Research Article

Potential Mechanisms Responsible for the Antinephrolithic Effects of an Aqueous Extract of Fructus Aurantii

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The potential effects of Fa extract on the prevention and treatment of CaOx nephrolithiasis were analyzed in an ethylene glycol-(EG-) induced CaOx crystallization model in rats and an in vitro assay. Multiple biochemical variables were measured in the urine and kidney. Kidney sections were subjected to histopathological and immunohistochemical analyses. Urolithiasis-related osteopontin (OPN) was evaluated by Western blotting. The in vitro assay revealed the significant inhibition of crystal formation (3.50 ± 1.43) and dilution of formed crystals (12.20 ± 3.35) in the group treated with 1 mg/mL Fa extract compared with the control group (52.30 ± 4.71 and 53.00 ± 4.54, resp.) (p < 0.05). The in vivo experiments showed that prophylactic treatment with Fa aqueous extract significantly prevented EG-induced renal crystallization and pathological alterations compared with nephrolithic rats (p < 0.05). Significantly lower levels of oxidative stress, oxalate, and OPN expression as well as increased citrate and urine output levels were observed in both the low- and high-dose prophylactic groups (p < 0.05). However, in the low- and high-dose therapeutic groups, none of these indexes were significantly improved (p > 0.05) except for urinary oxalate in the high-dose therapeutic groups (p < 0.05). Fa extract prevented CaOx crystallization and promoted crystal dissolution in vitro. Additionally, it was efficacious in preventing the formation of CaOx nephrolithiasis in rats.

1. Introduction

Urolithiasis is a worldwide public health problem, and the economic burden of urinary calculi is enormous. The lifetime incidence of urolithiasis is 5–12%, and approximately 7% of women and 13% of men are affected by this condition [1]. Approximately 80% of kidney stones are composed of calcium oxalate. The mechanisms involved in stone formation are not fully understood. It is generally agreed that urolithiasis involves events such as crystal nucleation as well as aggregation and growth of insoluble particles [2]. Urine is supersaturated with common stone-forming minerals; however, the crystallization-inhibiting capacity of urine does not permit urolithiasis in most individuals. This natural inhibition capacity is deficient in stone-forming individuals [3]. Stone formation has a multifactorial etiopathogenesis involving anatomic, environmental, genetic, infectious, metabolic, nutritional, and socioeconomic factors that are major areas of concern [4].

The current treatments for urolithiasis, including the use of extracorporeal shock wave lithotripsy and surgical intervention for stone destruction, are widely applied in clinical practice to effectively manage urolithic patients. However, the 1-year recurrence rate in the majority of patients after these treatments is approximately 10%, and these procedures are invasive and do not prevent the recurrence of urolithiasis [5]. Certain drugs, such as citrate and thiazide diuretic preparations, may have side effects that compromise their long-term use and are not consistently effective.

Medicinal plants have played a significant role in various ancient traditional systems of medication. Even today, plants provide a cheap source of medicine for the majority of
the world’s population, and such medicines are considered quite safe, with minimal or no side effects [6]. Studies indicate that numerous herbs, including Selaginella lepidophylla, Adiantum capillus veneris Linn., Rubia cordifolia roots, and Cynodon dactylon plants, can be used as antiureolithic agents [7–10]. Fructus Aurantii (Fa), the unripe fruit of Citrus aurantium Linn. (Rutaceae), is a Qi-regulating drug used in traditional Chinese medicine to treat urolithiasis clinically. Previous studies have shown that numerous bioactive compounds, including polymethoxylated flavones (PMFs), flavonoid glycosides, alkaloids, and coumarins, have been identified and isolated from Fructus Aurantii [11]. Studies have shown that certain components of Fructus Aurantii demonstrate pharmacological activity; antioxidant, antimicrobial [12], and anticarcinogenic properties [12, 13]; and neuroprotective effects [14, 15]. Fructus Aurantii has recently attracted the attention of many researchers, and the study of this herb is gradually increasing.

In the present study, we investigated the antiureolithiatic effects of an aqueous extract of Fructus Aurantii on calcium oxalate stones and its possible mechanisms of action using both in vitro and in vivo methods.

2. Materials and Methods

2.1. Plant Materials. Fructus Aurantii was purchased from the Second Hospital of Lanzhou University (Lanzhou, China) and graciously identified by members of the School of Pharmacy at Lanzhou University. A long-lasting decoction is used in traditional Chinese medicine, and the antioxidant levels from medicinal plants are not influenced by aqueous extraction; thus, the traditional decoction method was used to examine the effects of the aqueous extract [16]. Methods for extracting active ingredients from plants have been previously described [17]. Briefly, 500 g of raw material was mixed with 2.5 L of distilled water for 60 min and decoted for approximately 25 min. The supernatant was collected after filtration. Subsequently, 1.5 L of fresh distilled water was added to the residue, and the mixture was boiled. Finally, we combined the two aqueous extracts. After filtration, concentration, and lyophilization, the freeze-dried powder was stored at −80°C until use (yield = 23 g).

2.2. In Vitro Crystallization Assay. A previously described method was used to study the in vitro effects of the Fructus Aurantii aqueous extract on calcium oxalate crystals [18]. Briefly, stock solutions of 15 mL of calcium chloride (8.5 mM) and 15 mL of sodium oxalate (1.5 mM) containing 200 mM NaCl and 10 mM sodium acetate were adjusted to pH 5.7 to achieve calcium oxalate crystallization. A crystal formation inhibition test and crystal dissolution test were conducted. In the crystal formation inhibition test, solutions containing the Fructus Aurantii extract were added concomitantly to the solution containing the crystallization reagents at time 0 h (before crystallization) to obtain a final Fructus Aurantii concentration of 0.3, 0.7, or 1.0 mg/mL. The same amount of vehicle (distilled water) was added to solutions containing the crystallization reagents at time 0 h and was used for the control (0 mg/mL). All samples were maintained under 500 rpm agitation at 37°C for 24 h. For the dissolution test, crystal formation was induced over 24 h as previously described. After crystal formation, solutions of the Fructus Aurantii extract were added to obtain a final concentration of 0.3, 0.7, or 1.0 mg/mL. The same amount of vehicle (distilled water) was added to the formed crystals and was used for the control group (0 mg/mL). Each sample was observed under an inverted microscope (Nikon Corporation, Tokyo, Japan) to determine crystal morphology (400x). Five randomly selected fields were counted 24 h after the addition of the Fructus Aurantii extract or distilled water. The tests were performed as independent triplicates for each extract concentration.

2.3. Animals and Experimental Protocols. The study was approved by the Institutional Animal Care and Use Committee of Lanzhou University. Adult male Sprague-Dawley rats (200–220 g) from the Animal Experimental Center of Lanzhou University were maintained under standard laboratory conditions (temperature: 25 ± 2°C, humidity: 50–60%, and light regime: 12 h light–dark cycle). The animals had free access to tap water and a standard diet. After acclimatization for 1 week, the rats were divided into 6 groups of 10 animals each. The control group received only filtered water ad libitum. The EG group received 1% (V/V) EG in filtered water ad libitum for 28 days. The two preventive groups included an EG + PFa220 group, which received 1% (V/V) EG in filtered water ad libitum for 28 days along with Fructus Aurantii extract at a dose of 220 mg/kg BW per day, and an EG + PFa660 group, which received 1% (V/V) EG in filtered water ad libitum for 28 days along with Fructus Aurantii extract at a dose of 660 mg/kg BW per day. The two therapeutic groups included an EG + TFa220 group, which received 1% (V/V) EG in filtered water ad libitum for 28 days with a Fructus Aurantii extract at a dose of 220 mg/kg BW per day, and an EG + TFa660 group, which received 1% (V/V) EG in filtered water ad libitum for 28 days with a Fructus Aurantii extract at a dose of 660 mg/kg BW per day.

All extracts and standards were given orally once per day.

2.4. Collection and Analysis of Urine. On day 28 of the experimental period, the rats were placed in separate metabolic cages to collect 24 h urine samples, and 0.04% sodium azide was added to the urine to prevent bacterial growth. Following volume and pH determinations, urine samples were centrifuged at 2,000 × g for 5 min to remove debris, and the supernatants were stored at −80°C for the subsequent determination of calcium, magnesium, uric acid, oxalate, and citrate using commercially available kits (Biovision Ltd., Milpitas, CA, USA; Instruchemie Ltd., Netherlands). Urinary 8-IP (a product of lipid peroxidation) was measured using an enzyme-linked immunosorbent assay kit (Quantikine Assay, R&D Systems, USA) following the manufacturer’s instructions.

2.5. Kidney Homogenate Analysis. A 0.1 g sample of renal tissue was mixed with 9 volumes of cold phosphate buffer...
(pH 7.4) and then homogenized. The homogenates were centrifuged at 16,000 xg for 5 min at 4 °C. The supernatants were measured for superoxide dismutase (SOD) and malondialdehyde (MDA) using two kits (Jiancheng Bioengineering Ltd., Nanjing, China).

2.6. Kidney Crystal Deposits and Pathological Examination. Histological studies were performed using a Nikon Eclipse E400 light microscope. Crystal deposits in the kidney were evaluated as previously described [19] using the following point values: no deposits = 0 points; crystal deposits in the papillary tip = 1 point; crystal deposits in the corticomedullary junction = 2 points; and crystal deposits in the cortex = 3 points. If crystal deposits were observed at multiple sites, the points were combined to provide a total score for each pathological section. The pathological alterations were semiquantified based on the area of injury, from 0 to 3, as follows: 0 = invisible lesions; 1 = tubule interstitial inflammatory infiltration with lesion area <20% and mild dilation of the tubules; 2 = tubule interstitial inflammatory infiltration with lesion area <40% and obvious dilation of the tubules; and 3 = tubule interstitial inflammatory infiltration with lesion area >40% and severe dilation of the tubules [17].

2.7. Microbiological Studies. At the time of sacrifice, microbiological studies were carried out on urine aspirated from the bladder. The kidneys were aseptically removed and transected. One-half of each kidney was homogenized separately in 5 mL of sterile normal saline solution (TRI Instruments). Dilutions (10−1, 10−3, and 10−5) of the tissue homogenates were cultured on plates, and the bacteria were enumerated after correction for the dilution factor. Standard microbiological techniques were used. The presence of more than 105 colony-forming units per mL (cfu/mL) of urine indicated the presence of a urinary tract infection. Pyelonephritis was defined as the presence of ≥106 cfu/g of kidney tissue [20].

2.8. Immunohistochemical Staining. After preservation, the kidney sections were boiled for antigen retrieval and treated with 3% H2O2 to remove endogenous peroxidases. After rinsing with PBS, goat serum was used to block the sections. The sections were incubated at 4 °C overnight with a polyclonal OPN antibody (rabbit polyclonal to OPN, ab8448, Abcam, Cambridge, UK) (5 μg/mL). After rinsing with PBS, the sections were incubated at 37 °C with a polymer helper for 20 min. After rinsing three times with PBS, the slides were treated with poly-peroxidase-anti-rabbit/mouse IgG for 2 h before a final wash with PBS and staining with DAB.

2.9. Western Blot Analysis for OPN Protein in the Kidney. Kidney tissue (100 mg) was homogenized in 1 mL of lysis buffer mixed with 10 μL of phenylmethanesulfonyl fluoride (100 mM) before being centrifuged (12,000 xg 5 min) at 4 °C. The supernatants were collected to determine the protein concentration. The proteins were mixed with SDS-PAGE sample loading buffer. The mixture was heated at 100 °C for 5 min before performing SDS-PAGE with a 10% acrylamide resolving gel. The separated proteins were transferred to polyvinylidene fluoride (PVDF) membranes. The PVDF membranes were blocked with 5% skim milk powder for 2h at room temperature before being incubated with OPN antibodies at 4 °C overnight. The PVDF membranes were washed three times with TBST for 10 min each rinse before being incubated with antibodies labeled with horseradish peroxidase for 2 h. The membranes were then washed as previously described, and the proteins were detected by enhanced chemiluminescence.

2.10. Statistical Analyses. All of the data are expressed as the mean ± standard deviation except where otherwise is indicated. Significant differences were determined by Student’s t-test for comparisons between two groups or one-way ANOVA for comparisons of three or more groups. The rates of urinary tract infection or pyelonephritis were compared between groups by χ2 analysis. A p value of <0.05 was considered statistically significant.

3. Results

3.1. In Vitro Assay. The crystal formation inhibition test showed numerous crystals composed predominantly of claviform calcium oxalate in the control group (0 mg/mL) (Figure 1(a)). Crystal formation was significantly inhibited 24 h after the addition of the Fa extract at a concentration of 1.0 mg/mL to the solution containing the crystallization reagents at time 0 h (Figure 1(b)). The Fa extract caused a more prismatic shape in the calcium oxalate crystals (Figure 1(b)). The crystal counts obtained using a light microscope (400x) were 25.10 ± 7.09 (p < 0.05), 15.40 ± 4.32, and 3.50 ± 1.43 (p < 0.05) for solutions incubated with Fa at 0.3, 0.7, or 1.0 mg/mL, respectively, compared to the control (52.30 ± 4.71) (Figure 2). The crystal dissolution test revealed that the number and morphology of crystals from the control group were similar to those of crystals from the control group based on the crystal formation inhibition test (Figure 1(c)). After the addition of the Fa extract at 1.0 mg/mL to formed crystals for 24 h, the number of crystals was significantly reduced (Figure 1(d)). The results were significant at doses of 0.7 (22.80 ± 3.42) and 1.0 mg/mL (12.20 ± 3.35) when compared with the control group (control: 53.00 ± 4.54) (Figure 2).

3.2. Urinary Biochemical Variables. As shown in Table 1, the EG group had significantly higher urinary oxalate when compared with the control group. In addition, the urine volume was increased, and citrate levels were decreased. Preventive treatment with the Fa extract significantly decreased urinary oxalate and significantly increased urine output and citrate levels when compared with the EG group (p < 0.05). However, in the EG + TFa220 and EG + TFa660 groups, except for the significantly lower oxalate levels in the EG + TFa660 group, the above-mentioned variables did not improve significantly compared to those in the EG group.

3.3. Histology. As shown in Figure 3(a), severe swelling, focal hemorrhage, and pale yellow needle-point crystals were
Figure 1: The CaOx crystals were observed under an inverted microscope (400x). (a) A control group exhibits crystal formation during the inhibition test. (b) The Fa extract was concomitantly added at a concentration of 1.0 mg/mL to the solution containing the crystallization reagents at time 0 h for a duration of 24 h. (c) A control group exhibits crystal formation during the crystal dissolution test. (d) The Fa extract was added at a concentration of 1.0 mg/mL to the formed crystals for 24 h.

Figure 2: Number of crystals in crystal formation inhibition and crystal dissolution tests.

Figure 3: Representative images from each group. (a) Observation of the kidneys in the EG group shows some pathological alterations similar to those observed in the kidneys of the EG group. Gross alterations similar to those observed in the EG group were observed in the EG + TFa220 and EG + TFa660 groups. However, only mild swelling and a small number of yellow needlepoint crystals were observed in the EG + PFa220 and EG + PFa660 groups (Figure 3(a)). Severe dilation of the tubules and massive inflammatory infiltration (black arrows) were observed in the EG group (Figure 3(b)). The pathological alterations observed in the EG + PFa220 and EG + PFa660 groups included a slight dilation of the tubules and mild inflammatory infiltration (Figure 3(b)), and the pathological scores of the two groups were reduced compared with the EG group (Figure 4) (p < 0.05). However, no significant differences were observed between the EG + PFa220 and EG + PFa660 groups (Figure 4) (p > 0.05). The pathological scores of the EG + TFa220 and EG + TFa660 groups were significantly elevated compared with that of the control group (p < 0.05) (Figure 4).

3.4. Scoring of Kidney Crystal Deposits. Abundant crystals were observed in all regions of the kidney in the EG group, particularly in the cortex region, using a Nikon Eclipse E400 light microscope (green arrows) (Figure 3(b)). Compared with the EG group, significantly lower scores were not
respectively (EG + TFα220 and TFα660 groups were 80% (8 of 10) and 90% (9 of 10), rates of urinary tract infection in rats in the EG + TFα220 and TFα660 groups (0%, 100%, 20%, 10%, 80%, and 90%, respectively) were consistent with the rate of urinary tract infection in each group.

### 3.5. Microbiological Studies

None of the rats from the control group (0%) suffered from a urinary tract infection, whereas the rate of urinary tract infection in rats in the EG group was 100% (10 of 10). After preventive treatment with Fa extracts, the rates of urinary tract infection in rats in the EG + PFα220 and EG + PFα660 were 20% (2 of 10) and 10% (1 of 10), respectively (p < 0.05 compared with EG group). However, after therapeutic treatment with Fa extracts, the rates of urinary tract infection in rats in the EG + PFα220 and EG + PFα660 groups were 80% (8 of 10) and 90% (9 of 10), respectively (p > 0.05 compared with EG group). The pyelonephritis rates in the control, EG, EG + PFα220, EG + PFα660, EG + TFα220, and EG + TFα660 groups (0%, 100%, 20%, 10%, 80%, and 90%, resp.) were consistent with the rate of urinary tract infection in each group.

### 3.6. Immunohistochemistry

Immunohistochemical staining for osteopontin revealed OPN expression in all groups (Figure 5); however, OPN levels were barely detectable in the control group (Figure 5(a)). The EG group (Figure 5(b)) exhibited OPN expression that was distributed throughout the renal tubular cells of the whole kidney, particularly in the distended tubules. The increased OPN synthesis in the renal tubules of the EG + TFα220 and EG + TFα660 groups (Figures 5(c) and 5(d), resp.) was similar to that of the EG group. However, the EG + PFα220 and EG + PFα660 groups (Figures 5(e) and 5(f), resp.) demonstrated mild OPN staining that was barely detectable.

### 3.7. Western Blot

Western blot analysis of OPN revealed distinct bands at 41 kDa in the EG group and the therapeutic groups (EG + TFα220 and EG + TFα660). No clear bands were observed in the control group or the prophylactic groups (EG + PFα220 and EG + PFα660) (Figure 6(a)). The corresponding band intensities of GAPDH are presented in Figure 6(a). The results of quantitative analysis of OPN expression by densitometry are expressed as the ratio of the OPN band intensity relative to the intensity of GAPDH. These data indicated that the EG group exhibited relatively high levels of OPN expression (Figure 6(b)). The therapeutically treated rat tissues (EG + TFα220 and EG + TFα660) exhibited OPN expression levels that were similar to those of the EG group. In contrast, the control and prophylactic groups (EG + PFα220 and EG + PFα660) exhibited minimal OPN expression (Figure 6(b)).

### 3.8. Oxidative Studies

As shown in Table 2, the EG group exhibited increased renal MDA and urinary 8-IP content and decreased SOD levels when compared with control animals (p < 0.05). Preventive treatment with the Fa extract protected against the oxidative stress induced by the lithogenic treatment (p < 0.05). The above three indicators were nearly restored to normal levels in the EG + PFα660 group. However, there were no statistically significant alterations in MDA, SOD, or 8-IP levels in any of the therapeutic groups when compared with the EG group.

### 4. Discussion

Based on its medicinal use, we evaluated the antiurolithic potential of a Fructus Aurantii extract using different models. An *in vitro* crystal formation inhibition test showed that Fa decreased the crystal count and modified the morphology of CaOx crystals. Similar alterations in calcium oxalate crystal morphology have been reported with Mg$^{2+}$ and citrate [21]. In the crystal dissolution study, Fa decomposed formed CaOx crystals in a concentration-dependent manner. These data are similar to those observed for potassium citrate, a well-known inhibitor of CaOx crystallization that is widely used for the clinical management of urolithiasis [3]. The crystal dissolution study suggests that the potential activity of Fa as a kidney stone inhibitor is associated with complexion of nonionized Ca$^{2+}$ in CaOx crystal embryos, which ultimately leads to the dissolution of CaOx. Indeed, we are not aware of any study in which certain components of Fa aqueous extract have been reported to dissolve CaOx. However, it must be recognized that, under *in vitro* conditions, Fa may dissolve crystal embryos that might grow into stable nuclei, ultimately forming solid bonds. Although crystal production is not always equivalent to stone formation, crystal formation along the urinary tract is the primary requisite for subsequent formation.
Figure 3: Kidney crystal deposits and pathological examination. (a) Gross anatomy of the kidney. (b) Micrograph of renal tissue (100x magnification).

Table 2: Oxidative stress variables in urolithiasis rats treated with Fa extract (X ± s).

| Parameter (unit) | Control | EG      | EG + TFa220 | EG + TFa660 | EG + PFa220 | EG + PFa660 |
|------------------|---------|---------|-------------|-------------|-------------|-------------|
| 8-IP (pg/mL)     | 3.03 ± 0.60 | 9.34 ± 1.77 | 8.70 ± 2.16* | 9.58 ± 2.51* | 3.64 ± 1.19# | 2.59 ± 1.07# |
| SOD (U/mg)       | 302.87 ± 65.57 | 148.16 ± 31.84* | 162.13 ± 45.17* | 145.78 ± 45.80* | 284.99 ± 66.02* | 311.69 ± 25.67* |
| MDA (nmol/mg)    | 1.07 ± 0.30 | 3.17 ± 0.45* | 3.33 ± 0.62# | 3.14 ± 0.88# | 0.98 ± 0.24# | 1.09 ± 0.18# |

Values are expressed as the mean ± SD. *p < 0.05 when compared with the control, #p < 0.05 when compared with the EG group, and @p < 0.05 for the EG + PFa660 group compared with the EG + PFa220 group. P: preventive; T: therapeutic.
stone supersaturation via the formation of soluble calcium oxalate crystals from the urine. Some studies have suggested that disorders of renal tubular calcium reabsorption are the major cause of hypercalciuria. In the present study, hypercalciuria was not observed in the EG group as previously described. A possible explanation for this result is that the formation of CaOx consumes free calcium (Ca) and normalizes Ca levels in the urine. Most importantly, the oxalate values were approximately normal in the EG + Pfa660 group. Oxalate is a more important risk factor in the process of urinary calculi formation than hypercalciuria [28]. The normalization of oxalate excretion levels in the prophylactic groups suggests that the Fa extract exerts a protective effect against stone production. Additionally, the approximate 2-fold increase in urinary volume in the EG + Pfa660 group compared with the EG group revealed distinct diuresis, which effectively diluted the oxalate, increased urine flow, and prevented CaOx deposition.

In addition, crystals can induce the production of reactive oxygen species (ROS) in kidney tissue. The production of ROS can lead to renal epithelial injury, which increases the areas available for crystal attachment and eventual retention within the kidney [29, 30]. Antioxidant therapy has been used to manage CaOx nephrolithiasis [31, 32]. In previous studies, Selvam reported that vitamin E administration prevented crystal precipitation in the rat kidney [33]. Vitamin E has also been shown to decrease the urinary excretion of oxalate and calcium and restored antioxidant ability in the blood in patients who underwent surgical nephrolithiasis removal [34]. A previous study of antioxidant enzyme levels in lithic rats indicated that almost all the antioxidant enzyme activity was attenuated [35]. In our study, SOD activity was restored, and MDA and 8-IP were increased in EG-treated rats when compared with untreated rats (p < 0.05). After preventive treatment with Fa, SOD activity was restored, and MDA and 8-IP levels declined when compared with those in EG-treated rats (p < 0.05). Particularly in the EG + Pfa660 group, the OS marker levels were approximately normal, demonstrating an optimal antioxidative effect of Fa at the chosen dose. This effect may be due to the flavonoids in the extract, which protect the kidney against the damaging effects of ROS, including peroxynitrite, peroxyl radicals, superoxide, singlet oxygen, and hydroxyl radicals [36].

Based on chemical compositional analysis and bacteriology of metabolic stones, studies have suggested that some metabolic stones, particularly CaOx nephrolithiasis, originate from bacterial infections [37–39]. Among these studies, Sohshang et al. reported that approximately 47% of calculi obtained from 100 kidney stone patients had positive bacterial cultures in stone samples and urine; in addition, *Escherichia coli* was the most common bacteria causing infection [38]. Chutipongtane et al. [40] demonstrated
that CaOx crystal growth and aggregation can be promoted directly by bacteria in vitro and that CaOx stone promotion by bacteria is comparable to fragmented red blood cell membranes, which were recently proposed to promote CaOx stones [41]. Coumarins, bioactive compounds isolated from Fructus Aurantii, demonstrate exceptional antimicrobial activity. In the present study, the inhibition of CaOx crystal growth and aggregation by Fa extract may be due to its antimicrobial activity. In addition, in the in vivo experiments, all of the rats in the EG group (100%) suffered from urinary tract infections, which could have contributed to crystal nucleation [42]. Preventive use of Fa extract containing bioactive compounds that show exceptional antimicrobial activity can significantly reduce urinary tract infection, which may inhibit crystal formation. However, therapeutic treatment with Fa extracts did not significantly improve urinary tract infection, likely because the dose of drug we used did not reach therapeutic levels.

OPN is an important protein component of the stone matrix and is thought to be involved in many pathologic and physiologic processes, including cell migration, adhesion, inflammation, and renal injury [43–45]. OPN is localized in specific sites in the kidney and is mainly restricted to renal epithelial cells in the cortex. However, elevated OPN expression in the renal cortex and medulla was observed after CaOx crystal deposition [25, 45]. Immobilized OPN increases crystal aggregation, and OPN adherence to the surface of collagen granules causes an increase in calcium oxalate crystal adherence and aggregation [46]. Thus, factors that modulate renal OPN expression are expected to regulate the growth and aggregation of crystals in the kidney. In our study, we determined that preventive treatment with Fa

Figure 5: Immunostaining of OPN for each group (Figures 5(a)–5(f)) in kidney tissue (100x magnification).
Figure 6: Western blots for OPN expression. (a) Analysis of OPN protein expression in the SD rat kidney was conducted using conventional western blotting. GAPDH was used as the internal control. The following groups are shown: (1) control group, (2) EG group, (3) EG + TFa220 group, (4) EG + TFa660 group, (5) EG + P Fa220 group, and (6) EG + P Fa660 group. (b) Quantitative densitometric analysis of OPN protein levels. The values are reported as the mean ± SD of three experiments (*p < 0.05 when compared with the control, #p < 0.05 when compared with the EG group, and @p < 0.05 for the EG + P Fa660 group compared with the EG + P Fa220 group).

extract significantly lowered OPN expression compared with EG-treated groups.

In the therapeutic groups, certain urolithiasis-related indicators were improved. However, the number of crystals was not significantly reduced. It is well known that the etiology of CaOx urolithiasis is complex and involves multiple factors. The improvement of certain indicators may not have been sufficient to reduce crystals that were already formed by day 14 in the EG + TFa220 and EG + TFa660 groups. Thus, renal crystal deposition did not significantly improve in the EG + TFa220 and EG + TFa660 groups. However, further studies are needed to confirm this hypothesis.

5. Conclusion

In conclusion, our results indicate that Fa aqueous extract prevents CaOx crystallization and promotes crystal dissolution. The extract also prevents CaOx calculi in the rat kidney, possibly through a combination of increasing antioxidant levels and antimicrobial activity while decreasing urinary stone-forming constituents, OPN expression, and hypercitraturic effects. Further studies are necessary to identify the active components in the extract and the mechanisms responsible for the observed pharmacological activity.

Abbreviations

Fa: Fructus Aurantii
OS: Oxidative stress
SD: Sprague-Dawley
SOD: Superoxide dismutase
ROS: Reactive oxygen species
MDA: Malondialdehyde
MAPK: Mitogen-activated protein kinase
ERK: Extracellular signal-regulated kinase
OPN: Osteopontin
BUN: Blood urea nitrogen
Cr: Serum creatinine
JNK: c-Jun NH2-terminal protein kinase
CaOx: Calcium oxalate
EG: Ethylene glycol.

Conflict of Interests

The authors declare that they have no conflict of interests.

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