sitting and standing blood pressures were recorded before and 1 hour after dosing. At visits 3 and 4 the dose was titrated up or down if necessary, depending on efficacy and tolerability. At visit 5, after 10 weeks of treatment, patients attended for the crossover and the same procedure was repeated, visits 6 and 7 being similar to visits 3 and 4. At the final visit, the patients repeated the IIEF questionnaire and had a further blood test.

Medication was kept in the pharmacy together with the randomisation detail and only enough was dispensed until the next visit. Patients were required to keep a dosing log.

Statistical analysis was carried out by Pfizer, New York, USA. A crossover analysis of variance (ANOVA) model was utilised with terms for sequence, patient, period, and
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in one patient) until the drop in blood pressure

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and two at visit 5), the standing blood pressure

pressure 1 hour after sildenafil (one at visit 2

1).

showed no postural drop in mean blood

pressure for the placebo medication

was a significant improvement in the quality of sex life. The p

values are based on a crossover ANOVA model.

The same responses after 10 weeks of active

medication starting with 50 mg and titrated up

to 100 mg in most patients showed a marked

and significant improvement in being able to

obtain (question 3) and maintain an erection

(question 4) for sexual intercourse. Although

there was little change in the whole life QoL

there was a marked improvement in sex life

(table 1). The partner questionnaire response

confirmed the improvement. There were no

serious adverse events over the study period.

One patient reported headache and a warm

feeling after taking the active medication but

did not discontinue it.

The predose and postdose measurements of

blood pressure for the placebo medication

showed no postural drop in mean blood

pressure (fig 1). One hour after active medi-

cation there was a change in the average mean

blood pressures of 5 mm Hg, 9 mm Hg, and 9

mm Hg in the lying, sitting, and standing posi-

tions respectively.

MULTIPLE SYSTEM ATROPHY

In the multiple system atrophy group, six

patients were studied before recruitment was

stopped. Four of six were randomised to

receive the placebo for the first 10 weeks and

active medication for the second 10 weeks.

Baseline scores for IIEF questions 3 and 4 were

low, with marginal improvement after placebo

but they improved significantly after active

medication, as did the quality of sex life (table

1).

Three patients had a severe fall in blood

pressure 1 hour after sildenafil (one at visit 2

and two at visit 5), the standing blood pressure

falling from 128/85 to 65/55, 104/60 to 56/32,

and 115/70 to 55/39 respectively. All three

remained conscious but had severe symptoms

and signs of orthostatic hypotension, feeling

unwell and unable to stand for more than a few

seconds. Each patient had to wait (for 4 hours

in one patient) until the drop in blood pressure

became tolerable. Figure 1 shows the average

mean blood pressure changes in this group. On

the basis of this adverse event a decision was

taken to unblind the study and discontinue

recruitment.

Discussion

We have shown that sildenafil citrate is

efficacious in the treatment of erectile dysfunction

in patients with parkinsonism having Parkinson’s disease or multiple system atrophy.

Results

PARKINSON’S DISEASE

Of the 12 men with Parkinson’s disease

recruited, 10 completed the study, one with-
drew consent at visit 3 and another was found to

have carcinoma of the lung during the run in

period and was not treated. Four of 10 patients

were randomised to receive the placebo for the

first 10 weeks and the remainder for the second

10 weeks. Nine of 10 patients reported a good

response to sildenafil citrate, eight titrated up to

100 mg and one titrated down to 25 mg. One

patient reported lack of efficacy and returned to

intracavernosal injections at the conclusion of

the study.

Overall the baseline and 10 week postpla-

cebo data for IIEF (questions 3 and 4) and the

QoL for whole life and sex life were similar.

Table 1 The responses to Q3 and Q4 of the IIEF

questionnaire and the quality of whole life and sex life for

placebo medication and sildenafil citrate in men with

parkinsonism.

<table>
<thead>
<tr>
<th></th>
<th>Placebo (n=16)</th>
<th>Sildenafil (n=14)</th>
<th>Crossover ANOVA model</th>
</tr>
</thead>
<tbody>
<tr>
<td>IIEF:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Question 3</td>
<td>1.56</td>
<td>3.71</td>
<td>p=0.0095</td>
</tr>
<tr>
<td>Question 4</td>
<td>1.44</td>
<td>3.79</td>
<td>p=0.0041</td>
</tr>
<tr>
<td>Quality of life:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Whole life</td>
<td>4.69</td>
<td>4.93</td>
<td>p=0.3505</td>
</tr>
<tr>
<td>Sex life</td>
<td>2.19</td>
<td>4.21</td>
<td>p=0.0073</td>
</tr>
</tbody>
</table>

There was a significant improvement over placebo for both Q3

(contraining an erection) and Q4 (maintaining an erection).

Although sildenafil did not affect the quality of whole life there

was a significant improvement in the quality of sex life. The p

values are based on a crossover ANOVA model.

Figure 1 Average mean (diastolic+one third difference in

systolic and diastolic pressure) lying (ly), sitting (si), and

standing (st) blood pressures in 10 patients with

Parkinson’s disease who tolerated the medication with

minimal effect on the blood pressures (top panel); three

patients with multiple system atrophy who tolerated the

medication without an adverse effect on their blood

pressures (middle panel); and three patients who developed

a symptomatic fall in their blood pressures 1 hour after

treatment with Sildenafil (bottom panel).

Mean blood pressure, the variable shown in

figure 1, was calculated as the diastolic pressure

plus one third of the difference between systo-

lic and diastolic pressures.
Sildenafil enhanced erectile function, enabling intercourse, which resulted in an improved quality of sex life. It was well tolerated in Parkinson’s disease and the minor adverse effects of headache and flushing were transient and required no treatment. However, in multiple system atrophy, sildenafil caused a severe fall in blood pressure in half of the treated patients, resulting in early termination of the study. The adverse event was recognised because the first dose of all medication was taken under supervision in the department. Two of these patients were already receiving treatment with ephedrine and midodrine for orthostatic hypotension and all three patients with this adverse event had a significant fall in postural blood pressure before sildenafil, which potentiated the response. However, on treatment their lying, sitting, and standing blood pressures were within the inclusion criteria for the study protocol. Despite severe and symptomatic hypotension, all three wanted to continue to use the medication and were dismissive of the suggestion that sildenafil was unsafe for them. One patient obtained it via the internet although he was made aware of the possible hazards. The three men with multiple system atrophy who tolerated the medication well had minimal postural fall in blood pressure and although the lying, sitting, and standing pressures were lower with active medication there was no postural accentuation (fig 1). Thus, the presence of orthostatic hypotension and its magnitude was predictive of a further fall and adverse reaction to sildenafil.

Debate continues about the mechanisms causing erectile dysfunction in multiple system atrophy, and a mechanism which includes impairment of dopaminergic mediated pathways in the CNS—which are being increasingly recognised as important for erectile function—should be considered. The demonstrated efficacy of sildenafil in the presence of orthostatic hypotension indicates that hypotension itself, with reduced filling pressures, cannot be causal.

There are often difficulties in early diagnosis and separation of Parkinson’s disease from multiple system atrophy. Although orthostatic hypotension is uncommon in early Parkinson’s disease, this is not the case in multiple system atrophy. We recommend measurement of lying and standing blood pressure before prescribing sildenafil to men with parkinsonism. Furthermore, such men should be followed up with blood pressure measurements and made aware of the need to seek medical advice if they develop symptoms suggesting orthostatic hypotension.

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