Male Sexual Disorders in Patients with Parkinson Disease: Treatment with Natural Remedies

Abstract

Parkinson’s disease (PD) is a neurodegenerative disorder caused by a progressive deterioration of midbrain dopamine neurons in the substantia nigra. The incidence of PD in male is higher than that in women, but psychological symptoms are as varied as the motor symptoms in both gender. Psychological symptoms encompass a decrease in sexual desire with a long list of thymic, cognitive, behavioral, and neuropsychiatric complications. Their origin can be attributed to the natural course of the disease, side effects of treatment, or both. Clinical researches are mainly focused on the dominant motor symptoms of PD, but the nonmotor features of PD also need attention. Sexual dysfunctions (SD) are one of the most neglected nonmotor symptoms in PD. SD usually begin after the onset of motor disorders and many patients receiving DA agonists as treatment manifest uncontrollable sexual desire, playing thus a major role in the deterioration of the life’s quality of patients and their partners. Research articles were retrieved from PubMed and Google using relevant keywords. Overall, this review emphasize on the sexual disorders widely reported in patients with PD and the types of natural products that are potential future supplementary agents in their control.

Keywords: Hyper sexuality; Erectile dysfunction, Plants; Aphrodisiac; Neuroprotective

Abbreviations: PD: Parkinson’s Disease; ED: Erectile Dysfunction; HS: Hypersexuality; NADH: Nicotinamide Adenine Dinucleotide; 6-OHDA: 6-hydroxydopamine; MPTP: 1,2,3,6-Methyl-Phenyl-Tetrahydropyridine; MAO: Monoamine Oxidase

Introduction

Parkinson’s disease (PD) is a common neurodegenerative movement disorder characterized by extensive degeneration of dopamine (DA) neurons in the nigrostriatal system [1]. It is the second most common human neurodegenerative disorder after Alzheimer’s disease and is characterized by progressive motor disability and cognitive dysfunction [2,3]. PD inflicts a tremendous social and economic burden on modern society due to its disabling nature and high prevalence in the aging population. Currently, the mean age of onset is around 55 years [4].

Clinical studies support that the incidence of PD in male is higher than that in women [5], but psychological symptoms are as varied as the motor symptoms in both gender. Autonomic dysfunction symptoms significantly impair the quality of life of PD patients, even more than motor symptoms [6]. Autonomic dysfunctions are part of a spectrum of non-motor symptoms in patients with PD [7]. Nonmotor symptoms may precede typical motor features of PD by several years [8,9]. The spectrum of nonmotor symptoms encompasses constipation, bladder dysfunction, daytime somnolence, delusions/hallucinations, difficulty in concentration, dribbling, dysphagia, episodes of confusion, fatigue, impulse control disorders, memory problems, mood disorders (depression, anxiety), orthostatic hypotension, pain, paranoia, sensation of breathlessness, sleep disturbances, sweating, and also sexual disorders [10,11].

The conventional strategies of PD treatment are oriented towards increasing the level of the striatal DA; this is achieved either by increasing the DA precursor (levodopa) supply or by inhibiting the DA breakdown by monoamine oxidase (MAO) [12]. However, the therapeutic dose of levodopa is adapted to the neuronal degeneration progresses, leading thus to an increasing demand of Levodopa administration [13]. The resulting insatiable supply of DA is linked to many undesirable side effects, which compromise the benefits and limit of the medical treatment. According to Bhattacharyya & Rosa-Grilo [14], the extent of the DA agonist side effects especially on sexual function is still one of the most under recognized aspects of the condition after 200 years since the very first description by James Parkinson. The focus of management is the relief of the clinically dominant motor symptoms of PD, but the non-motor features of PD also need attention [12]. In particular, sexual disorders (SD), those are commonly reported in patients with PD [15,16] and have attracted attention since the important publication of Giovannoni et al. [17]. The incidence of SD related to the increasing use of DA medications [18] has been explained by the main role of DA in sexual function. DA is a key neurotransmitter in the control of sexual functions, potentially involved in both sexual motivation and control of sexual performance [19].

In recent years, novel therapeutic approaches are being investigated with the intention of influencing pathways leading to hinder or alleviate PD symptoms. Some active compounds derived from plants have been found to exert neuroprotective,
Most men with PD consider erectile dysfunction to be the most distressing of their various disabilities imposed by the disease [25]. A total of 60% of men with PD reported erectile dysfunction, as compared with 37.5% in age-matched controls [26,27]. Many males were also unable to ejaculate and to achieve an orgasm [10]. Recent efforts to understand the physiological basis underlying ED in PD have evidenced an unrecognized high prevalence of testosterone deficiency in elderly male patients with PD, similar to that found in the general population. Testosterone deficiency is also a well-documented cause of decreased libido, erectile dysfunction and decreased work performance [28]. Further erectile function is closely related to sexual motivation during normal intercourse and sexual motivation is partially under the control of dopaminergic system [29]. With regard to the neurodegenerative nature of PD on this system, we can easily understand the occurrence of ED in PD.

Hypersexuality: Hypersexuality (HS) is usually considered to be the result of dopamine control of sexual functions. Hypersexuality is usually considered to be a dopaminergic disorder, and is probably around 20% [22]. In patients with PD, sexual disorders usually begin after the onset of motor disorders. It can be a decrease in desire, a difficulty in reaching orgasm and, for men, disorders of erection or ejaculation [16]. These sexual problems can be related to age (as for PD, the risk of developing erectile dysfunction increases after age 50) or to the disease itself. Indeed, dopamine plays a major role in libido and the occurrence of erection. As a result, the decline in dopamine production suffered by patients with PD may be associated with deterioration in their sex lives [20], but the motor symptoms of PD like tremors, hypertonia or akinesia (rarefaction and slowing of movements) are not likely to promote a satisfied sexuality. Psychological causes (depression and loss of self-esteem) have undoubtedly a negative effect on libido. Sexual desire also disappears because the sexual function is disrupted by disappointment, frustration, and anxiety/depression, all main sources of mental health stress that turn off the libido as well sexual performances. Further, the direct and indirect effects of the disease such as loss of sexual desire does not arise of a simple decline in sexual motivation due to age or PD, it may be also related to medications.

Impact of the treatment against PD

Side effects of Dopamine agonists: Although the cause remains unknown, several pathological processes and central factors such as protein aggregation, mitochondrial dysfunction, iron accumulation, neuroinflammation, and oxidative stress have been reported. Current treatment is primarily symptomatic using anti-Parkinson drugs namely levodopa, carbipoda, DA agonists, monoamine oxidase type B inhibitors and anticholinergics to replace DA [3]. Many patients receiving DA agonist treatment for the severe neurological conditions of PD manifest uncontrollable sexual desire [21]. Effectively, some anti-Parkinson treatments can alter the control of impulses in the brain leading thus to dramatic consequences like sexual assault of partner, family members, or carers [22]. Recent studies showed that 15% of patients receiving a dopaminergic agonist exhibited an impulse control problem like hypersexuality, gambling, compulsive shopping, compulsive eating, or compulsive hobbyism, but this figure is underestimated and is probably around 20% [22].

The occurrence of such side effects is closely related to the nature of PD: the progressive loss of the neurons responsible for the production of dopamine in the nigrostriatal system [1].

The resulting drop of DA level in this brain region is compensated with two kind of treatments: bringing L-dopa, a mixed solution of dopamine in the central nervous system, or administrating dopaminergic agonists, molecules whose structure is close to that of dopamine [12]. The effect of the latter is similar to that of L-dopa but their duration of action is longer. Moreover, the risk of dyskinesias is less, which justify their prescription in first intention.

Mechanisms of action: Each DA receptor is implicated in a wide range of physiological functions. Five DA receptors have been cloned. D1-like receptors (D1 and D5) are positively coupled to adenylate cyclase, and D2-like receptors (D2, D3 and D4) are negatively or not coupled to this enzyme [19]. The main targets of the various dopaminergic agonists used in the treatment of Parkinson’s disease are the dopamine receptors (D2 and D3) located in different areas of the brain. D2 receptors are numerous in structures involved in motricity, whereas D3 are also expressed in other regions, in particular the limbic system [18]. But the latter plays an important role in emotions and pleasure [19]. Thus, the affinity of an agonist to bind D3 instead of D2 represents a greater risk of having impulse control disorder [18]. D2 receptors have also been found to be particularly abundant in the dorsomedian and the dorsolateral nucleus, which innervates the bulbospongiosus and ischiocavernosus striated muscles, involved in penile rigidity in the rat [23].

Some neuroleptics have the ability to block D2 receptors in the anterior pituitary, and correlated to erectile dysfunction [24]. Many others psychoactive substances (heroin, cocaine, cannabis, alcohol) can stimulated the reward system provoking an increasing release of DA. Impulse control problems have been associated to an excessive stimulation by the agonists of the D3 receptors, which may stop inhibiting mechanisms behind compulsive behaviors whilst stimulating reward system circuitry. Scientific research on active ingredients blocking D3 receptors or targeting specifically D2 should get more attention because their identification can improve drug manufacturing.

Main sexual disorders associated to PD

Erectile disorders: Most men with PD consider erectile dysfunction (ED) to be the most distressing of their various Disabilities imposed by the disease [25]. A total of 60% of men with PD reported ED, as compared with 37.5% in age-matched controls [26,27]. Many males were also unable to ejaculate and to achieve an orgasm [10]. Recent efforts to understand the physiological basis underlying ED in PD have evidenced an unrecognized high prevalence of testosterone deficiency in elderly male patients with PD, similar to that found in the general population. Testosterone deficiency is also a well-documented cause of decreased libido, erectile dysfunction and decreased work performance [28]. Further erectile function is closely related to sexual motivation during normal intercourse and sexual motivation is partially under the control of dopaminergic system [29]. With regard to the neurodegenerative nature of PD on this system, we can easily understand the occurrence of ED in PD.

Hypersexuality: Hypersexuality (HS) is usually considered to
constitute a marked increase in sexual interest, arousal, and behavior, which has adverse consequences for the patient and their partner or carers, and is out of keeping with premorbid personality. It is often characterized by a preoccupation with sexual thoughts, frequent demands, and desire for sexual practice that might be quite different from those previously engaged in, and currently, habitual use of sex lines and Internet pornography or contact with sex workers [30,31]. HS is often a consequence of dopaminergic dysregulation associated to higher doses of DA treatment.

Clinical and neuropsychological features of PD cases of patients with HS revealed that some of them developed paraphilias; described a completely new interest in masochistic sex; admitted to indecently exposing them self on several occasions; or drilled holes in the bathroom walls of their house to watch partner undress, or accused of sexually assaulting daughter by inappropriate physical contact. The behaviors led to the breakup of the relationships with partners in the majority of the cases [22].

Treatments of sexual disorders associated to PD

Treatments of erectile dysfunction: ED is the only SD with evidence-based drug treatment available [32]. Sildenafil is an effective treatment for ED in men with several neurological disorders, also PD [33,34]. Its efficacy in PD patients with ED and depression was reported to be 85% [33]. Symptoms of Parkinson’s disease and decreased libido in the non-depressed patient can be treated in different ways. Either by an L-Dopa treatment that will restore the dopamine level and relieve motor symptoms or by using type 5 phosphodiesterase inhibitors, such as Cialis, Levitra, or Viagra, which are also useful against erectile dysfunction. Studies in humans have shown some effect of amphetamine [35] and yohimbine [36] both of which are known to be dopaminergic, in the treatment of erectile dysfunction. Treatment with apomorphine sublingually is another therapeutic option for PD patients with ED. The action is through a dopaminergic effect in the hypothalamus. Doses of 2-4 mg have been recommended, with the erection occurring within 10-25 minutes [10].

Treatments of Hypersexuality: Hypersexuality requires an adaptation of the prescribed dopaminergic treatment to the patient [10]. Introduction of an atypical neuroleptic treatment, or even antiandrogenic should reduced or even stopped HS. However this introduction will slightly conduce to motor function degradation. DA agonist could also be replaced by an anticholinergic, an MAO-B or an atypical neuroleptic at low dose. Moreover, Deep Brain Stimulation of the subthalamic nucleus has also demonstrated positive influence on sexual well being [37].

Natural products used against PD and its complications

Several pathological processes and central factors such as protein aggregation, mitochondrial dysfunction, iron accumulation, neuroinflammation and oxidative stress have been reported as the possible causes of PD [30]. Current treatment is primarily symptomatic using anti-Parkinson drugs namely levodopa, carbidopa, dopamin (DA) agonists, monoamine oxidase type B inhibitors and anticholinergics to replace DA. When drug therapy is not satisfactory, surgical treatments are recommended. Unfortunately, the existing conventional strategies against PD are with numerous side effects, and possess an economic burden. Therefore, novel therapeutic approaches which can regulate pathways leading to neuronal death and dysfunction are needed.

For many years, nature has been providing the primary resource for the discovery of potential therapeutic agents. Remarkably, many natural products from medicinal plants, fruits, and vegetables have been shown to be good anti-Parkinson agents [3]. Since they possess neuroprotective properties not only due to their well-recognized anti-oxidative and anti-inflammatory activities but also their inhibitory roles on iron accumulation, protein misfolding and maintenance of proteosomal degradation as well as mitochondrial homeostasis. Effectively, some natural remedies are specifically sought after for their neuroprotective action and can improve the therapeutic effects of traditional therapies. Among the most effective natural treatments to fight Parkinson’s disease, there are dietary supplements and herbal remedies.

Vitamin D: Essential to prevent osteoporosis and maintain maximum effectiveness of the immune system, vitamin D is a micronutrient also fundamental to the brain. Several studies have shown not only that low levels of vitamin D are associated with an increased risk of neurodegenerative diseases, but also that taking vitamin D supplements helps to contain, and may even prevent, Parkinson’s disease.

NADH: The dinucleotide nicotinamide adenine (NADH) also helps to fight against the manifestations of PD; it is a coenzyme involved in the energy production cycle and able to stimulate the synthesis of dopamine. The NADH dosage is normally 5 milligrams per day.

The most effective herbal remedies for PD and related sexual dysfunctions: The variety of neuroprotective mechanisms of natural plant extracts allowed researchers to target PD progression in different pathological stages through multiple pathways. Interestingly, various plants extracts that have been investigated in preclinical trials for their neuroprotective effects against neurotoxicity induced by PD, also demonstrated aphrodisiac or an aphrodisiac properties in normal condition (Table 1). However the both, neuroprotective and aphrodisiac/ an aphrodisiac properties, have never been tested together on the same model of PD.

Mucuna pruriensis a natural source of levodopa, the most commonly used medication to cure PD. It is also an established herbal drug used for the management of male infertility, nervous disorders, and also as an aphrodisiac [39]. M. pruriens has been used traditionally as an aphrodisiac for both men and women [40]. One of the scientific reasons behind the effectiveness of this herb for this purpose is due to its activity on hormonal profile. M. pruriens reduces excessive prolactin and increases testosterone in the body. Prolactin tends to counteract testosterone in the body, thus reducing sexual interest.

Ginkgo biloba therapeutic effects on cerebral blood circulation and cognitive functions are now known and recognized [41]. G.
gilboa extract contains flavonoid glycosides, mainly composed of kaempferol, quercetin glucorhamnose esters, and terpenes of ginkgolides and bilobalides. These ingredients have been suggested to have the ability to inhibit MAO enzyme and uptake of certain neurotransmitters such as noradrenaline and serotonin in the central nervous system [42]. A recent meta-analysis of 28 clinical studies with Ginkgo has found no significant effect on cognitive function [41]. Further the herb Ginkgo biloba has long been considered as one of the most potent aphrodisiacs for men. Ginkgo extracts have been suggested to improve sexual function by improving the blood flow to the brain as well as to the genital organs. These extracts have also been suggested to increase NO bioavailability, which may have positive impact on sexual function [43].

### Table 1: Potential herbal remedies against Parkinson’s disease related sexual disorders.

<table>
<thead>
<tr>
<th>Plants</th>
<th>Common name</th>
<th>Therapeutic activity on PD</th>
<th>Model</th>
<th>Benefits for sexual function</th>
<th>Studies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mucuna pruriens</td>
<td>Velvet bean</td>
<td>Natural source of levodopa, improved climbing behaviour and restored damaged mitochondria</td>
<td>Mutant model of Drosophila melanogaster; MPTP induced neurotoxicity</td>
<td>Aphrodisiac, mood and libido enhancer; reduce prolactin, enhance testosterone</td>
<td>[38,39,47]</td>
</tr>
<tr>
<td>Curcuma longa</td>
<td>Curcumin</td>
<td>Reduced depletion of dopamine, reversal of glial fibrillary acidic protein (produced by astrocytes during degeneration) and reduced pro-inflammatory cytokines; reduced the iron chelation in the dopaminergic neurons.</td>
<td>MPTP induced model; 6-OHDA mouse model of PD</td>
<td>Aphrodisiac</td>
<td>[48,49]</td>
</tr>
<tr>
<td>Albizia adianthifolia</td>
<td>Sene (Bwondo-Cameroon)</td>
<td>Attenuates 6-OHDA induced anxiety, depression, and oxidative stress in rat amygdala, cognitive-enhancing effects</td>
<td>6-OHDA-lesioned rat model of PD</td>
<td>Anaphrodisiac endowed with estrogenic properties</td>
<td>[50]</td>
</tr>
<tr>
<td>Pausinystalia yohimbe</td>
<td>Yohimbe</td>
<td>Low affinity to some of the dopamine receptors. Antidepressant; Patients demonstrated a vulnerability to yohimbine-induced somatic symptoms (Panick attack)</td>
<td>Parkinsonian patients; chronic mild stress rat model,</td>
<td>Most widely recognized natural aphrodisiac, inhibit testicular lipid peroxidation, increase arterial blood pressure</td>
<td>[44,51,52]</td>
</tr>
<tr>
<td>Carthamus tinctorius L.</td>
<td>Safflower</td>
<td>Reversed the decreased protein expression of tyrosine hydroxylase, dopamine transporter and DJ-1 and increased the levels of dopamine and its metabolite</td>
<td>Rotenone-Induced Rat Model</td>
<td>Very effective aphrodisiac, Improves sexual vigor and thickens semen</td>
<td>[53]</td>
</tr>
<tr>
<td>Humulus japonicus</td>
<td>(Japanese) hops</td>
<td>(in vitro) increased glutathione levels and decreased phosphorylation of ERK1/2 in SH-SY5Y cells exposed to 6-OHDA (in vivo) improved the motor dysfunction and notably reduced dopaminergic cell death and fiber loss in the SNc and striatum caused by 6-OHDA</td>
<td>In vitro and in vivo 6-hydroxodopamine (6-OHDA) models</td>
<td>Possesses estrogenic and vasodilatory properties, treat anxiety and insomnia</td>
<td>[54]</td>
</tr>
</tbody>
</table>

6-OHDA: 6-Hydroxydopamine; SNc: Substantia Nigra Pars Compacta

Yohimbine is a natural tryptamine alkaloid, which can be extracted from the bark of Pausinystalia yohimbe, a tree from Africa and Asia origin. Yohimbe bark extract is traditionally used in Africa as an aphrodisiac [44]. Yohimbine has high affinity to human alpha-2 adrenoceptors, moderate affinity to alpha-1 adrenoceptors, and low affinity to some of the serotonin and dopamine receptors in the central and peripheral nervous systems [45]. Alpha-2 adrenoceptors mediate erection-inhibiting impulses in the central nervous system and Yohimbine is generally believed to enhance central sexual impulse by blocking the alpha-2 adrenoceptors in the locus coeruleus in the brain [46].

*Bacopa monnieri* also called ‘Brahmi’ in India is a brain tonic mainly used to improve memory and concentration, and in general to support cognitive function. Animal model studies have shown that *Bacopa* extracts help prevent degeneration of dopaminergic neurons and may represent good supplementation in the treatment of Parkinson’s disease. *B. monnieri* is also considered in certain part of Asia, as brain tonic, aphrodisiac, memory booster, and tonic for many ailments.

### Conclusion

Sexuality is a basic human right and essential part of healthy life. Sexual disorders, particularly erectile dysfunction and Hypersexuality, are significantly frequent among patients with PD, but they remains underestimated and commonly neglected, despite their considerable impact on the deterioration of the life quality of patients with PD. However, Hypersexuality is more and more recognized and it has been evidenced that it is often associated with PD treatment. Until now, there is no effective cure for PD, and this review propose to search for neuroprotective strategies to stop or slow the disease progression taking into consideration the importance of the development of safe sexual
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Conflict of Interest

No conflict of interest exists.

References


