Effect of Extracts of the Aerial Parts and Roots from Four Ferulago Species on Erectile Dysfunction in Rats with Streptozotocin-Induced Diabetes

Songül KARAKAYA1*, Didem YILMAZ ORAL2, Serap GÜR2, Hayri DUMAN3, Ceyda Sibel KILIÇ4

1Atatürk University, Faculty of Pharmacy, Department of Pharmacognosy, Erzurum, Turkey
2Ankara University, Faculty of Pharmacy, Department of Pharmacology, Ankara, Turkey
3Gazi University, Faculty of Science, Department of Biology, Ankara, Turkey
4Ankara University, Faculty of Pharmacy, Department of Pharmaceutical Botany, Ankara, Turkey

ABSTRACT

Objectives: The extracts of Ferulago species are used as aphrodisiacs in Turkey and so we aimed to demonstrate in vivo and in vitro the relaxant effect of four Ferulago species’ extracts on the corpus cavernosum (CC).

Materials and Methods: A total of 30 adult male Sprague Dawley rats were divided into control and diabetic groups. Diabetes was induced by a single intraperitoneal injection of 40 mg/kg streptozotocin. In vivo erectile responses were obtained by stimulation of the cavernosal nerves and repeated after intracavernosal injection of extracts in rats, and the data were expressed as intracavernosal pressure (ICP)/mean arterial pressure and total ICP. The relaxant and contractile responses of CC strips were analyzed in the presence or absence of extracts.

Results: The extracts were active in both control and diabetic rats. The extract-induced maximum relaxation responses (especially of methanol extract of the root of Ferulago bracteata) (98.30±2.6%) were decreased after incubation with L-NAME (44.8±1.8). ODQ, a soluble guanylate cyclase inhibitor, inhibited 77% of extract-induced maximum relaxation in the CC from the control rats.

Conclusion: These species can be utilized in erectile dysfunction and may be an herbal alternative to synthetic drugs.

Key words: Aphrodisiacs, Apiaceae, Ferulago, erectile function

Amaç: Ferulago türlerine ait ekstreler Türkiye’de afrodizyak olarak kullanılmaktadır, bu nedenle in vivo ve in vitro olarak dört Ferulago türüne ait ekstrelerin korpus cavernosum (CC) üzerindeki gevşetici etkisini göstermeyi amaçladık.

Gereç ve Yöntemler: Kontrol ve diyabetik gruba ayrılan toplam 30 yetişkin erkek Sprague Dawley sıçanı, 40 mg/kg Streptozotocin ile intraperitoneal olarak tek seferlik enjeksiyon ile diabetik olarak indüklenmiştir. Kavernosal sinirlerin uyarılmasıyla in vivo erekt yanıtlandırılması elde edildi ve sıçanlarda intrakavernozal ekstraktların enjeksiyonu sonrasında tekrarlanması ve veriler intrakavernozal basınç (ICP)/ortalama arteriyel basınç ve toplam ICP olarak ifade edildi. CC striplerin gevşetici ve kasılma yanıtları, ekstraktların varlığı ve yokluğunda analiz edildi.

Bulgular: Ekstraktların hem kontrol hem de diyabetik sıçanlarda aktif olduğu bulundu. Ekstraktlar (özellikle Ferulago bracteata kök metanol ekstresi) ile maksimum gevşeme yanıtları (%98,30±2,6) L-NAME (44,8±1,8) ile inkübasyondan sonra azalmıştır. ODQ, çözünebilir guanilat siklaz inhibitörü, kontrol sıçanlarından CC’de ekstraktların indüklediği maksimum gevşemenin %77’sini inhibe ettiği görülmüştür.

Sonuç: Sonuç olarak bu türler erektil disfonksiyonu önleme ve sentetik ilaçların alternatif olarak kullanılabilir.

Anahtar kelimeler: Afrodizyak, Apiaceae, Ferulago, erekt disfonksiyon

*Correspondence: E-mail: ecz-songul@hotmail.com, Phone: +90 442 231 52 50 ORCID-ID: orcid.org/0000-0002-3268-721X
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INTRODUCTION

Diabetes is one of the most prevalent causes of erectile dysfunction (ED), which eminently influences the quality of life, and the risk of developing ED in diabetic men is threefold higher than that in healthy men. As compared with the other complications of diabetes, the development of ED begins at an earlier age. Moreover, the incidence and severity of ED increase with the duration of diabetes and multifactorial mechanisms including neurogenic and vasculogenic factors are involved in diabetic ED. The efficacy of some ED treatments is limited for diabetes-associated ED. For example, men with diabetes frequently show a poor response to first-line oral phosphodiesterase type 5 (PDE-5) inhibitors. An alternative therapy choice may be phytotherapy for diabetic ED.

In the present study, we examined the effect of lyophilized aqueous and methanol extracts of Ferulago species growing naturally in Turkey on erectile tissue. In Turkey these species are known as “çağşır” or “çaşşır” and are utilized conventionally as an aphrodisiac in South and Southeast Anatolia. Actually, many species that belong to the genera Ferulago, Prangos, and Ferula have been utilized for this aim. These species are utilized in rutting of goats and sheep, and water decoctions of the roots and aerial parts are administered orally as aphrodisiacs. In Turkey Ferulago species are usually well known for their aphrodisiac activities like various plants in other countries. Apart from their medicinal usage, they have been consumed in salads or as spices due to their special odor, and used as food for goats and deer.

Ferulago W. Koch. (Apiaceae) is represented by 34 taxa in Turkey, 19 of which are endemic. For this reason Anatolia is considered to be the gene center of this genus. Ferulago blancheana Post ex Boiss., Ferulago pachyloba (Fenzl) Boiss., and Ferulago bracteata Boiss. & Hausskn. are endemic perennial species growing only in Kayseri, Central Anatolia; Niğde, Central Anatolia; and Gaziantep, Southeastern Anatolia, Turkey, respectively, but Ferulago trachycarpa Boiss. is not an endemic species, growing in Antalya. During our studies, we found that aqueous and methanol extracts of the roots and aerial parts from Ferulago species produced relaxation in precontracted rat corpus cavernosum (CC). Therefore, we planned to investigate the pharmacological profile of their relaxant effect by using isolated CC tissue in vivo and in vitro. This study aims to give the first report to evaluate the effect of extracts from F. blancheana, F. pachyloba, F. trachycarpa, and F. bracteata on ED in rats with streptozotocin (STZ)-induced diabetes.

MATERIALS AND METHODS

Plant material

Flowering plants of F. blancheana, F. pachyloba, F. trachycarpa, and F. bracteata were collected in 2014 from Kayseri, Niğde, Antalya, and Gaziantep (Turkey), respectively, and identified by Prof. Dr. Hayri Duman, a plant taxonomist at the Department of Biology, Faculty of Science, Gazi University. The voucher specimens are kept in the Herbarium of Ankara University, Faculty of Pharmacy (herbarium numbers AEF 26673, AEF 26674, AEF 26677, and AEF 26676, respectively).

Extraction

The air-dried roots and aerial parts of these species were powdered and macerated three times with methanol for 8 h in a water bath not exceeding 45°C (3×200 mL) using a mechanical mixer at 300 rpm, separately. The extracts were filtered and concentrated until dryness by rotary evaporator (Heidolph VV2000, Germany). Moreover, 50 g of roots and aerial parts from these plants were ground and macerated with 200 mL of distilled water for 8 h/3 days at 30 to 35°C, separately. The aqueous extract was filtered, frozen (Sanyo Medical Freezer, Germany), and lyophilized (Christ® Gamma 2-16 LSC, Germany) to give aqueous extracts from the roots and aerial parts. The amounts of the powdered plants and extracts obtained are given in Table 1.

Animals

Adult male Sprague Dawley rats (350–400 g) received a dose of streptozotocin (STZ, 40 mg/kg, i.p.) within a citrate buffer (pH 5.5) on the day of use. Measurement of blood glucose levels was carried out using an Accu-Chek glucometer (Roche Diagnostics, Indianapolis, IN, USA) after the induction of diabetes. The animals were housed in separate cages on a 12-h light–dark cycle and were fed standard water and chow ad libitum. This study was approved by the Institutional Animal Care and Use Committee of Ankara University (2014-15-86).

In vivo assessment of erectile function

To assess erectile function in vivo, intracavernosum pressure (ICP) (ICP, mmHg) was monitored in the rats. The rats were anesthetized with ketamine (50 mg/kg, i.p.) and the trachea was cannulated [polyethylene, (PE)-240 tubing] to keep the airway open, and the carotid artery was cannulated (PE-50 tubing) to measure the main arterial pressure (MAP, mmHg), by a transducer (Statham, Oxnard, CA, USA) attached to a data acquisition system (Biopac MP 100 System, Santa Barbara, CA, USA). A 25-gauge needle filled with 250 μL heparin and connected to polyethylene-50 tubing was placed in the right crus of the penis connected to a pressure transducer to measure ICP indissolubly. The right major pelvic ganglion and cavernosal nerve (CN) were represented. A stainless-steel bipolar hook electrode for stimulation was installed around the CN postero-lateral to the prostate on one side, and the MAP

| Species         | Used parts | Powdered (g) | MeOH (g) | Lyophilized aqueous (g) |
|-----------------|------------|--------------|----------|-------------------------|
| F. blancheana   | Root       | 50           | 6.62     | 5.78                    |
|                 | Aerial part| 50           | 3.22     | 4.78                    |
| F. pachyloba    | Root       | 50           | 7.25     | 6.98                    |
|                 | Aerial part| 50           | 3.32     | 4.01                    |
| F. trachycarpa  | Root       | 50           | 6.77     | 7.76                    |
|                 | Aerial part| 50           | 3.41     | 3.67                    |
| F. bracteata    | Root       | 50           | 7.94     | 5.99                    |
|                 | Aerial part| 50           | 3.65     | 4.88                    |
(mmHg) and ICP (mmHg) were indissolubly measured with pressure transducers. The CN was stimulated (2.5, 5, and 7.5 V, 15 Hz, 30 s train duration) with a square pulse stimulator (Grass Instruments, Quincy, MA, USA) and electrical stimulation was inducted distally to the ligature. The measurements were repeated after intracavernosal administration of extracts (1 µM) in groups.

Isometric tension measurements
Cavernosal tissue (CC) strips were placed in organ bath chambers and maintained in Krebs-bicarbonate solution (containing, mM: KCl 4.7, NaCl 118.1, MgSO₄ 1.0, KH₂PO₄ 1.0, NaHCO₃ 25.0, glucose 11.1, and CaCl 22.5, pH 7.4). The strips (1×1×9 mm³) were dissected and combined under 1 g of resting tension in a 20 mL organ bath. The organ chamber temperature was kept at 37°C by a circulating water bath and continuous bubbling with a mixture of 95% O₂ and 5% CO₂. The tissues were permitted to equilibrate for a minimum of 60 min, and the bath solution was changed every 15 min. Electrical field stimulation (EFS) of the autonomic nerves (duration: 15 s; amplitude: 50–90 V; frequency: pulse width: 5 ms) was achieved by the use of platinum electrodes, placed on either side of the tissue strip (Grass Instruments, Quincy, MA, USA).

In the first series of trials, CC strips were precontracted with phenylephrine (Phe, 10⁻⁵ M) and allowed to relax after administration of the extracts. The relaxation response curves to the extracts were also acquired in the presence of the nonspecific nitric oxide (NO) synthase inhibitor L-NAME (L-N(G)-nitroarginine methyl ester, 100 µM) and soluble guanylate cyclase inhibitor ODQ (1H-[1,2,4]-oxadiazolo[4,3-a] quinoxaline-1-one, 30 µM).

In the second series of trials, acetylcholine (ACh)-, EFS-, sildenafil-, and sodium nitroprusside (SNP)-induced relaxation responses were stimulated after precontraction of CC strips with Phe (10⁻⁵ M) in the presence or absence of the extracts (100 µM).

Statistical analysis
All results are expressed as mean ± standard error and differences between means were statistically analyzed using one-way ANOVA followed by Bonferroni’s complementary analysis, with p<0.05 considered to indicate statistical significance. At the end of the experiment, each CC strip was weighted. All contractile responses were expressed as mg of tension developed per mg of corporal tissue and relaxant responses were calculated as a percentage of Phe-contraction.

Drugs
All drugs were purchased from Sigma Chemical Co. (St. Louis, MO, USA).

RESULTS
Extraction
Methanol and lyophilized aqueous extracts of the roots and aerial parts from Ferulago species were evaluated for their effect on ED.

Characteristics of animals
Body weight of the diabetic rats was considerably lower than that of the control rats (Figure 1a, p<0.001). Blood glucose levels in the diabetic group were considerably higher than those in the control group (Figure 1b, p<0.001).

In vivo erectile responses in both groups
ICP/MAP values in the control rats were higher than in the diabetic rats (p<0.001; Figure 2), which was reversed by intracavernosal administration of the extracts (1 µM). Moreover, total ICP values were decreased in the diabetic group compared with the control group (p<0.001; Figure 2). After the intracavernosal administration of the extracts (1 µM) total ICP values were restored in the diabetic group at all voltage levels, except for the 7.5 voltage level (Figure 2).

In vitro responses of CC strips
The extract-induced maximum relaxation responses (especially methanol extract of the roots from F. bracteata) (98.30±2.6%) were decreased after incubation with L-NAME (44.8±1.8, Figure 3a). ODQ, a soluble guanylate cyclase inhibitor, inhibited 77% of extract-induced maximum relaxation in the CC from the control rats (Figure 3).

The endothelial-dependent relaxation response to ACh (1 mM) in the control rats was higher than in the diabetic rats, which was increased after the incubation of the extracts (100 µM) in the control and diabetic groups (Figure 4).
EFS-induced relaxation response at 20 Hz was decreased in the diabetic group compared with the control group, which was restored by the incubation with the extracts (100 µM). There was no difference in EFS-induced relaxation response in the control rats between the presence and absence of the extracts (Figure 5).

SNP-induced endothelial-independent relaxation response at 0.1 µM dose relaxation was not different in the control rats when compared with the diabetic rats (Figure 6). However, relaxation responses to SNP were enhanced in the presence of the extracts (100 µM) in the diabetic and control rats.

The relaxation response induced by the PDE-5 inhibitor sildenafil at 1 µM dose was considerably reduced in the diabetic rats when compared with the control rats (Figure 4d). After incubation of the extracts (100 µM), relaxation responses to sildenafil were higher in the diabetic and control rats (Figure 7).

DISCUSSION

In the present study, we aimed to examine the relaxant effect of methanol and lyophilized aqueous extracts of the roots and aerial parts of \textit{F. blancheana, F. pachyloba, F. trachycarpa}, and \textit{F. trachycarpa} in the CC with \textit{in vivo} and \textit{in vitro} studies. Corporal smooth muscle relaxation plays a significant role in erection. Smooth muscle relaxation, which is interceded by NO throughout sexual stimulation, is synthesized in the nerve terminals of parasympathetic noncholinergic and non-adrenergic nerves in the penis as well as by the endothelial cells lining the blood vessels and lacunar spaces of the CC.\textsuperscript{11} The first data provide basic mechanistic information concerning the extract-induced dose-dependent relaxation in rat CC. The major findings of the study show that (i) the extracts relax rat CC in a concentration-dependent manner; (ii) the NO-cGMP pathway plays an important role in mediating extract-induced relaxation; and (iii) they partially restore \textit{in vivo} erectile function in diabetic rats.

Penile erection in response to CN stimulation was confirmed \textit{in vivo} in a diabetic animal model. Our data showed that diabetes reduced the \textit{in vivo} erectile response and the \textit{in vitro} relaxant response of the CC to EFS. Amazingly, erectile responses (ICP/MAP and total ICP) gained after cavernous nerve stimulation except 7.5 V were augmented in the extract-injected diabetic group, as compared with the vehicle-injected diabetic group. In \textit{in vitro} studies, the nitricergic relaxation response to EFS in the diabetic rats was increased by the incubation of extracts. There were no previous data to evaluate the effect of these species on erectile function. However, the extract treatment reduced the diabetes-induced renal damage related to the diabetic nephropathy.\textsuperscript{15} Moreover, the treatment improved the activities of enzymatic and nonenzymatic antioxidants,\textsuperscript{13} and also \textit{in vitro} increased the glycolytic activities.\textsuperscript{14} These results indicate a rationale for more studies using combinations of extracts and phosphodiesterase-5 inhibitors in diabetes-induced ED.

The present study showed that extract-induced relaxation in the CC from the diabetic group was not changed compared with that from the control group. The data support the intracavernosal administration of extracts to augment erectile responses. It seems that the extract responses serve as the normal activity in
vivo and *in vitro* in diabetes. Moreover, relaxation to the extracts was calmly inhibited after precontraction with KCl. Potential sensitive calcium channels are forced by depolarization of the plasma membrane when the extracellular K\(^+\) concentration is augmented. Potential sensitive calcium channels were activated by depolarization of the plasma membrane when the extracellular K\(^+\) concentration was enhanced. In the current study, we researched the underlying mechanism of the extracts’ effects on erectile responses that can be mediated by the NO/cGMP-dependent pathway, which is damaged in diabetes. No earlier study appears to have been done on the mechanism of the extracts in penile tissue. The extracts are most likely to have a role in the NO-cGMP signaling pathway, mediating CC relaxation responses.

**Figure 4.** Relaxation responses to single doses of ACh in the presence of extract of FBlR, FBlH, FPR, FPH, FTR, FTH, FBrR, and FBrH, respectively. Data represent mean ± standard error of mean of 6–8 observations. *p<0.05, **p<0.001 vs. control value, §p<0.05, §§p<0.01 vs diabetic value.

ACh: Acetylcholine, FBlR: Root of *F. blancheana*, FBlH: Aerial part of *F. blancheana*, FPR: Root of *F. pachyloba*, FPH: Aerial part of *F. pachyloba*, FTR: Root of *F. trachycarpa*, FTH: Aerial part of *F. trachycarpa*, FBrR: Root of *F. bracteata*, FBrH: Aerial part of *F. bracteate*
In the isolated CC from the diabetic group, the endothelium-dependent relaxation response to ACh was considerably reduced, which was potentiated in the presence of the extracts. There were no previous supporting data similar to these findings.

There was no difference in the endothelial-independent relaxation response to SNP between the control and diabetic rats, which was enhanced in the groups after incubation of the extracts. In previous studies, SNP-induced relaxant responses did not change in diabetic rats when compared with the controls.15,16

In the present study, relaxation responses to the PDE-5 inhibitor sildenafil in CC strips were lower in the diabetic rats than in the control rats. There was no difference in relaxant response to sildenafil between the control and diabetic rats’ CC after incubation of the extracts. This finding indicates that these species have a potential effect on penile function by means of various pathways to contribute to erectile function in diabetic rats.

As shown in Figure 1, among the extracts, the methanol extracts of roots (especially roots of *F. bracteata*) showed the best activity. On the other hand, lyophilized aqueous extracts...
of the aerial parts (especially *F. blancheana*) showed the worst activity. EFS relaxation responses decreased from 40% in the controls rats to 3% in the diabetes rats. However, as a result of 15-min incubation of the extracts, the EFS relaxation responses increased to 21%. Similarly, acetylcholine relaxation responses decreased from 38% in the controls to 13% in the diabetic rats. However, as a result of 15-min incubation of the extracts, acetylcholine relaxation responses were increased by 40% and were higher than those in the controls. Sildenafil relaxation responses were 92% in the controls and 74% in the diabetic rats, but, as a result of 15-min incubation of the extracts, acetylcholine relaxation responses were increased by

![Figure 6. Relaxation responses to single doses of SNP in the presence of extract of FBIR, FBlH, FPR, FPH, FTR, FTR, FBrR, and FBrH, respectively. Data represent mean ± standard error of mean of 6-8 observations. *p<0.05, **p<0.001 vs control value. §p<0.05, §§p<0.01 vs diabetic value](image)
95% and were higher than those in the controls. SNP relaxation responses were 90% in the controls and 85% in the diabetic rats. However, as a result of 15-min incubation of the extracts, acetylcholine relaxation responses were increased by up to 94% and were higher than those in the controls. The results are shown in Figures 1-7.

Figure 7. Relaxation responses to single doses of sildenafil in the presence of extract of FBIR, FB1H, FPR, FPH, FTR, FTH, FBrR, and FBrH, respectively. Data represent mean ± standard error of mean of 6-8 observations. *p<0.05, **p<0.001 vs control value. §p<0.05, §§p<0.01 vs diabetic value.

FBIR: Root of F. blancheana, FB1H: Aerial part of F. blanchean, FPR: Root of F. pachyloba, FPH: Aerial part of F. pachyloba, FTR: Root of F. trachycarpa, FTH: Aerial part of F. trachycarpa, FBrR: Root of F. bracteata, FBrH: Aerial part of F. bracteate
CONCLUSIONS
The present study primarily revealed the useful effect of intracavernosal administration of extracts in improving erectile function in diabetic rats, which is dependent on the NO/cGMP pathway. The preclinical findings should extend our information of the beneficial effects of the extracts on penile function to develop preventive or therapeutic agents and combinations of them, and phosphodiesterase-5 inhibitors may be a beneficial option for diabetes-induced ED.

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Conflict of Interest: No conflict of interest was declared by the authors.

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