INTRODUCTION

Aristolochic acid (AA)-containing compounds have been used as antiviral, antibacterial, anti-inflammatory, anti-neoplastic, and slimming agents [1,2] for a long time. However, these compounds have become notorious after the breakout of interstitial nephritis in Europe-an women who took these compounds to reduce body weight [3]. The first case of aristolochic acid nephropathy (AAN) was reported in Belgium [3], and several cases with irreversible renal toxicity after intake of AA-containing compounds have been reported. More recently, AA has also been identified as a carcinogen responsible for urothelial carcinoma [2,4-6]. Accordingly, the World
Health Organization classified AA as a type 1 carcinogen [2].

Due to both the renal toxicity and potential carcinogenic effects of AA, the Food and Drug Administration of the United States advised caution regarding the consumption of AA-containing compounds in 2000. In Korea, the Korea Food and Drug Administration (KFDA) prohibited herbal ingredients containing AA in 2005. However, even after AA-containing ingredients were prohibited in many countries, cases identified as AAN have been sporadically reported [3,7-15].

A wide range of plants contains AA [1]. Therefore, AA-containing compounds can still be acquired in traditional medicine clinics and through internet markets even though many countries have designated AA-containing compounds as prohibited substances [1,16]. Therefore, it is expected that more cases of AAN have developed in Korea. Herein, we collected cases of AAN by searching past publications in Korea and summarized a total of 16 cases with a review of the literature.

METHODS

Study population
We searched the Research Information Service System, Korean Medical Database, and the Korean Studies Information Service System using the phrases “Chinese herb nephropathy (CHN)” or “AAN.” We reviewed the medical records of all patients with CHN or AAN [11,17-22]. We also reviewed the medical records of patients proven to have AAN in four hospitals from January 2002 to December 2015. This study was approved by the Institutional Review Board of Seoul St. Mary’s Hospital (IRB No. KC16RISI0735) and the Institutional Review Boards of each participating hospital. The informed consent was waived.

Clinical parameters
From the medical records, we obtained clinical information, including age, gender, past medical history, initial symptoms and physical examination, treatment, and clinical course. We also collected hematological and biochemical parameters and histopathologic findings, including hemoglobin, blood urea nitrogen (BUN), creatinine, AA level by phytochemical analysis, urinalysis, and histopathologic findings of the kidney.

Phytochemical analysis and renal biopsy
For the phytochemical analysis, we were able to obtain the remaining herbal medication from six of the patients diagnosed with AAN. The procured medical ingredients were analyzed by HPLC (high-performance liquid chromatography), and the results were expressed as mg/kg.

AA is a mixture of structurally related nitrophenanthrene carboxylic acids that consist of several subtypes, mainly 8-methoxy-6-nitro-phenanthro-(3,4-d)-1,3-dioxolo-5-carboxylic acid (AAI) and 6-nitro-phenanthro-(3,4-d)-1,3-dioxolo-5-carboxylic acid (AAII). Renal biopsy was conducted in all patients at first hospitalization. Renal tissue was

Table 1. Suggested diagnostic criteria for aristolochic acid nephropathy

|         | Impaired renal function (eGFR < 60 mL/min/1.73 m²) plus |
|---------|-------------------------------------------------------|
| Definitive | Any two of the following three criteria               |
|          | 1. Characteristic renal histopathology\(^a\)          |
|          | 2. Ingestion of products containing AA confirmed by phytochemical analysis |
|          | 3. Presence of AA-DNA adducts in renal or urinary tract tissue |
| Probable | Impaired renal function (eGFR < 60 mL/min/1.73 m²) plus |
|          | Only one of the above three criteria plus             |
|          | Urothelial cancer at the time of presentation        |
| Possible | Impaired renal function (eGFR < 60 mL/min/1.73 m²) plus |
|          | No alternative explanation for renal dysfunction plus |
|          | History of use of herbal remedy likely to contain AA but without phytochemical confirmation |

\(^a\)Hypocellular interstitial fibrosis associated with tubular atrophy decreasing from the outer to the inner renal cortex.

eGFR, estimated glomerular filtration rate; AA, aristolochic acid.
prepared for observation with light microscopy using hematoxylin and eosin, periodic acid-Schiff reaction, and silver methenamine and trichrome staining, and was also investigated via electron microscopy and immunofluorescence staining.

Diagnosis of AAN

We diagnosed AAN based on the diagnostic criteria of AAN suggested by Gokmen et al. (Table 1) [6]. The details included impaired renal function, renal histopathology, confirmation of AA by phytochemical analysis, and proof of AA-DNA adducts in renal or urinary tract tissue. If unexplained acute kidney injury (AKI) was accompanied by characteristic renal histology findings (tubulointerstitial nephritis, interstitial fibrosis, and tubular atrophy with preserved glomeruli by light microscopy without other abnormalities), it was considered to be compatible with the histopathology of AAN. Depending on the compatibility of our findings with the diagnostic criteria, patients were divided into definitive, probable, and possible AAN groups.

Statistical analysis

Statistical analyses were performed using SPSS version 18.0 (SPSS Inc., Chicago, IL, USA). Data were presented as the mean ± standard deviation or the count and percentage to the nearest tenth. Chi-square analysis was used for comparisons between the two groups, and a p value less than 0.05 was considered statistically significant.

RESULTS

Classification of patients based on diagnostic criteria

All cases satisfied the diagnostic criteria for AAN. Of the 16 cases, six (37.5%) were classified as definite AAN and 10 (62.5%) were classified as possible AAN. There were no cases of probable AAN.

Occurrence and route of obtaining AA-containing drugs

Fig. 1A shows the occurrence of AAN over a period of 20 years in Korea. When we divided the occurrence of AAN by every 5 years since 1996, 13 patients (81.3%) had developed AAN before 2005, which is the year the KFDA prohibited AA-containing ingredients in herbal medicine. After prohibition, there was no occurrence of AAN between 2006 and 2010. However, three cases of AAN developed between 2011 and 2015. This finding clearly shows that the KFDA probation on the use of AA-containing ingredients in herbal medicine is not sufficient to prevent the occurrence of AAN in Korea. With respect to the route of obtaining AA-containing remedies, 11 cases (68.7%) had received prescriptions from official oriental clinics, and five patients (31.3%) had purchased herbal medicine from a traditional market (Fig. 1B). This finding shows that patients have easy access to AA-containing herbal medicine.

Basic characteristics of patients

Thirteen patients (81.3%) were originally healthy, and three patients (18.7%) had past history of medical dis-
ease (minimal change of disease in two patients, and type 2 diabetes with hypertension in one patient). Of the 16 patients, most were female (n = 14, 87.5%). The occurrence of AAN based on age showed an even distribution, although the fourth and fifth decade groups were shown to be the most vulnerable (Fig. 2A). The reasons for taking oriental medicine were, in order of frequency, slimming (n = 5, 31.3%), control of postpartum edema and breast-feeding (n = 4, 25%), chronic illness (n = 4, 25%), and health promotion (n = 3, 18.8%) (Fig. 2B). These findings suggest that all age groups are exposed to the risks of AAN, and healthy females are at the greatest risk of AAN.

**Initial symptoms and diagnosis at hospitalization**

Fig. 3 shows the symptoms and diagnosis at initial hospitalization of all patients. Initial symptoms at admission were mostly nausea and vomiting (n = 8, 50%) and subsequent weakness (n = 5, 31.3%). Other symptoms that presented irregularly included edema, febrile sense, poor oral intake, flank pain, polydipsia, oliguria, and quadriplegia. All patients were diagnosed with AKI and had significantly elevated serum creatinine at initial hospitalization. Among them, seven patients were diagnosed with Fanconi syndrome and two patients presented with nephrotic syndrome.

![Figure 2. Age distribution of patients with aristolochic acid nephropathy (AAN) and reasons for taking herbal medicine.](image1)

![Figure 3. Symptoms and diagnosis of aristolochic acid nephropathy patients at initial hospitalization.](image2)
Laboratory findings of patients
The main laboratory findings of the patients are summarized in Table 2. All patients presented with anemia and renal dysfunction, including elevated serum creatinine and proteinuria at admission. The mean value of total patients characterized the lower limit of the reference range for serum sodium, potassium, calcium, and phosphate. A larger proportion of total subjects had acidic urine pH, glucosuria, microscopic hematuria, pyuria, and subnephrotic range proteinuria. Seven patients had Fanconi syndrome, which was diagnosed by characteristic laboratory findings such as hypokalemia, alkaline urine pH with metabolic acidosis, glucosuria, and aminoaciduria.

Histologic findings
The renal histopathologic findings were similar in all patients. All patients revealed characteristic findings such as severe tubulointerstitial nephritis, interstitial fibrosis, and tubular atrophy, whereas glomeruli were preserved in light microscopy. In addition, many cases also displayed disruption of the tubular epithelium, interstitial edema, infiltration of inflammatory cells (including lymphocytes and monocytes in the interstitium), and partial acute tubular necrosis. Fig. 4 shows the representative renal histologic findings observed by light microscopy.

Concentration of AA and its clinical implication
We obtained AA-containing remedies from six patients and measured the concentration of AA by phytochemical analysis. The concentrations of AA I, AA II, and cumulative AA I in each patient are summarized in Table 3. The mean levels of AA I and AA II were 2,070.0 mg/kg (range, 1.9 to 8,020.0) and 222.8 mg/kg (range, 1.3 to 740.0), respectively. The mean prescribed duration was 38.6 days (range, 10 to 120). The mean cumulative concentration of AA I was 204,898 mg/kg (range, 67.9 to 962,400). Of the

Table 2. Laboratory findings in patients with aristolochic acid nephropathy

| Parameter                  | Mean ± SD | Range   |
|----------------------------|-----------|---------|
| Hemoglobin, g/dL           | 9.7 ± 2.1 | 6.7–12.2|
| BUN, mg/dL                 | 44.7 ± 32.2| 5.0–129.0|
| Serum creatinine, mg/dL    | 5.1 ± 4.4 | 1.3–13.9 |
| eGFR*, mL/min/1.73 m²      | 20.1 ± 14.1 | 2.8–47.5 |
| Serum sodium, mEq/L        | 138.4 ± 46 | 128–143 |
| Serum potassium, mEq/L     | 3.5 ± 1.2 | 1.5–5.5 |
| Serum chloride, mEq/L      | 111.9 ± 6.9 | 105–126 |
| Uric acid, mg/dL           | 5.1 ± 3.7 | 1.0–11.9 |
| Serum calcium, mg/dL       | 8.2 ± 0.7 | 7.3–9.9 |
| Serum phosphate, mg/dL     | 3.6 ± 2.0 | 0.4–8.1 |
| Total cholesterol, mg/dL   | 175 ± 71 | 124–313 |
| Triglyceride, mg/dL        | 116 ± 62 | 76–226 |
| LDL-C, mg/dL               | 78 ± 11  | 62–87  |
| hs-CRP, mg/dL              | 0.7 ± 0.9 | 0.1–1.8 |
| Urine pH                   | 5.8 ± 0.9 | 5.0–7.5 |
| Urine glucose (+)          | 1.7 ± 1.6 | 0–4    |
| Urine occult blood (+)     | 1.4 ± 1.3 | 0–3    |
| Urine leukocyte (+)        | 1.2 ± 1.3 | 0–3    |
| Proteinuria, mg/day        | 1,882 ± 1,580 | 426.7–6,270.0 |

BUN, blood urea nitrogen; eGFR, estimated glomerular filtration rate; LDL-C, low density lipoprotein cholesterol; hs-CRP, high-sensitivity C-reactive protein.

* eGFR was calculated by the Modification of Diet in Renal Disease equation based on serum creatinine and patient age.
six cases, four with high AA I concentration and cumulative dosage showed rapid progression to ESRD within 6 months after AAN diagnosis.

**Clinical course of patients**

Patients showed diverse clinical courses. Progression to ESRD was observed in seven patients (43.8%), and nine patients (56.2%) showed progression of chronic kidney disease (CKD). The interval between AAN diagnosis and ESRD was 0 to 14 months, and five of seven patients (71.4%) had progressed to ESRD within 6 months of diagnosis. Seven patients were treated with steroids at diagnosis of AAN, but it did not decrease the incidence of ESRD compared with patients who were not treated with steroids (14.3% vs. 44.4%, \( p = 0.23 \)).

**DISCUSSION**

We examined the occurrence of AAN in Korea during the past 20 years. The results of our study show that the outbreak of AAN decreased after the ban of AA-containing ingredients by the KFDA in 2005, but AAN still occurred sporadically after the KFDA prohibition. AA-containing herbal medicine was available through official (oriental medicine clinics) and non-official routes (traditional drug markets). The results of our study revealed that the risk of AAN is still present in Korea even after the legal prohibition of AA-containing herbal medicine, and people are still exposed to AA-containing herbal compounds.

The reason for the high occurrence of AAN in young females is that herbal medicine is thought to be an effective treatment for weight control without complications [23]. Therefore, most reports of AAN are related to herbal remedies used for slimming purposes. Furthermore, AAN due to herbal medicine is also known to occur in a wide variety of remedies for diseases or poor health conditions such as eczema, hepatitis B, liver function enhancement, arthritis, rheumatism, and pain relief [1,2]. The use of Chinese herbal medication to improve health is popular in Korea. For this reason, herbal medicine is favored in a wide range of age groups, as observed in this study (Fig. 2A). Compared to reports in other countries, AAN in Korea was characterized by a high occurrence in females during the postpartum period. The main purpose of herbal medicine in this group was for alleviation of post-partum edema and to improve breast-feeding. Therefore, the possibility that AA exposure to mothers can be transferred to babies via breast feeding should be taken into consideration, and further studies are needed on this point.

Reported cases of AAN were usually asymptomatic on initial presentation, and renal failure was often discovered by routine blood testing [1,23]. Therefore, it is difficult to diagnose AAN if clinicians do not consider the association between AKI and herbal medicine. In our study, we identified some clues to clinically suspect AAN. First, the most important symptoms to suspect AAN were gastrointestinal (GI) symptoms such as nausea and vomiting. Many patients stopped taking herbal medicine due to these symptoms. Second, laboratory findings of Fanconi syndrome were helpful in suspecting AAN. In our study, seven patients showed both AKI and Fanconi syndrome, which is quite unusual in young people. Finally, renal biopsy findings of interstitial ne-

### Table 3. Individual data of AA level in six patients measuring AA level (n = 6)

| Patient no. | AAI, mg/kg | AAII, mg/kg | Prescribed duration, day | Cumulative AAI, mg/kg |
|-------------|------------|-------------|--------------------------|-----------------------|
| 1\(^a\)     | 10.1       | 1.3         | 10                       | 100.6                 |
| 2\(^a\)     | 4,357.0    | 147.3       | 14                       | 60,998.0              |
| 3           | 1.9        | 2.7         | UK                       | UK                    |
| 4\(^a\)     | 8,020.0    | 740.0       | 120                      | 962,400.0             |
| 5           | 4.9        | ND          | 14                       | 67.9                  |
| 6\(^a\)     | 26.4       | ND          | 35                       | 924.0                 |

*a* Patients with progression to end stage renal disease are indicated by a star.

Note that the amount of AA contained in herbal remedies is not consistent, and the amount of exposure to patients is diverse. AA, aristolochic acid; AAI, 8-methoxy-6-nitro-phenanthro-(3,4-d)-1,3-dioxolo-5-carboxylic acid; AAII, 6-nitro-phenanthro-(3,4-d)-1,3-dioxolo-5-carboxylic acid; UK, unknown; ND, not detected.
phritis were strongly suggestive of AAN. In our study, pathologists strongly suggested the diagnosis of AAN based on characteristic histologic findings. Therefore, we suggest that clinicians should suspect AAN in patients presenting with acute GI symptoms, unexplained AKI, and/or Fanconi syndrome with interstitial nephritis on kidney biopsy.

In our study, only 37.5% (6 out of 16 cases) of the cases were able to undergo phytochemical analyses. This small percentage can be explained in two ways. First, many individuals discard the remainder of their herbal medicine once complications arise. Therefore, in many cases, we can suspect AAN but are unable to make a definite diagnosis. Second, measurement of AA in herbal medicine is not available in clinical practice because of technical difficulties and high costs for measuring the concentration of AA. Indeed, in our study, phytochemical analysis was performed at the researcher’s laboratory or at a public toxicology laboratory. Based on phytochemical analysis, we found that the amount of AA contained in herbal remedies is not consistent (ranging from 4.9 to 43,570 mg/kg), and the mean concentration of AA was higher than the dose used for developing acute AAN in experimental animal models [1]. This can explain the wide range of renal dysfunction and the high proportion of acute GI symptoms in patients at admission.

The exact mechanisms of AAN have not been fully understood. As reported in several in vivo and in vitro studies, the proximal tubule is the target of AA. Atrophy and apoptosis is involved in the progression of AA-induced proximal tubular denudation [1,24]. In clinical studies, cumulative AA dose is the only risk factor for the development and progression of renal disease. The Belgian cohort showed that the ingested dose of emerged Aristolochia fangchi was strongly associated with the progression of renal failure [25], and retrospective cohort and case control studies carried out in China and Taiwan have reported median doses of AA associated with renal impairment [26,27]. We also calculated the cumulative doses of AA and found that patients who progressed to ESRD received higher cumulative doses than patients who received lower doses. Taken together, results show that the cumulative dose of AA is a risk factor for the progression of renal failure, and the formation of AA-DNA adducts potentially leads to atrophy and apoptosis in proximal tubules [24].

In terms of clinical outcomes in our study, seven patients (43.8%) showed progression to ESRD, and nine patients (56.2%) had ongoing CKD during the follow-up period. This finding was consistent with previous reports showing that AAN has variant phenotypes with distinct prognosis. In AAN patients with progression to CKD, steroid therapy is recommended for patients with an estimated glomerular filtration rate greater than 20 mL/min/1.73 m² [6,23], and its use is extended to patients with advanced renal dysfunction [28]. In our study, we administered steroid treatment to seven patients, but this did not significantly delay progression to ESRD compared with non-treated patients. However, because our study included a small number of AAN patients with varying degrees of renal dysfunction, this result does not imply that steroid treatment is not effective in delaying the progression of AAN. Another important factor is the presence of Fanconi syndrome. Previous reports have suggested that the presence of Fanconi syndrome predicts favorable clinical outcomes [26]; however, our study showed that 42.9% of AAN patients with Fanconi syndrome progressed to ESRD. This finding suggests that the presence of Fanconi syndrome does not indicate a benign clinical course, and we recommend more frequent follow-up of renal function in AAN patients with Fanconi syndrome.

There are several reasons for the persistent occurrence of AAN after KFDA prohibition of AA in Korea. First, AA-containing compounds were not properly screened. It is strongly suspected that Aristolochia spp. plants were mistakenly identified as Tetrpanax papyrifera and Stephania tetrandra because of similarities in their appearance. Second, Chinese medicines can be easily acquired in traditional markets and on the Internet [1,16], since there is not a clear distribution channel for herbal medicine. Third, oriental medicine in Korea lacks scientific evidence on its safety and effectiveness. The use of herbal remedies is based on knowledge obtained through historic medical texts written in the 17th century. Therefore, the safety and effectiveness of herbal medicine are still unknown. Furthermore, there is a risk of outbreak of AAN because several oriental clinics function as dispensary centers for Chinese medicine. Unfortunately, it is quite difficult to avoid AAN under the current screening system and distribution route, and it might be necessary to perform toxicity tests before the ingredients of...
traditional medicine are distributed to clinics in order to prevent AAN.

Our study has some limitations. First, this study is retrospective. Second, the included number was too small to evaluate the clinical outcome. Third, only six cases were available to measure AA levels in remained herbal medicine. Fourth, data on baseline renal function were not available. Fifth, there is a selection bias since we included only four hospitals. We expect that more cases may be included if a national survey is conducted. In spite of these limitations, this report suggests that there is a possibility of AAN among patients previously diagnosed as having chronic tubulointerstitial nephritis of unknown origin, and more patients will be included if clinicians pay attention to the possibility of AAN.

In conclusion, AA is one of the most important causes of renal tubular damage and interstitial nephritis leading to progressive kidney injury. We believe that AAN is a preventable cause of ESRD via a strong national supervising system for AA-containing herbal medicine in oriental clinics and traditional markets.

### KEY MESSAGE

1. Aristolochic acid (AA)-containing herbal medicine is still available via various routes.
2. Because the definite incidence of aristolochic acid nephropathy (AAN) is unknown and there is a possibility of underdiagnosis, clinicians should suspect AAN in patients presenting with acute gastrointestinal symptoms, unexplained acute kidney injury, and/or Fanconi syndrome with interstitial nephritis.
3. A stronger national supervising system on herbal ingredients and remedies in oriental medicine is needed to prevent AAN.

### Conflict of interest

No potential conflict of interest relevant to this article was reported.

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