A novel therapy with testosterone and sildenafil for erectile dysfunction in patients on renal dialysis or after renal transplantation

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Abstract

Background We undertook a prospective pilot study in a small cohort of patients with renal replacement therapy to determine the cause of erectile dysfunction (ED) and evaluate the role of testosterone replacement therapy and sildenafil.

Methods We investigated 12 patients (eight post-transplant and four on haemodialysis) who presented with ED for hypogonadism and cavernosal insufficiency. We assessed sexual performance before and after treatment by a questionnaire method based on the modified International Index of Erectile Function (IIEF) and National Institutes of Health (NIH) rating. Patients received 250 mg intramuscular monthly injections of testosterone cypionate and 50–100 mg sildenafil orally once or twice weekly for 12 months. Therapeutic response was considered good if the patient could maintain an erection adequate for successful sexual intercourse (NIH criteria) and had a marked improvement in the overall sexual performance (IIEF scoring).

Results Before treatment all patients had severe ED with a poor IIEF score while 11 also had diminished libido. Eleven patients had diminished testicular volume and six had elevated follicle-stimulating hormone levels suggestive of germ cell damage. All patients had a good response to the therapeutic trial of testosterone and sildenafil.

Conclusions Therapy with testosterone and sildenafil may be indicated for those with both cavernosal arterial insufficiency and reproductive hormone abnormalities. Further longer-term data are needed to determine the safety and efficacy of this novel regimen.

Key message points

- Patients with erectile dysfunction (ED) who were receiving dialysis or had renal transplantation were shown to have cavernosal arteriogenic insufficiency and abnormal reproductive hormone profiles.
- Investigations to determine the aetiology of ED may help to guide therapeutic options.
- Therapy with sildenafil and testosterone should be considered in patients who have cavernosal arteriogenic insufficiency and abnormal reproductive hormone profiles. Patients receiving immunosuppressant drugs require monitoring as they are at higher risk of the carcinogenic and other side effects of testosterone.

Introduction

Erectile dysfunction (ED) affects 40–100% of men on renal replacement therapy, including dialysis and transplantation. The aetiology is usually multifactorial. The main organic factors are primary or secondary hypogonadism (hypothalamic pituitary dysfunction, hyperprolactinaemia) and penile arterial insufficiency.

Previous attempts at therapy have included intracavernosal injections of vasoactive agents, such as alphaprostil, papaverine or penile prosthesis. All are associated with poor results due to poor tolerance, prosthetic infection and cylinder leak. The use of the oral agent, type 5 phosphodiesterase inhibitor, sildenafil citrate, has been used successfully in patients on renal dialysis and after renal transplantation.

Vasoactive drugs, however, may fail to correct sexual dysfunction if libido is also affected. Patients with renal transplant and dialysis may have testosterone deficiency, which can cause reduced libido. Reports are conflicting on the value of testosterone in renal patients, especially as testosterone levels may return to normal in some patients after transplantation. However, patients may have normal testosterone levels with diminished Leydig cell reserve, which can contribute to symptoms of androgen insufficiency including diminished libido. Other advantages of testosterone include its potential cavernosal vasodilator activity. A meta-analysis of studies confirms the beneficial effects of testosterone supplementation in patients with ED. A similar improvement is expected in patients with renal replacement, especially if they have hypogonadism.

We undertook a pilot study in a small cohort of patients with renal replacement or on dialysis to determine the cause of ED and to institute a trial of sildenafil and testosterone therapy, as this treatment had been successful in a group of patients with haematological malignancies.

Methods

We prospectively studied 12 patients over the period 1997–2001, who presented with ED, either after renal transplantation (n = 8) or while on haemodialysis (n = 4). Clinical data are given in Table 1. All patients older than 50 years and those with a history of cardiovascular disease were screened for potential contraindications before treatment.

We assessed testicular function by measuring testicular volume (ultrasound and orchidometer). Hyperplasia or adenoma was excluded by prostate ultrasound. All patients had an endocrine profile (Table 2). All hormones were assayed by standard radio-immunoassay using double antibody techniques.
Colour flow Doppler was undertaken to assess haemodynamic function of the penis after injection of a vasoactive agent to induce an erection. The response to intracavernosal injection was graded as: 0 = nil erection; I = tumescence only; II = partial and III = full rigidity according to our previously published data.

All patients had a therapeutic trial of 12 months of 250 mg intramuscular monthly injections of testosterone cypionate and 50–100 mg sildenafil orally, once or twice weekly. Therapeutic response was considered good if the patient could maintain an erection adequate for successful sexual intercourse measured by the National Institute of Health (NIH) rating and International Index of Erectile Function (IIEF) scores. The regime was in accordance with our previously published data. All patients gave written informed consent to participate in this study as approved by the local research ethics committee of the University College London Hospitals Trust.

Results
Table 1 shows the clinical features and Table 2 the baseline endocrine and investigative data of the patients at the onset of the study. All transplant patients had renal transplantation at least 12 months before and had stable renal function for 6 months before and during the study period. No patients had contraindications to therapy. Only one patient (Patient 10) did not have diminished testicular volume and six had elevated follicle-stimulating hormone levels. While only two patients had low testosterone levels, others had a variety of reproductive hormone abnormalities (Table 2). All patients responded to treatment (Table 3). There were no adverse effects or deterioration of renal function and compliance was excellent.
**Table 3 Endocrine and sexual function pretreatment vs post-treatment**

<table>
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<tr>
<th>Patient</th>
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<th>NIH rating</th>
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<tr>
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</tbody>
</table>

IIEF, International Index of Erectile Function; NIH, National Institutes of Health; P, poor; S, satisfactory.

**Discussion**

We found reproductive hormonal abnormalities and cavernosal insufficiency in renal patients presenting with ED. We used combination therapy with testosterone and sildenafil successfully in these patients. We found a similar favourable outcome for erectile performance in cancer patients using the same therapeutic regimen. The main advantage of sildenafil is that it effectively treats ED, whatever the cause. This is relevant in renal diseases where multiple organic and psychogenic disorders are present. Although cardiac, retinal and vasomotor symptoms have been described with sildenafil, the adverse effects are infrequent and generally mild. Sildenafil monotherapy is likely to be less effective in patients who have diminished libido due to Leydig cell insufficiency. Since sildenafil only works on a sexually stimulated penis, addition of testosterone can be synergistic. Testosterone can improve symptoms such as energy, drive and generalised symptoms of depression and fatigue. Some trials have shown the efficacy of testosterone in ED patients. The mixed aetiology of ED in these patients suggests that testosterone and sildenafil may be indicated for those with erectile impotence in renal transplant patients with intracavernosal vasoactive drugs.

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Competing interests. None identified.

**References**