Aconiti Lateralis Radix Preparata, the Dried Root of Aconitum carmichaelii Debx., Improves Benign Prostatic Hyperplasia via Suppressing 5-Alpha Reductase and Inducing Prostate Cell Apoptosis

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Benign prostatic hyperplasia (BPH) is a common disease in elderly men which can be characterized by an abnormal enlargement of the prostate associated with lower urinary symptoms. Current medications available for BPH treatment display several adverse effects; thus, the search for effective treatments with less side effects is still ongoing. In this study, we investigated the effect of Aconiti Lateralis Radix Preparata (dried root of Aconitum carmichaelii Debx.; AL), which is an herb used to treat extremely cold symptoms in traditional Korean medicine, on BPH using a testosterone propionate- (TP-) induced BPH rat model. Eight-week inguinal injection of TP induced BPH in rats, the prostate of which was displaying an abnormal proliferation. The pathological proliferation of the prostate was ameliorated by AL treatment of 4 weeks. Pathohistological changes in the prostate including epithelial thickness and lumen area were restored in AL-treated rats. Furthermore, 5α-reductase (5AR) and androgen receptor (AR), the two main factors in the pathogenesis of BPH, were decreased. In addition, the ratio of BAX and Bcl-2, an indicator of apoptosis, was increased by AL as well. Similar results were observed in AL-treated LNCaP prostate cancer cells. AL treatment suppressed the expression of the 5AR-AR axis and increased the ratio of BAX and Bcl-2. Apoptosis in the testis is considered a crucial side effect of finasteride, a 5AR inhibitor used to treat BPH. Our results showed that AL treatment did not display such effects, while finasteride treatment resulted in loss of spermatogenic cells within the prostate. Overall, these results suggest AL as a potentially safe nature-derived therapeutic agent for BPH treatment.

1. Introduction

Benign prostatic hyperplasia (BPH) is a chronic disease commonly found especially in aging men [1]. It is reported that about 90% of males over an age of 80 suffer from BPH and lower urinary tract symptoms (LUTS) [2]. The main causes of BPH are pathologically increased proliferation of the smooth muscle in the prostate, leading to an abnormal enlargement of the prostate. Patients suffering from serious BPH display symptoms such as urinary intermittency, incomplete emptying, weak stream, staining, urgency, and nocturia [3].

The detailed mechanism of BPH pathogenesis is not fully revealed, but testosterone and dihydrotestosterone (DHT) are well known as inducers of BPH [4]. After the male sex hormone testosterone is produced, it is converted into DHT by 5α-reductase (5AR). DHT is considered a main factor of BPH pathogenesis, as it is expressed five times more than testosterone in BPH patients [4]. DHT has 2- to 10-fold higher binding affinity towards androgen receptor (AR) [5], a nuclear receptor which acts as a transcription factor after activation, and in result accelerates the proliferation of prostate cells. After conversion by 5AR, DHT combines with the AR and
induces the expression of the growth factor, which leads to prostatic proliferation [6].

There are two mainly considered medical treatments for BPH. Alpha-blockers, which are inhibitors of α1-adrenergic receptors, relax the smooth muscle tone in the prostate and improve the urinary flow [7, 8]. However, the use of α-blockers is limited because of side effects such as headaches, hypotension, or ejaculation changes (absence of seminal emission, reduced ejaculation volume and force, etc.). Furthermore, the α-blocker itself cannot regulate the size of the prostate [9, 10]. Another class of medication is 5AR inhibitors, such as finasteride (Fi) and dutasteride. Suppression of 5AR by these inhibitors decreases the conversion of testosterone to DHT [11] and eventually leads to shrinking of prostate tissues. However, 5AR inhibitors can also cause side effects, such as loss of libido and erectile dysfunction [12]. Because of the side effects of existing medications, the need for alternative treatment is constantly growing [13].

Aconiti Lateralis Radix Preparata (AL), the processed daughter root of Aconitum carmichaelii Debx., of family Ranunculaceae, has been traditionally used as an essential herbal drug for cold sensation by stimulating heat generation in East Asia [14]. Studies have reported the anti-inflammatory, antitumor, and analgesic effects of AL [15–18], and moreover, its usage to treat colds, polyarthralgia, diarrhoea, heart failure, beriberi, and edema are well known as well [19]. Nowadays, AL is clinically prescribed for diarrhea, syncope, joint pain, rheumatoid arthritis, and other inflammations [20].

The antiproliferative effect of AL on the prostate has been first reported by Dumbre et al. [21]. Dumbre’s team showed that three ayurvedic plants including Aconitum heterophyllum suppressed prostate enlargement in testosterone-treated rats. However, this study only reports the antiproliferative effect lacking the related action mechanism. Therefore, we conducted a study to evaluate the effect of AL on BPH using testosterone propionate- (TP-) induced BPH rats and LNCaP prostate cancer cells.

2. Materials and Methods

2.1. Preparation of AL. The water extract of AL was obtained by boiling 100 g of dried roots of Aconitum carmichaelii Debx. in 1 L of distilled water at 100°C for 150 min based on the traditional extraction method of herbal prescriptions used in Korean medicine [22]. The solution was freeze-dried (yield 12.8%), filtered through a 0.22 μm syringe filter, evaporated, and then stored at −20°C until usage.

2.2. Chemical Reagents. TP was purchased from Wako Pure Chemical Industries (Osaka, Japan), and Fi (≥97% pure) was purchased from Sigma-Aldrich Inc. (St. Louis, MO, USA). Antibody for AR was from Pierce Biotechnology (Rockford, IL, USA), and antibody for 5AR was from Abcam Inc. (Cambridge, MA, USA). Antibody for glyceraldehyde 3-phosphate dehydrogenase (GAPDH) was from Santa Cruz Biotechnology (Santa Cruz, CA, USA), and antibodies for BAX and Bcl-2 were purchased from Cell Signaling Technology (Danvers, MA, USA).

2.3. Animals. Twelve-week-old male Sprague Dawley (SD) rats (body weight 200–220 g) were purchased from the Daehan Experimental Animal Center (Daehan Biolink, Eumsung, Korea). The animals were all maintained in conditions in accordance with the regulation issued by the Institutional Review Board of Kyung Hee University (confirmation number KHUASP(SE)-P-034). The rats were housed in a pathogen-free room maintained at 23 ± 2°C under a 12 h light/dark cycle. Water and standard laboratory diet (CJ Feed Co., Ltd., Seoul, Korea) were provided ad libitum. BPH was induced as previously described [23], that is, by a 4-week pretreatment with daily subcutaneous injections of TP (5 mg/kg/day) in the inguinal region of rats. Fifty milliliters of the injection formula were prepared by dissolving 750 mg of TP in 5 ml of 100% ethanol and then mixing with 45 mL of corn oil. Each rat was injected with 100 μl of this injection formula in the inguinal area. Normal control rats were injected with the same solvent without TP included. Then, the rats were divided into the following four groups (n = 5 per group) and were treated for 4 additional weeks: (a) a normal control (NC) group that received ethanol with corn oil, (b) a BPH group that received TP with corn oil, (c) a positive control group that received finasteride (Fi) (1 mg/kg/day) with TP (5 mg/kg/day), and (d) a group that received AL (20 mg/kg/day) with TP (5 mg/kg/day). AL and Fi were administrated via oral gavage. After the final treatment, animals were fasted overnight and euthanized using CO2, and the ventral region of prostate tissues was obtained as described previously [24]. The relative prostate weight index (prostate index (PI)) was calculated as the ratio of prostate weight (mg) to body weight (100 g). The prostate tissue was divided in half; one half was fixed in 10% formalin and embedded in paraffin for histomorphological assays, and the other was stored at −80°C for further assays.

2.4. Hematoxylin and Eosin (H&E) Staining. The prostate tissue section preparation and H&E staining were performed as described previously [25]. The slides were examined using the Olympus IX71 inverted phase microscope (Olympus Co., Tokyo, Japan). Epithelial thickness and lumen area fold were measured using ImageJ 1.47v software (National Institutes of Health, Bethesda, MD, USA).

2.5. Western Blotting Assay. Protein expression analysis was performed as previously reported [26]. In brief, homogenized prostate tissues or harvested LNCaP cells were lysed with ice-cold RIPA buffer, the insoluble materials were removed, and the proteins were separated by 8% sodium dodecyl sulfate-polyacrylamide gel electrophoresis and transferred onto polyvinylidenefluoride (PVDF) membranes (Billerica, MA, USA). The membranes were then incubated with the primary antibody at 4°C overnight and subsequently incubated with HRP-conjugated AffiniPure Goat Anti-Rabbit IgG (Jackson ImmunoResearch Labs) or
HRP-conjugated AffiniPure Goat Anti-Mouse IgG (Jackson ImmunoResearch Labs). The immunoblot intensity was quantified using ImageJ 1.47v software (National Institutes of Health).

2.6. Cell Culture. The human prostatic cancer cell line LNCaP was obtained from the Korean Cell Line Bank (Seoul, Republic of Korea). LNCaP cells were cultured in the Roswell Park Memorial Institute (RPMI) medium (Gibco, Big Cabin, OK, USA) supplemented with 100 mg/mL penicillin/streptomycin (HyClone, Logan, UT, USA) and 10% FBS (Sigma-Aldrich Inc.).

2.7. MTS Assay. LNCaP cells were seeded (2.5 x 10^4 cells/well) and incubated in the RPMI medium plus 10% FBS for 24 h. Then, the cells were incubated in fresh media containing various concentrations of BBR for an additional 24 h. Cell viability was monitored using the cell proliferation MTS kit by the Promega Corporation as recommended by the manufacturer as previously described [27]. The absorbance was measured at 490 nm in a VersaMax microplate reader (Molecular Devices, Sunnyvale, CA, USA).

2.8. Ultrahigh-Performance Liquid Chromatography (UHPLC) Analysis. An Agilent 1290 Infinity II LC system (Santa Clara, CA, USA) with an Agilent 6550 iFunnel Q-TOF LC/MS system was used to perform high-resolution liquid chromatography-mass spectrometry (LC-MS) of AL according to the manufacturer’s instructions by the Western Seoul Center of Korea Basic Science Institute. Data acquisition was performed using 6200 series TOF/6500 series Q-TOF B.06.01 (B6172 SP1) software (Agilent, Santa Clara, CA, USA). Specific conditions for LC-MS analysis are described in Table 1.

2.9. Statistical Analysis. The data were presented as mean ± standard deviation (SD). Statistical significance (P < 0.05) was analyzed by the Kruskal–Wallis H test followed by Bonferroni’s method as a post hoc test using GraphPad Prism 5 for Windows (GraphPad Software, San Diego, CA, USA).

3. Results

3.1. AL Attenuates Prostatic Hyperplasia in TP-Induced BPH Rats. Changes in prostate tissue size, prostate weight, and prostate index are shown in Figure 1. Rats treated with TP showed an increase in prostate weight and PI by 1.74-fold and 1.88-fold compared to the normal control. AL treatment significantly decreased the prostate weight by 70% and PI by 74%. This effect was relatively higher than that in the Fi group, rats of which showed decreased prostate weight by 62% and PI by 70%. Body weight of rats did not show any difference among all groups (Figure 1(e)).

3.2. AL Restores Histological Changes of Prostate Tissues in TP-Induced BPH Rats. As shown in Figure 2(a), histology of the prostate was assessed by H&E staining. TP administration caused various histological changes in the prostate. The epithelium of the prostate gland with BPH displayed signs of proliferation such as increased epithelial thickness, over-formed acinus area, and decreased lumen area. However, AL-treated rats and Fi-treated rats showed recovered histology similar to NC rats, with less pathological tissue structures (Figures 2(b) and 2(c)).

3.3. AL Regulates 5AR-AR Axis in TP-Induced BPH Rats and LNCaP Cells. The factors related to prostatic hyperplasia were examined by western blot assays using antibodies against 5AR and AR. These factors were increased in the TP-treated BPH group compared to those in the NC group. However, as shown in Figure 3(a), 5AR and AR were significantly decreased by AL treatment when compared to the TP-treated BPH group (0.73-fold and 0.63-fold, respectively). This was further confirmed in vitro, as AL treatment inhibited the expression of 5AR and AR in LNCaP prostate cells (Figure 3(b)), even though AL treatment up to 1000 μg/mL did not affect cell viability of LNCaP cells (Supplementary Figure S1).

3.4. AL Elevates BAX/Bcl-2 Ratio in TP-Induced BPH Rats and LNCaP Cells. We then evaluated the effect of AL on the ratio of BAX and Bcl-2. BAX/Bcl-2 ratio is considered an indicator of cell apoptosis, and Saker et al. have shown this ratio is important in the pathogenesis of both BPH and prostate cancer [28]. The ratio of BAX and Bcl-2 was decreased in prostate tissues of TP-induced rats, and this was restored to a level similar to that in the NC group by AL and Fi treatment (Figure 4(a)). Similar effects were observed in AL-treated LNCaP cells as well (Figure 4(b)), suggesting an apoptotic induction by AL in prostate cells.
3.5. AL Treatment Does Not Cause Pathological Changes in Testis. One of the fatal side effects of 5AR inhibitors such as Fi is apoptotic enhancement in the testis [29, 30]. Based on these studies, we compared the histology of the testis of Fi- and AL-treated rats. As shown in Figure 5, the testicular tissue showed a proper arrangement of germinal cells in the NC and BPH groups, with no histopathologic lesions. However, in the Fi group, lower spermatogenic cell density was observed, which was improved in AL-treated rats. These results show that AL does not display toxicity in the testis and does not impact spermatogenesis.

3.6. Chromatographic Analysis of AL. We used the UHPLC method to analyze components of AL. As a result, we could identify three components with molecular weights of 239.0575, 634.2889, and 648.305 (Figure 6). Further information on the spectrum peaks is presented in Table 2.

4. Discussion

Prostate diseases can be categorized into three groups: prostatitis, BPH, and prostate cancer. Among the three diseases, BPH is the most common disease in men [31]. Prostate tissue consists of four areas: the central zone, peripheral zone, transition zone, and anterior fibromuscular stroma. BPH associates with an overproliferated transition zone and stiffness of the prostatic smooth muscle tone. These abnormal changes in the size and muscle tone cause compression on the urethra, thus leading to LUTS, a common class of symptoms in BPH [32]. The main reason which makes BPH a serious health issue is epidemiology: around
50% of males over 50 and 90% over 80 suffer from the symptoms of this chronic illness [2].

The pathological process of BPH associates with several molecular mechanisms. As described above, the 5AR-AR axis is considered the most crucial pathway of BPH. Besides 5AR and AR, the balance between cell proliferation and apoptosis is also closely related [33]. Apoptosis starts by two different programmed mechanisms. The extrinsic apoptosis is initiated by activation of Fas and tumor necrosis factor receptor 1, followed by cleavage of caspase-8 [34], while the intrinsic pathway begins by release of cytochrome c from the mitochondria and a subsequent activation of caspase-9 [35],
and both pathways meet at the final few stages including cleavage of caspase-3, the effector caspase [36]. The intrinsic apoptosis is regulated by Bcl-2, a mitochondrial outer membrane-located protein which inhibits release of the proapoptotic factor cytochrome c [37]. The BAX gene was the first to be identified as a proapoptotic member among the Bcl-2 family [38]. BAX protein forms a heterodimer with Bcl-2 and then activates the apoptosis signal. Thus, the ratio of BAX and Bcl-2 determines the apoptotic sensitivity [39].

Under the normal condition, prostate tissues express relatively low levels of Bcl-2 and caspase-3, associated with a low level of apoptosis [40].

AL is the dried root of *Aconitum carmichaelii* Debx., a flowering plant of the genus *Aconitum*. As a native plant of East Asia, AL in traditional medicine has been used for centuries to treat extremely cold symptoms in China and Korea [41]. Although AL is extensively used as a potent herb in traditional Korean medicine, its toxicity requires careful preparation. Aconitine and related alkaloids are toxins which can impact the sympathetic-parasympathetic nerve systems and cardiovascular system [42]; however by heat-processing, they can be changed into rather safer alkaloids [43]. In this study, we investigated the possible use of AL as a novel herbal medication for BPH treatment.

The TP-induced BPH animal model [44] has been widely used for BPH studies because this model displays similar histologic characteristics and pathological abnormalities to humans as described by McNeal [45]. AL treatment suppressed the TP-induced enlargement of the prostate by 70% in prostate tissue weight and 74% in PI. The inhibition rate on both factors was higher than that of Fi treatment (62% in prostate tissue weight and 70% in PI). The pathohistological changes such as epithelial thickness and lumen area were recovered by AL as well. TP injection resulted in an increase of epithelial thickness and decrease of lumen area, associated with the increase of acinus numbers. AL-treated rats showed decreased thickness of the epithelium and increased lumen area while reducing the number of acini.

In addition, the two main pathways responsible for abnormal proliferation of the prostate, androgen-related pathway and apoptosis pathway, were regulated by AL treatment. As thoroughly described, the androgen-related pathway is a crucial pathogenic pathway of BPH. When testosterone is converted into the highly affinitive hormone DHT [5] by 5AR, it binds to the nuclear receptor AR and results in increased proliferation by the transcriptional activity of AR [6]. Studies including our previous ones showed improvement of BPH by suppressing these two factors [23–27]. Our current results also demonstrated that AL can reduce the expressions of 5AR and AR in both TP-induced BPH rats and LNCaP prostate cancer cells, suggesting the inhibitory effect of AL on the androgen pathway during the pathogenesis of BPH.

The balance between proliferation and apoptosis, which can be identified by BAX/Bcl-2 ratio, is also closely related to BPH [33]. While normal prostate shows a relatively low level of apoptosis, in BPH situations, imbalance between apoptosis and proliferation is displayed either by decreased apoptotic signals [40] or by increased proliferative factors such as proliferating cell nuclear antigen (PCNA) [46] and Ki67 [47]. In this study, AL treatment showed the restoration of BAX/Bcl-2 ratio which was significantly decreased by TP. In LNCaP cells, AL treatment increased the ratio of BAX and Bcl-2, indicating induced apoptosis in this prostate cancer cell line.

In addition to the suppressive effect of AL on 5AR and AR, AL treatment also significantly increased the ratio of BAX and Bcl-2, indicating upregulated apoptosis. Corresponding to previous reports [40], prostate tissues of NC rats showed relatively lower levels of Bcl-2 and higher levels of BAX, which were pathologically changed by TP injection. AL treatment restored the BAX/Bcl-2 ratio similar to that of NC rats. Similar results were observed in LNCaP prostate cancer cells, and AL treatment dose-dependently increased the ratio of BAX and Bcl-2.

5AR inhibitors such as Fi reduce the size of the prostate gland by inhibiting the activity of 5AR. However, sex-related side effects are frequently reported, including pathological apoptosis in the testis [29, 30]. Our results also demonstrated decreased cell density caused by Fi on the testis tissue, but AL-treated rats did not show any abnormal histological changes in the testis.
Hit identification is the basic level of small molecule discovery, and therefore, it is necessary to identify effective compounds to value the effects of natural products. Our LC-MS analysis identified three constituents which may possibly be responsible for the beneficial effect of AL on BPH. Although the molecular weight of identified compounds did not match exactly with that of previously known constituents of AL, based on previous reports [48, 49], there is a
posibility that component 1 (molecular weight: 239.0575) could be higenamine (C_{16}H_{17}NO_{3}; molecular weight: 271.316), component 2 (molecular weight: 634.2889) could be hypaconitine (C_{33}H_{45}NO_{10}; molecular weight: 615.72) or mesaconitine (C_{33}H_{45}NO_{11}; molecular weight: 631.719), and component 3 (molecular weight: 648.305) could be aconitine (C_{34}H_{47}NO_{11}; molecular weight: 645.746). However, to clarify, further studies should be carried out with different fractions and/or identified constituents to confirm which is the active compound responsible for the effects of AL.

5. Conclusions
In conclusion, we confirmed increased levels of 5AR and AR and a decreased ratio of BAX and Bcl-2 as well as the changes of the histological structure in BPH rats. These changes were reversed by AL treatment, and in addition, AL-treated LNCaP cells also showed similar results. Furthermore, AL treatment did not induce any abnormal changes in the histology of the testis as Fi treatment did. These results suggest AL as a potentially safe nature-derived therapeutic agent for BPH treatment.

Data Availability
All raw data supporting the results of this study are available from the corresponding author upon request.

Conflicts of Interest
The authors declare that they have no conflicts of interest.

Authors’ Contributions
Jinbong Park and Dong-Hyun Youn contributed equally to this paper and are co-first authors. Jae-Young Um developed the experimental design and conducted the study. Jinbong Park wrote the manuscript. All authors read and approved the final manuscript for submission.

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Supplementary Materials
Supplementary Figure S1: effects of AL on cell viability in LNCaP cells. The MTS assay was performed in order to measure the effect of AL on cell viability in LNCaP cells. (Supplementary Materials)

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