Injection of Aluminum Potassium Sulfate and Tannic Acid in the Treatment of Fecal Incontinence: A Single-Center Observational Study

Tatsuya Abe, Masao Kunimoto, Yoshikazu Hachiro, Kei Ohara, Mitsuhiro Inagaki
Department of Proctology, Kunimoto Hospital, Asahikawa, Hokkaido, Japan

Purpose: Perianal injection of bulking agents is an attractive treatment option for patients with mild to moderate fecal incontinence (FI). Various bulking agents have been used for injection therapy, but the optimal injection materials and methods are yet to be standardized. This study aimed to evaluate the effects of injection therapy using aluminum potassium sulfate and tannic acid (ALTA) in the management of FI.

Methods: This study included consecutive patients who underwent ALTA injection therapy for FI at our institution. The procedure was performed with the patient in the jackknife position, under caudal epidural anesthesia. The procedure consisted of a 4-step injection to the 3 main cushions and a multipoint injection to the remaining submucosa of the anal canal.

Results: Seventy-seven patients (mean age, 76 years) were enrolled in the study. The mean Cleveland Clinic incontinence score of 11.9 ± 4.1 at baseline significantly improved to 7.3 ± 5.2 at 3 months following treatment. The mean maximal resting pressure also increased significantly 3 months after the intervention. Postoperative complications were observed in 3 patients (3.9%), and all events were mild. The mean duration of postoperative follow-up was 17.5 months. The cumulative recurrence-free rate at 3 years was 72.4%.

Conclusion: ALTA injection for FI is safe, easy to perform, and provides reasonable mid-term outcomes. Moreover, concomitant anorectal diseases that may be contraindicated by other injectable bulking agents could be treated simultaneously. Therefore, ALTA injection is a promising alternative in the absence of other injectable agents.

Keywords: Fecal incontinence, Anal canal, Sclerotherapy, Aluminum potassium sulfate and tannic acid

INTRODUCTION

Fecal incontinence (FI) is the involuntary loss of liquid or solid stool, which is a social or hygienic problem [1]. FI has a negative effect on the quality of life and may lead to severe social restrictions. The management of FI can be challenging owing to its multifactorial etiology. Conservative treatments such as dietary modifications, pharmacological treatment, and biofeedback are recommended as first-line therapies [2, 3]. However, if conservative measures fail, surgical intervention may be required.

FI can be divided into 3 clinical subtypes: urge incontinence, passive incontinence, and mixed presentation [4]. Urge incontinence is suggestive of poor external anal sphincter (EAS) function, whereas passive incontinence is usually associated with internal anal sphincter (IAS) dysfunction [1]. The IAS has a greater impact on resting anal pressure. Endoanal sonography has enabled increased detection of FI secondary to IAS injury or degeneration [5]. IAS dysfunction is the most common cause of FI in Japan (35%) [6]. Direct surgical repair of IAS defects is difficult, and the outcomes remain poor [7]. Further, the role of sacral neuromodulation in isolated IAS dysfunction has not yet been proven [8]. Therefore, the treatment modalities for IAS dysfunction are...
very limited.

Perianal injection of bulking agents is therefore an attractive alternative for patients with IAS dysfunction. Injectable bulking agents are intended to expand tissue in the anal canal and enhance resting anal pressures, thereby improving FI symptoms [2]. Various bulking agents have been utilized for injection therapy, but the optimal injection materials and methods have yet to be standardized [9]. In a recent Cochrane review of perianal bulking agents, the authors concluded that there is limited evidence to demonstrate the clinical benefit of silicone biomaterials and carbon-coated beads, but dextranomer in hyaluronic acid (non-animal stabilized hyaluronic acid/dextranomer, NASHA Dx) gel improved FI in 52% of patients, and these results were maintained during long-term follow-up [9-11]. NASHA Dx is a relatively new material that is injected into the submucosa of the anal canal, and it is currently the only product approved by the U.S. Food and Drug Administration (FDA) [2]. The Gatekeeper and Sphinkeeper (THD SpA, Correggio, Italy) are the latest self-expandable prostheses that can be implanted in the intersphincteric space of the anal canal and have produced favorable mid-term results [12, 13].

Although the use of perianal injections for treating FI is increasing in Europe and North America, no bulking agent has been approved in Japan thus far [14]. Injection sclerotherapy is an effective office procedure for treating hemorrhoids. Aluminum potassium sulfate and tannic acid (ALTA), an injectable sclerosant based on Xiaozhiling (Xiaozhiling zhusheye, Beijing, China) developed in China in the 1970s [15], has been used in Japan since 2005 [16]. ALTA injection proved to be highly effective even for grade III to IV hemorrhoids, and the effects were comparable with those of hemorrhoidectomy [17, 18]. ALTA solution is not currently approved by the FDA; however, it is used in major Asian countries, including South Korea [19], and has also been applied in the treatment of rectal prolapse and rectocele [20-23].

The active ingredients of ALTA are alum and tannic acid. While alum stimulates a strong local inflammatory reaction, tannic acid suppresses any excessive inflammation that may be caused by the alum and plays a critical role in alleviating any consequential tissue damage. Therefore, ALTA causes sclerosis and retraction of hemorrhoids through fibrosis associated with alum-induced chronic granulomatous inflammation [16]. In case of rectal prolapse or mucosal prolapse, ALTA is also injected into the region without hemorrhoidal cushions [20, 21]. Hence, persistent fibrosis and sclerosis develop circumferentially in the submucosal layer, which may lead to thickening and toughening of the anal canal and rectal wall. Furthermore, we previously reported that when ALTA injection was administered in patients with low resting anal pressure, the maximal resting pressure (MRP) significantly increased following treatment [24]. Therefore, we believe that ALTA injection has a similar mechanism to injectable bulking treatment.

Accordingly, we hypothesized that ALTA injection would be as effective as an injectable bulking agent for the treatment of FI. This study aimed to examine whether ALTA injection could be an alternative to injectable bulking treatment.

**METHODS**

This retrospective observational study analyzed the efficacy and safety of ALTA injections for FI. The study was approved by the Institutional Review Board of Kunimoto Hospital (No. K09-002), and written informed consent was obtained from all the patients.

**Study population**

We included consecutive patients who visited our hospital and underwent ALTA injection therapy for FI between January 2009 and December 2019. All the patients had a history of failed standard conventional treatments for FI, including antidiarrheal agents, anal electrical stimulation, and biofeedback. Patients with urge incontinence with significant EAS defects were excluded from the study. ALTA injection therapy has been approved only for internal hemorrhoids by the Japanese Health Insurance [16]. Therefore, FI patients with internal hemorrhoids (up to grade III) were selected as candidates and were first treated with topical ointments concurrent with conventional treatment for FI. Patients whose FI symptoms persisted even after the conventional treatment were included in the study. Patients who received combination therapy with hemorrhoidectomy and ALTA injection due to hemorrhoid progression despite topical ointment therapy were excluded. Patients with concomitant large rectocele (> 4 cm) or grade V intussusception (full-thickness rectal prolapse) [25] were also excluded from the study because those symptoms may obscure the FI symptoms.

**Clinical assessment**

Preoperative assessment included a questionnaire regarding FI, Cleveland Clinic incontinence score (CCIS) [26], digital rectal examination, and anoscopy, followed by anal manometry, endoanal sonography, and barium defecography, if required. The CCIS was the sum of 5 parameters scored on a scale of 0 (absent) to 4 (daily). A total score of 0 suggested full continence and a score of 20 indicated complete FI [26].

Anal manometry was performed using a 5-mm diameter, 1-channel, solid-state catheter with the patient in the left lateral position. The MRP was recorded using a rapid pull-through technique and was defined as the highest recorded resting pressure. Next, the maximal squeeze pressure (MSP), defined as the highest recorded pressure above the baseline (zero) at any level within the anal canal, was measured. Subsequently, the IAS and EAS were scanned by endoanal sonography using a 7-MHz rotating probe.

**Treatment procedure**

ALTA injection was administered in the operating room under caudal epidural anesthesia with the patients in the jackknife position. In patients taking anticoagulants, local anesthesia was ad-
ministered to avoid spinal epidural hematoma. The ALTA injection method consisted of a 4-step injection for the 3 main hemorrhoids and a multipoint injection for the other area within the anal canal. Using a Z-type proctoscope (Arakawa Seisakujyo, Tokyo, Japan), ALTA solution (Zione, J-Dolph Corp., Osaka, Japan) was slowly injected into 4 parts of each internal pile; submucosa of the superior pole, submucosa and mucosal layer of the central pile, and submucosa of the inferior pole (4-step injection) [17]. For small hemorrhoids, injection to the superior pole or inferior pole was omitted. Next, ALTA (0.5–1.0 mL) was evenly injected into the remaining submucosa of the anal canal at 5 to 10 locations (multipoint injection) [20, 21]. In cases with coexisting rectal intussusception or rectocele, a multipoint injection was performed on the corresponding rectal submucosa. The injected tissues were gently kneaded to evenly distribute the ALTA solution for each injection. Postoperatively, the patients who underwent caudal epidural anesthesia were hospitalized overnight.

Outcome measures
The patients were followed up at 2 weeks, then at 1, 3, and 6 months, and at 1, 2, and 3 years after ALTA injection. Subsequent follow-ups were voluntary. The follow-up comprised a questionnaire based on the CCIS, digital rectal examination, anoscopy, and anal manometry. Recurrence was defined as a re-increase in CCIS to a level at or above the baseline, or cases requiring additional operative interventions for FI. All data were retrospectively collected from the medical records. The CCIS, MRP, and MSP at baseline and 3 months after treatment were compared. The outcome measures were changes in CCIS, MRP, and MSP, postoperative complications, and cumulative recurrence rates.

Statistical analysis
Categorical variables are reported as frequencies and percentages, whereas continuous variables are reported as means and standard deviations (SD). The Wilcoxon signed-rank test was used to comparatively analyze the CCIS, MRP, and MSP values. The results are presented as means ± SD values. The 2-sided significance level was set at 5%. The cumulative recurrence-free rate was assessed using the Kaplan-Meier method. All statistical analyses were performed using EZR software ver. 1.11 (Saitama Medical Center, Jichi Medical University, Saitama, Japan).

RESULTS
From January 2009 to December 2019, ALTA injection therapies were performed in overall 6,236 patients at our hospital. Of these, 4,727 patients (75.8%) underwent combination therapy with ALTA injection and hemorrhoidectomy, 1,509 (24.2%) underwent ALTA injection therapy alone. Of these, 1,509 patients, 77 (5.1%) (63 females; mean age, 76.4 years) met the inclusion criteria and were included in the study. Patient characteristics at baseline are summarized in Table 1. ALTA injection was administered under caudal epidural anesthesia in 74 patients (96.1%) and under local anesthesia in 3 patients (3.9%). The planned injection procedure was successful in all the patients. The mean operative time was 7.8 ± 3.7 minutes, and the mean dose of ALTA per patient was 21.2 ± 6.4 mL.

CCIS was analyzed in 69 patients who completed the questionnaire, and 57 patients (82.6%) had a reduction in CCIS after treatment. Of these, 31 (44.9%) had a ≥50% reduction in CCIS, and 7 (10.1%) regained full continence (CCIS score of 0). The mean CCIS of 11.9 ± 4.1 at baseline significantly improved to 7.3 ± 5.2, 3 months after treatment. All the CCIS sub-scores also significantly improved (Table 2). Changes in manometric data were analyzed in 71 patients who completed the tests. Both the mean MRP and MSP increased significantly 3 months after treatment (Table 3).

Postoperative complications were observed in 3 female patients (3.9%). One patient experienced fecal impaction, 1 had a fever of ≥38°C, and the other patient had perianal dermatitis. All events were mild and ameliorated within days after treatment with sodium bicarbonate suppository, antipyretics, and topical ointments.

Table 1. Baseline patient characteristics (n = 77)

<table>
<thead>
<tr>
<th>Variable</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (yr)</td>
<td>76.4 ± 7.7</td>
</tr>
<tr>
<td>Female sex</td>
<td>63 (81.8)</td>
</tr>
<tr>
<td>Height (cm)</td>
<td>150.3 ± 7.8</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>50.7 ± 8.1</td>
</tr>
<tr>
<td>Vaginal delivery (n = 63)</td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>6 (9.5)</td>
</tr>
<tr>
<td>1–2</td>
<td>34 (54.0)</td>
</tr>
<tr>
<td>≥3</td>
<td>23 (36.5)</td>
</tr>
<tr>
<td>Stool frequency</td>
<td></td>
</tr>
<tr>
<td>&lt; 3 times/wk</td>
<td>12 (15.6)</td>
</tr>
<tr>
<td>≥3 times/wk to ≤ 3 times/day</td>
<td>54 (70.1)</td>
</tr>
<tr>
<td>&gt; 3 times/day</td>
<td>11 (14.3)</td>
</tr>
<tr>
<td>Bristol stool form scale</td>
<td></td>
</tr>
<tr>
<td>Type 1, 2</td>
<td>17 (22.1)</td>
</tr>
<tr>
<td>Type 3–5</td>
<td>49 (63.6)</td>
</tr>
<tr>
<td>Type 6, 7</td>
<td>11 (14.3)</td>
</tr>
<tr>
<td>Clinical subtype</td>
<td></td>
</tr>
<tr>
<td>Urge incontinence</td>
<td>2 (2.6)</td>
</tr>
<tr>
<td>Passive incontinence</td>
<td>51 (66.2)</td>
</tr>
<tr>
<td>Mixed incontinence</td>
<td>24 (31.2)</td>
</tr>
<tr>
<td>Coexisting anorectal disease</td>
<td></td>
</tr>
<tr>
<td>Hemorrhoids (≤ grade III)</td>
<td>77 (100)</td>
</tr>
<tr>
<td>Intussusception (≤ grade IV)</td>
<td>24 (31.2)</td>
</tr>
<tr>
<td>Rectocele (≤ 4 cm diameter)</td>
<td>15 (19.5)</td>
</tr>
</tbody>
</table>

Values are presented as mean ± standard deviation or number (%).
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Of the 77 patients at baseline, 33 were available for follow-up after 12 months and 15 were continued follow-ups after 36 months. The mean duration of postoperative follow-up was 17.5 months (range, 1–84 months). Fig. 1 depicts the Kaplan-Meier plots of the recurrence-free probability for patients after treatment. Cumulative recurrence-free rates (95% confidence interval) at 3 and 5 years were 72.4% (56.7%–83.3%) and 57.0% (33.1%–75.2%), respectively.

DISCUSSION
In this retrospective study of 77 patients, we demonstrated that ALTA injection for FI was safe, easy to perform, and achieved reasonable mid-term outcomes. This was confirmed by improvement in the validated severity index and objective manometric data.

The advantages of ALTA injection for FI are that it can be safely administered even to patients with concomitant anorectal diseases and those taking anticoagulants. ALTA injection has no risk of migration or dissipation like prostheses. Unlike other injectable bulking agents, repeated injections are not required. Furthermore, ALTA injection does not preclude future surgical options because there is no residual biocompatible material.

A Cochrane review of perianal bulking agents analyzed 5 randomized trials involving 382 patients [9]; most patients (n = 206) were derived from the recent placebo control trial by Graf et al. [10]. They reported that 71 of 136 patients (52.2%) who received NASHA Dx had a ≥ 50% reduction in the number of incontinence episodes at 6 months [10]. In a review and meta-analysis of 23 articles (3 randomized controlled trials and 20 case series) in which 11 different bulking agents were used, 19 of the studies (637 patients) used the CCIS as a measure of therapeutic efficacy [27]. The pooled mean CCIS was 12.4 preoperatively and 7.7 at the last follow-up. The weighted mean difference in CCIS was 4.9, and the rate of improvement was 39.5% based on the CCIS [27]. In our study, 45% of patients exhibited a ≥ 50% reduction in CCIS and 10% achieved full continence; the rate of improvement was 38.7% based on CCIS. Therefore, we conclude that ALTA injection has a comparable effect on FI as other injection therapies.

In the above-mentioned review of 23 articles, 16 studies evaluated changes in manometric data after injection therapy. Mean MRP and MSP were the most commonly reported parameters [27]. In terms of change in MSP following injection, mixed results were observed, and no trend was noted. Meanwhile, 7 studies reported a significant increase in MRP during short-term follow-up [27]. Similarly, in our study, the mean MRP increased significantly after injection, whereas the MSP increased modestly.

Most studies suggest that the benefits of bulking agents appear to decline by 6 to 12 months...
[28]. Loss of treatment efficacy is likely related to the migration or dissipation of the bulking agents [9]. Meanwhile, the studies that evaluated the submucosal injection of NASHA Dx demonstrated that significant improvement in FI was sustained for up to 3 years [10, 11]. In our study, the 3-year recurrence-free rate of ALTA injection for FI was as high as 72%. We have vast experience in treating hemorrhoids with ALTA, and the long-term results were good [16, 18]. Therefore, we expect that the long-term results of the ALTA injection for FI will also be favorable.

The limitations of this study include its retrospective, single-institution, and observational design without a control group. The observation period was shorter than that of studies of other injectable bulking agents. Another weakness is that no specific score was used to evaluate the quality of life before and after the treatment. While all patients had hemorrhoids, some had comorbidities such as rectal intussusception or rectocele, which can by themselves cause FI [1, 4]. Consequently, our findings may have included the effects of correcting these comorbidities. Therefore, the results need to be confirmed by randomized controlled trials. Comparative studies with other injectable bulking agents are also required.

Patients with mild to moderate FI who experience mainly passive incontinence without significant EAS defects are suitable candidates for treatment with bulking agents [4, 9]. Most of these patients are elderly and often have comorbidities, such as rectal mucosal prolapse or rectocele [1, 6]. ALTA injection can simultaneously treat concomitant anorectal diseases that may be contraindicated by other injectable bulking agents. In conclusion, ALTA injection for FI is a promising alternative when current injectable bulking agents are not available.

CONFLICT OF INTEREST

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

REFERENCES


