INTRODUCTION

Among etiologies of erectile dysfunction (ED), arteriogenic ED is the most common etiology. Considering that increasing population of aging groups have ED, atherosclerosis (AS) originated arteriogenic ED could be the most popular etiology among older groups. Treating ED could be a medical burden [1,2].
Considering the pivotal role of preclinical studies in investigating the fundamental mechanism of arteriogenic ED with AS, an in vivo model that is broadly validated is essential for further research. Currently, there is limited report on such effort. Moreover, there exist difficulties regarding feasibility and reproducibility of such model in the literature [3-10]. Furthermore, reported models have limitations including no endothelial dysfunction unless genetic modification or a special diet is required to evoke endothelial injury. Inducing mechanical injury is feasible by ligation or clamping of iliac arteries [3-5]. However, to induce endothelial dysfunction is not easy. Moreover, ligation or clamping is far from physiologic AS which reflects progressive luminal narrowing.

To overcome limitations of previous studies validating an aging arteriogenic ED model, we adopted chronic pelvic ischemia (CPI). Our previous experience has shown that such CPI model is a sound in vivo model of an aging ischemic bladder [11-14]. In addition, we have examined an arteriogenic ED model with AS in a pilot preclinical study [15].

The aim of the present study was to validate our novel arteriogenic ED model with AS based on molecular and histologic evidence and determine the effect of treatment with a phosphodiesterase-5 (PDE-5) inhibitor.

MATERIALS AND METHODS

1. Animal subjects
   Male Sprague–Dawley rats (400–450 g) at 16 weeks of age were obtained, randomized, and acclimated for two weeks in plastic cages (two rats per cage). Rats were provided free access to purified water and standard compressed feed. Room air temperature was maintained at 22°C±1°C and light was provided from 7 a.m. to 7 p.m.

2. Ethics statement
   This study was approved by the Institutional Animal Care and Use Committee of Korea University (No. KUIACUC-20130523-3). Animal handling followed animal experimentation guide provided by the Animal Laboratory of Korea University, Ansan Hospital.

3. Experimental protocols
   There were three experimental groups: untreated sham-operated rats with a regular diet (Group I, control [n=7]), CPI with cholesterol diet (Group II, ischemia [n=6]), and CPI model with cholesterol diet and mirodenafil treatment (Group III, ischemia with treatment [n=7]). Rats of Groups II and III received an endothelial injury of iliac arteries and 2% cholesterol diet for eight weeks. Group I rats underwent a sham operation. They were maintained with a regular diet. Groups II and III rats received an endothelial injury with cholesterol diet during the eighth week. Eight weeks after operation, Group III rats were fed with mirodenafil plus normal saline for four weeks. Group II rats were fed with normal saline only. Twelve weeks after operation, in vivo measurements of parameters associated with erectile function (maximum intracavernous pressure [ICP] and ICP/mean arterial pressure [MAP]) were performed. Mirodenafil (5-Ethyl-3,5-dihydro-2-[5-([4-(2-hydroxyethyl)-1-piperazinyl]sulfonyl)-2-propoxyphenyl]-7-propyl-4H-pyrrolo[3,2-d]pyrimidin-4-one) was provided by SK Chemical Ltd. (Seoul, Korea) with a purity of 99%. We dissolved mirodenafil using 40% polyethylene glycol-electrolyte solution in water and used it at a dose of 20 mg/kg. The solution (0.3–0.5 mL) was administered via an oral Zonde needle.

4. In vivo chronic pelvic ischemia model
   An in vivo CPI model followed the original model introduced by Nomiya et al [11-14]. After general anesthesia using isoflurane (1.5%–2.5%), an inguinal incision was made. Exposure of the iliac fossa and femoral artery was made and a 2-Fr Fogarty arterial embolectomy catheter (E-060-2F; Edwards Lifesciences LLC, Irvine, CA, USA) was inserted through a small incision in the femoral artery. After upward placement of the catheter to the fossa area, air inflation (0.2 mL) was done and pulled downward to the initiation location of the femoral artery. This manual pulling down was repeated 10 times to induce mechanical intimal injury. The same approach was performed on the contralateral region. For the control group, a sham operation was done in the same manner without inducing mechanical intimal injury.

5. In vivo measurement of intracavernous pressure
   In vivo measurement of ICP was performed under general anesthesia with isoflurane (1.5%–2.5%) at 12 weeks after establishment of CPI or sham CPI opera-
tion. To measure MAP, a 24-gauge angiocatheter was inserted into the left carotid artery. After exposure of the penis corpus cavernosum, an incision was extended to the low abdominal area to introduce the bladder and prostate. A 22-gauge needle was inserted into the corpus cavernosum artery and connected to a polyethylene tube (PE 50; Clay Adams, Parsippany, NJ, USA). After visual confirmation of the right major pelvic ganglion, a bipolar electrical stimulator (Grass SD9; Grass Instrument, Quincy, MA, USA) was placed and the cavernosal nerve was stimulated for 50 seconds with the condition of 5 v, 20 Hz, and 0.5 milliseconds. Two cycles of electrical stimulation were done and the interval of each stimulation was maintained for at least 5 minutes. Main parameters included maximal ICP, ICP/MAP, total ICP, and slope. Measurements were done using AD Instrument PowerLab™ (AD Instrument, Colorado Springs, CO, USA).

**Fig. 1.** Representative charts of intracavernosal pressure (ICP). Increase in ICP by electrical stimulation was observed in the sham group (n=7; A). Increases in ICP were clearly lower in the chronic pelvic ischemia (CPI) model group (n=6; B). Increases in ICP were higher than the CPI but lower than the sham group in CPI with treated group (n=7, C). The upper curve is mean arterial pressure and the lower curve is ICP.
6. Western blot analysis

Sample preparation was done by homogenizing sample tissues that were kept under -70°C and centrifuged at 12,000 rpm for 10 minutes. Protein concentrations were determined by Bradford solution (Sigma, St. Louis, MO, USA). The amount of each protein sample loaded was 40 µg. Sodium dodecyl sulfate (SDS) polyacrylamide gel electrophoresis was done for measuring levels of hypoxia inducible factor 1-alpha (HIF-1α), endothelial nitric oxide synthase (eNOS), and transforming growth factor beta-1 (TGF-β1) by western blotting. For HIF-1α and eNOS, 6% SDS acrylamide gel was used with sampling loading amount of 40 µg. For settings of stacking gel and separation gel, -80 V and -100 V were adopted, respectively. For TGF-β1, 12% SDS acrylamide gel was used with sampling loading amount of 40 µg. For settings of stacking gel and separation gel, -80 V and -120 V were adopted, respectively. Transfer was performed using a polyvinylidene difluoride membrane at 150 mA for 90 minutes. Immunoblotting was performed after blocking and incubation with primary antibodies and secondary antibodies. Primary antibodies specific for HIF-1α, eNOS, and TGF-β1 (Santa Cruz Biotechnology, Santa Cruz, CA, USA) at a dilution of 1:200 in Tris-buffered saline, 0.1% Tween 20 buffer were incubated with the membrane overnight. Actin antibody (Cell Signaling, Danvers, MA, USA) was used at a dilution of 1:3,000 in 10% formaldehyde solution, the specimen was used to prepare paraffin block. After preparing paraffin block sections (5 µm in thickness), H&E staining was done. Intimal injury and lumen were then inspected.

7. Vascular histology

After obtaining a vascular specimen including abdominal aorta and iliac fossa, histological analysis was done. After fixation with a phosphate-buffered 10% formaldehyde solution, the specimen was used to prepare paraffin block. After preparing paraffin block sections (5 µm in thickness), H&E staining was done. Intimal injury and lumen were then inspected.

8. Masson trichrome staining

After washing fixed fragment of corpus cavernosum, dehydration was done using serial ethanol solution. A paraffin block was made and sectioned to slices (10 µm in thickness). Slice was then placed on a gelatin-coated glass for Masson trichrome staining. Quantitative analysis was done using an ECLIPSE Ti-U imaging analysis (Nikon, Tokyo, Japan) and NIS Elements D30 (Nikon). Calculation of smooth muscle (SM)/collagen ratio was done by repeating measurements 10 to 15 times under magnifying powers of 40 or 200.

9. Statistical analysis

All data are described as mean±standard deviation. For analytic method, one-way ANOVA test was used. To simulate non-parametric test, a Kruskal–Wallis test was conducted. Differences were considered significant at p<0.05. All analyses were performed using STATA ver. 14 (Stata Corp. LP, College Station, TX, USA).

RESULTS

1. Intracavernous pressure

Although the slope of each group showed various patterns (Fig. 1), maximal ICP showed prominent differences among groups I, II, and III (at 73.92±22.12 mmHg, 27.50±11.80 mmHg, and 39.68±7.92 mmHg, respectively, p=0.001) (Table 1). ICP/MAP showed statistically significant between control group and ischemia with treatment group; a statistically significant between ischemia group and ischemia with treatment group; a statistically significant between control group and ischemia group.

<table>
<thead>
<tr>
<th>Group</th>
<th>Category</th>
<th>Maximal ICP</th>
<th>ICP/MAP</th>
<th>Total ICP</th>
<th>Slope</th>
<th>Total ICP/MAP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>Mean±SD</td>
<td>73.92±22.12</td>
<td>0.522±0.215</td>
<td>3,542±1,391.29</td>
<td>3.10±0.46</td>
<td>24.37±10.20</td>
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<tr>
<td>Median</td>
<td></td>
<td>77.30±0.60</td>
<td>0.449±0.07</td>
<td>3,266±910.37</td>
<td>2.31±0.60</td>
<td>18.99±1.39</td>
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<tr>
<td>Ischemia</td>
<td>Mean±SD</td>
<td>27.50±11.80</td>
<td>0.159±0.56</td>
<td>1,857±1.136</td>
<td>1.36±0.60</td>
<td>7.86±1.39</td>
</tr>
<tr>
<td>Median</td>
<td></td>
<td>29.51±0.14</td>
<td>0.142±0.07</td>
<td>1,829±10.8</td>
<td>0.60±0.07</td>
<td>7.39±0.14</td>
</tr>
<tr>
<td>Ischemia with</td>
<td>Mean±SD</td>
<td>39.68±7.92</td>
<td>0.281±0.145</td>
<td>1,512±376.6</td>
<td>0.71±0.06</td>
<td>11.72±3.39</td>
</tr>
<tr>
<td>treatment</td>
<td>Median</td>
<td>38.47±0.25</td>
<td>0.251±0.06</td>
<td>1,555±0.737</td>
<td>0.59±0.05</td>
<td>11.72±0.20</td>
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<tr>
<td>P-value</td>
<td></td>
<td>0.001</td>
<td>0.002</td>
<td>0.006</td>
<td>0.059</td>
<td>0.020</td>
</tr>
</tbody>
</table>

ICP: intracavernous pressure, MAP: mean arterial pressure, SD: standard deviation.

aStatistically significant between control group and ischemia with treatment group; bStatistically significant between ischemia group and ischemia with treatment group; cStatistically significant between control group and ischemia group.
prominent differences among groups I, II, and III (at 0.5223±0.2158, 0.1590±0.5667, and 0.2815±0.1425, respectively, p=0.002). In post hoc analyses, the maximal ICP, ICP/MAP, total ICP, and total ICP/MAP of group I were higher than those of group II or group III (all p<0.05). Between groups II and III, only maximal ICP and ICP/MAP showed significant differences (p<0.05).

2. Western blotting analysis
Densitometry was used to obtain quantitative results of Western blot. Results were normalized to levels of actin (Fig. 2). HIF-1α levels of group II were significantly higher than those of group I or group III (0.9051±0.2286 vs. 0.2933±0.0942 or 0.5675±0.2586, p=0.02). eNOS levels of group II were significantly lower than those of group I or group III (0.4527±0.2119 vs. 0.7247±0.0339 or 0.7144±0.3097, p=0.04). TGF-β1 levels of group I were significantly lower than those of group II or group III (0.2547±0.0925 vs. 0.4807±0.2284 or 0.4718±0.3212, p=0.03). There was no significant difference in TGF-β1 level between groups II and III.

3. Vascular histology
A longitudinal section of common iliac arteries (×400) revealed remarked luminal narrowing with intimal thickening and a degenerative change of media in the ischemia group compared to other groups (Fig. 3).

4. Masson’s trichrome staining
The SM/collagen ratio of group I was significantly higher than that of group II or III (0.182±0.05 vs.
DISCUSSION

There is no doubt that AS is a significant etiology for ED [1,2,16]. It could be a grave issue for both individuals and public health considering its deep relationship with the aging process. Establishing a validated arteriogenic ED model with AS is the first step to clearly determine the mechanism and potential treatment for arteriogenic ED with AS. Based on our previous pioneer approach on this issue [15], this study demonstrated its validity with molecular and histological evidences, like the first time. To overcome limitations of previous studies on this issue, reviews were done to systematically design this validation study.

The key pathophysiology of AS change lies in obstructing arterial inflow by narrowed lumen due to artherosclerotic lesions [1,4,10,17,18]. Hence, without intimal injury, an ED model could not show endothelial dysfunction. The main mechanism of arteriogenic ED with AS could be explained with two components: structural and functional changes. Structural change is induced by impaired arterial flow which results in hypoxia and fibrosis. In this change, TGF-β plays an important role [4,5]. Among previous studies, arterial ligation or partial obstruction was a good model to show structural change [4,5,10]. However, these studies did not try to show functional change associated with endothelial dysfunction. The key mechanism of endothelial dysfunction lies on NOS. Nitric oxide (NO) is involved in anti-inflammation, anti-platelet aggregation, and anti-oxidative function [8,9,18]. NOS is composed of three isoforms including neuronal NOS (nNOS), eNOS, and inducible NOS. eNOS is in the endothelium of the cavernosal artery. It has a critical role in the endothelial dysfunction mechanism.

Nowadays, these two main components (structural and functional changes) have a tendency to be overlapped [3,18]. Hence, validation of a promising ED model should include these two components together. Our model meets this demand well. Structural change was mediated by the progressive narrowing of the arterial lumen which is far more physiologic than previous artificial ligation or obstructions. To induce functional change, our model included a long-term cholesterol diet and progressive narrowing of the arterial lumen by intimal injury. This study had additional benefit by showing functional change. By in vivo measurement of ICP, western blotting of molecular markers, and histological analysis, our model was validated as a promising arteriogenic ED model with AS. Moreover, we used male rats at 16 weeks old in the randomization period, meaning that our model could reflect an aging model more properly than other studies.

Another feature of our study was that mirodenafil, a PDE-5 inhibitor, was used as an inner validation tool. Based on its expected effect on ICP, we could conclude that this model was properly established. By integrating all our results, we could conclude that impaired arterial flow was generated and it created secondary cavernosal dysfunction which was manifested by abnormal ICP and increased HIF-1α. This structural change was aggravated, resulting in fibrosis which also affected secondary cavernosal dysfunction and manifested by elevation of TGF-β and decreased SM/collagen ratio. Together with structural change, there was also a functional change in that eNOS was decreased by endothelial dysfunction. One notable finding of our study was that structural change could not be reversed by PDE-5 inhibitor treatment.

Although this study validated a novel ED model, it had several limitations. First, there was no information about the critical role of mRNA, meaning that detailed processes could not be explained. For example, this study could not assert that endothelial dysfunction was preceded by fibrotic change. Second, besides eNOS, there are more molecular markers that can explain functional change, such as nNOS, nerve growth factor, and vascular endothelial growth factor [3]. Third, considering cholesterol resistant features of rat species, other efforts including genetic and longer-term cholesterol diet are needed. Lastly, our control group could not fully show whether characteristics of this model are caused by mechanical injury or cholesterol diet. Hence, additional groups (i.e., a group with real operation plus regular diet and a group with sham operation plus cholesterol diet) are needed to define characteristics of this model in the future.

CONCLUSIONS

This study validated a novel arteriogenic ED model with AS which was successfully induced by CPI based
on molecular and histologic evidences. More promising preclinical studies using this model are needed to elucidate the fundamental pathophysiology and treatment for arteriogenic ED with AS.

ACKNOWLEDGEMENTS

This research was supported by Soonchunhyang University Fund (No. 20190005).

Conflicts of Interest

The authors have nothing to disclose.

Author Contribution

Conceptualization: JHK, DGM. Funding acquisition: JHK. Methodology: JSS, JWK, SWD. Supervision: DGM, JHB, JHL, YSS, JJK. Writing–original draft: JHK, DGM. Writing–review & editing: all authors.

Data Sharing Statement

The data analyzed for this study have been deposited in HARVARD Dataverse and are available at https://doi.org/10.7910/DVN/HBJCKF.

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