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Accessibility
Gastric antral vascular ectasia: the evolution of therapeutic modalities

Gastric Antral Vascular Ectasia (GAVE) may be an enigmatic source of non-variceal upper GI bleeding associated with various systemic diseases such as connective tissue disorders, liver disease, and chronic renal failure. Successful treatment of GAVE continues to be a challenge and has evolved through the years. Currently, given the rapid response, safety, and efficacy, endoscopic ablative modalities have largely usurped medical treatments as first-line therapy, particularly using argon plasma coagulation. However, other newer ablative modalities such as radiofrequency ablation, cryotherapy, and band ligations are promising. This paper is an overview of GAVE and its various endoscopic and medical therapies.

What is GAVE?

Gastric Antral Vascular Ectasia, or GAVE, can be an obscure cause of upper GI bleeding and was described as early as the 1950s by Rider et al [1], as ‘fiery red’ hypertrophic antral mucosa with scattered areas of bleeding and blood clots, named veno-capillary ectasia. Later studies detailed the endoscopic appearance of GAVE, coining the term “watermelon stomach” to describe the erythematous ectatic vessels in longitudinal stripes along the rugal folds of the antrum [2] or, rarely other parts of the gastric mucosa [3]. There are 2 types of GAVE based on distinctive endoscopic appearances. The classic manifestation consists of this “watermelon” appearance of multiple flat, linear, erythematous strips of ectatic vessels radiating from the pylorus to the antrum. The second type is punctate, where the ectasia manifests as diffuse antral angiofibromas and tends to be more associated with liver cirrhosis [3].

Although GAVE has characteristic endoscopic appearances, it can often be mistaken for other upper GI vascular malformations or sources of GI bleeding. For this reason, diagnostic confirmation is reached through biopsy. Histologically, GAVE appears as dilated, ectatic capillaries in the mucosal and submucosal regions, accompanied by the presence of microthrombi. In addition, there is fibromuscular hyperplasia of the lamina propria in the setting of edema, congestion, and reactive changes of the foveolar epithelium. Generally, very few inflammatory changes are seen. To identify GAVE objectively, a scoring system was devised, which involves the histological factors of ectasia, fibrinolysis, and spindle cell proliferation (Fig. 1).

Most cases of GAVE are associated with liver cirrhosis, autoimmune disease, chronic renal failure, heart disease, diabetes, hypothyroidism, and bone marrow transplantation. GAVE accounts for ~4% of non-variceal upper GI bleeding. Seventy-one percent of non-cirrhotic patients with GAVE are women in their early 70’s, and 75% of cirrhotic patients with GAVE are men in their mid 60’s [2]. Given its association with several common comorbidities, GAVE may become increasingly common in the aging population. The actual etiology of GAVE is unknown, however, multiple hypotheses have been proposed by small studies and case reports. Given the diversity of associated conditions, a singular etiology of GAVE is unlikely.

GAVE with cirrhosis

Much of the research involving the pathophysiology of GAVE has focused on its relationship with portal hypertension and cirrhosis. Although GAVE is found in many cirrhotic patients, no causal relationship has been established, and portal hypertension was not found to play an etiologic role in GAVE. Nevertheless, GAVE is often confused with portal hypertensive gastropathy.
(PHG) since both conditions tend to occur in cirrhotic patients. GAVE and PHG have similar endoscopic appearance, but may be distinguished based on location of the vascular ectasia. GAVE appears mostly in the antrum, whereas PHG is mainly manifest in the gastric fundus. Mucosal biopsies may help distinguish GAVE from PHG in cases that are atypical in presentation, such as in diffuse gastric ectasia. PHG histologically presents as mucosal and submucosal vascular dilatation without associated inflammatory changes. Fibrin thrombi, which are typical findings in GAVE, are generally absent in PHG [5]. These fibrin thrombi can be highlighted with a simple PAS stain with diastase digestion. Immunohistochemical studies suggest that CD61, a platelet marker, is more readily seen in the fibrin thrombi associated with GAVE. This marker was able to diagnose GAVE more accurately, and was positive in 100% of patients with a histological diagnosis of GAVE and in 60% of patients with an endoscopic diagnosis of GAVE. Using CD61, researchers were also able to reclassify incorrectly diagnosed PHG to GAVE and confirm the diagnosis with re-examination of histology. To confirm these findings, researchers used CD31 to determine the mucosal microvessel density, which was found to be significantly higher in cases of GAVE versus PHG ($p < 0.01$) [6]. Despite the similar gross and endoscopic appearance, these two conditions are not the same, and distinction between these two etiologies is important given their differing management strategies. Unlike PHG, GAVE does not respond to a reduction in portal hypertension with the transjugular intrahepatic portosystemic shunt (TIPS) procedure or β blockade [7]. A study has also demonstrated the presence of elevated prostaglandin E2 levels, mainly in the corpus and antrum, in cirrhotic patients with GAVE lesions compared to cirrhotic patients without GAVE, as well as healthy controls. Elevated levels of E2 are thought to have vasodilatory effects, in addition to acid inhibitory effects [8].

**GAVE without cirrhosis**

In non-cirrhotic patients with GAVE, autoimmune disorders and, more specifically, connective tissue diseases are commonly seen. Systemic sclerosis (SSc) is associated with telangiectasias in multiple parts of the GI tract, particularly esophageal (3.9% of SSc patients) or colonic (5.2% of SSc patients), but can also present with severe hemorrhage and anemia secondary to GAVE. A recent retrospective multi-center study noted the presence of endoscopic changes consistent with GAVE in 25% of its 103 subjects with early and severe, diffuse systemic sclerosis [9]. The pathophysiology of GAVE in the SSc population is unknown, but there are two leading hypotheses: autoimmune reaction to gastric vessels, or pathophysiological consequence of gastric dysmotility.

In support of an autoimmune reaction to gastric vessels, anti-RNA polymerase III (RNAP III) antibodies, which are highly specific for SSc, have been noted as present in SSc patients with GAVE. A cross-reaction between specific proteins of the vascular tissue of the gastric mucosa and these antibodies has been speculated to result in GAVE. In case reports, RNAP III antibodies have been described in up to 25% of SSc patients with GAVE [10]. While smaller studies showed RNAP III as a positive predictor of GAVE in SSc, a more recent, larger multi-center study showed no association between the two diseases [9].

Further studies at the molecular level have been conducted to understand the pathophysiology of GAVE. Valdez et al. isolated an RNA helicase from an autoimmune antibody of a patient with GAVE. These RNA helicases are part of the DEXD box family, and have been implicated in pre-mRNA splicing, translation, ribosomal processing, cell growth and development [11]. The group further identified unique epitopes in RNA helicase II/Gu protein specific to GAVE. The GAVE-specific serum was able to recognize...
epitopes near the carboxy-terminus of RHII/Gu, whereas antibodies from patients with other connective tissue disease recognized epitopes of the NH₂-termius. Two additional serum samples from patients with GAVE in this study did not recognize the RHII/Gu antigen [12]. It is currently unclear whether there is a causal relationship of this unique epitope with the symptoms of GAVE, and further studies are needed to understand the pathophysiological role of such antibodies in the autoimmune subtype of GAVE.

Gastric dysmotility has also been touted as an etiology for GAVE. Prolapse or intussusception of the antral mucosa into the pylorus in a chronic, recurrent fashion can result in trauma, causing fibromuscular hyperplasia and vascular ectasia. The discoordinate gastric antral contraction may cause elongation and dilation of mucosal vessels, resulting in the ectatic vessels of GAVE [13]. High levels of gastrin have also been noted in various GAVE cases, which may explain the angiodysplasia. In some studies, GAVE has also been associated with low pepsinogen, and achlorhydria, suggesting a hormonal connection, but conflicting results from other studies have made causal relationships unclear. Although GAVE is also found in many cirrhotic patients, no causal relationship has been established. Vasoactive substances, such as gastrin, SHT-3, and VIP, are secreted by surrounding neuroendocrine cells and may result in malfunction of precapillary sphincters at high levels [14]. This sphincter malfunction can, in turn, result in vasodilatation, ectasia, and a higher propensity for bleeding.

Several treatment modalities have been developed with the main focus on achieving hemostasis, since the underlying pathophysiology of GAVE remains largely unknown. Many patients present with severe gastric bleeding, requiring continuous transfusions. Thus, the first-line therapy is generally endoscopic ablation with medical therapies considered to be adjunctive.

**Endoscopic/surgical treatments**

**APC**

The currently embraced endoscopic treatment modality for GAVE is argon plasma coagulation (APC). APC is a thermoabative method, which causes thermocoagulation using a high frequency current that passes through argon gas. Similar to the YAG laser, APC is able to treat large areas of mucosa per treatment session. However, the perforation risk is lower since there is no direct contact with the mucosa and so it presumably avoids deeper mucosal injury [15]. Many studies have demonstrated the efficacy of APC, however, most have been single center trials with a low number of subjects [16–21]. One of the larger trials involving 50 cirrhotic patients with iron deficiency anemia or melena related to GAVE, found an increased mean hemoglobin of 1.35±0.24g/dl in ~8.5 months of follow-up after the last APC session. Patients were also noted to have undergone a mean of 5.06±1.5 treatment sessions, likely related to the severity of their cirrhotic disease [22]. Leclaire et al. corroborated these findings that cirrhotic patients require more sessions of APC to treat GAVE lesions adequately [23]. An Arabic study showed similar efficacy in 29 patients with endoscopically proven GAVE, with decreased transfusion requirements and an increase in baseline hemoglobin levels. However, the follow-up time was shorter at 3 months post-APC [24]. Most of the early studies on APC showed efficacy in short-term follow-up periods. As a result, additional studies were done to assess long-term efficacy [17,19,25–28]. Nakamura et al. showed that the recurrence-free rate and survival rate after APC declined over time. Cumulative recurrence-free rates were 49.7% after 1 year, 35.5% after 2 years, and 35.5% after 3 years with post-treatment survival rates of 94.4%, 75.8%, and 64.9% at 1, 2, and 3 years, respectively [27]. APC has some drawbacks, most notably, sepsis, antral stenosis, and gastric outlet obstruction as other post-procedure complications [29,30]. Although deeper mucosal injury is deemed less likely, APC has been shown to result in inflammatory or hyperplastic polyps, which can be additional sources of bleeding [31,32]. Financially, APC has a lower initial capital cost, however, the per treatment cost of APC probes is higher given the multiple treatment sessions needed per patient for treatment of GAVE-related hemorrhage [33].

**Radiofrequency ablation**

Radiofrequency Ablation (RFA) is a thermoabative modality, initially used in the treatment of Barrett esophagus. This technique uses high power energy (11–20J) for short periods of time (less than 1s) to ablate superficial mucosal lesions and allows for ablation of the muscularis mucosa, resulting in less damage to the submucosal layer [34]. The BARRx Halo90 system (Covidien, Sunnyvale, CA) has been used for RFA, and is comprised of an ablative device and energy generator. In addition to its use in esophageal procedures, RFA has also been reported in the treatment of lower GI bleeding secondary to radiation proctitis [35,36]. Zhou et al. demonstrated successful use of this modality in patients with lower GI bleeding from ectatic vessels secondary to chronic radiation proctitis, including those patients refractory to other treatment modalities [35]. GAVE presents with a similar clinical complication to radiation proctitis in that it results in severe gastric hemorrhage requiring endoscopic ablative treatment. The favorable outcomes and decreased number of complications reported in studies involving the use of RFA in radiation proctitis make RFA an attractive treatment modality for GAVE. In one study, 6 patients with chronic transfusion-dependent bleeding from GAVE, 4 of whom had failed prior treatment with APC, underwent RFA ranging from 1 to 3 ablative treatment sessions. The mean hemoglobin (Hgb) improved from 8.5 to 10.2 and 5 of the 6 subjects were no longer transfusion-dependent. No complications were noted in the study [37]. A more recent study on 21 patients with GAVE refractory to APC showed similar efficacy and lack of complications 6 months post-treatment with RFA [38]. In addition, a retrospective international study of 8 European centers and 1 U.S. center investigated the use of RFA in patients with GAVE who were mostly refractory to APC (17/18 subjects) and recently presented promising results. Results were significant for treatment of 97% of lesions without adverse effects, as well as decreased blood transfusion requirements 6 months post-procedure compared with transfusion requirements 6 months prior to RFA [39]. It is unclear, however, whether such superficial ablations sufficiently ablate the deeper submucosal vascular network of GAVE, and larger studies with a longer follow-up period are needed. Regardless, the results of these recent studies are promising for the future of RFA in the treatment of GAVE.

**Band ligation**

Endoscopic Band Ligation (EBL) has been used as standard treatment in other GI vascular disorders, such as esophageal varices, hemorrhoids, and Dieulafoy lesions, and has been shown to be safer than its surgical/thermoabative counterparts. The submucosal obliteration of the vascular network is thought to be safer using EBL than other treatment modalities. Given this background, EBL became a viable option for GAVE as well. A study by...
Wells et al. demonstrated the efficacy of EBL versus thermoablative modalities, specifically APC. Patients who were treated using EBL showed a higher rate of bleeding cessation (67% versus 23%) in fewer treatment sessions (1.9 versus 4.7). In addition, the EBL group needed fewer blood transfusions and demonstrated a higher level of baseline hemoglobin after treatment [40]. However, this study reported a higher number of treatment sessions and lower efficacy rate of APC compared with other published studies on APC and may reflect differences in cohort and technique from similar published data. Regardless, many studies and case reports continue to show decreased bleeding recurrence, fewer hospital admissions, and fewer blood transfusions after EBL, as well as better cost-efficacy, making it a more attractive option for health facilities with limited financial resources [41].

**YAG laser coagulation**

Earlier endoscopic treatment of GAVE spanning the 1990s–2000s focused on laser technology, particularly with Nd:YAG lasers which result in thermal destruction of tissue by absorption of laser light. Nd:YAG lasers exhibit deeper mucosal injury to 4–6 mm depth compared with other laser modalities, allowing coagulation of superficial and submucosal vessels without direct contact [42]. In a 2003 long-term retrospective study, 24 patients with GAVE were treated with Nd:YAG laser over an 18-year period with a median of 2 treatments and follow-up median of 55 months (range 9–127) after the last procedure. Twenty patients in this cohort experienced resolution of bleeding, and remained transfusion-free for a median of 16 months [43]. Deeper treatment also increases the risk of perforation, which occurred in 1/24 subjects in this study. Further disadvantages include the higher cost associated with the technology, making other treatment modalities with similar efficacy more attractive.

**Cryotherapy**

Cryotherapy applies extremely cold temperatures to the area of interest to cause thermal destruction or necrosis of the tissue. Initial studies by Kantsevoy et al. demonstrated success of cryotherapy in patients with refractory GI bleeding secondary to GAVE. Of the 7 patients with GAVE treated, 5 (71%) had cessation of bleeding with normal mucosal findings at 6 months post-treatment [44]. Another study focused on patients with GAVE and iron deficiency anemia, who required a mean number of 4.6 units of blood transfusion 3 months prior to treatment. Of 12 enrolled patients, 6 showed a complete response 4 weeks after completion of 3 treatment sessions as defined by improvement in endoscopic appearance, increase in Hgb level, and no requirement for blood transfusions. Five patients showed a partial response, i.e., incomplete ablation with stable Hgb and a reduced number of transfusions [45]. A potential advantage of cryotherapy over APC is the large mucosal areas that can be treated in a 5-min treatment session.

**Heater probe/sclerotherapy/mucosal resection**

Lesser-studied endoscopic modalities include heater probe, sclerotherapy, and mucosal resection. When using a heater probe, the end of the probe is moved along each area of vascular ectasia with continuous coagulation until the bleeding ceases and mucosal blanches. In the late 1980s, one study showed the efficacy of heater probe therapy with 8/10 transfusion-dependent patients no longer requiring transfusions after treatments [46]. A more recent case report also demonstrated, albeit anecdotal, that both heater probe and hot biopsy forceps did not result in the same side effects as APC, namely bleeding, antral scarring, hyperplastic polyps, gastric outlet obstruction, and pneumoperitoneum. The use of heat forceps biopsy was also touted to be more efficient, given its dual ability to biopsy and provide hemostasis when needed during each 20-min session [47]. While these studies have provided some encouraging evidence with regard to the endoscopic heater probe, there is insufficient evidence to suggest that this modality has advantages over other endoscopic modalities and requires more study. Similarly, snare coagulation, which involves sweeping a snare over the mucosal surface, has been effective in a case report, but needs further assessment [48]. Endoscopic mucosal resection (EMR) has been used mostly for resection of superficial dysplastic lesions of the gastrointestinal tract such as Barrett’s esophagus and adenoma. It has also been used as a novel endoscopic modality for the treatment of GAVE in some case reports, with the thought being that some patients tend to hemorrhage with endoscopic modalities such as sclerotherapy and photocoagulation. This tends to occur more readily when larger vessels are present in the area of treatment [49]. While case reports, such as that of Okamoto et al., demonstrate resolution of symptoms after EMR [49], much more investigation is required before it can become a more routine treatment modality.

**Surgical antrectomy**

Surgical intervention with modalities such as antrectomy have been shown in small studies and case reports to be the definitive treatment for GAVE. Antrectomy is often considered for patients whose disease consists of more extensive vascular malformations, which are refractory to medical or endoscopic therapies [50,51]. The morbidity and mortality of the procedure outweigh the benefits, and laparoscopic antrectomy has attempted to decrease the risk involved with a less invasive approach [50]. Unfortunately, the morbidity and mortality associated with abdominal surgical procedures are even higher in patients with severe liver disease, such as cirrhosis, owing to increased bleeding risk from abdominal collateral vessels in the setting of severe portal hypertension [51] (Table 1).

**Medical therapies**

Many medical therapies have been proposed over the years as a non-invasive alternative for the treatment of GAVE-related hemorrhage. Therapies, such as cyclophosphamide, estrogen, progesterone, corticosteroids, tranexamic acid, octreotide, cyproheptadine, and thalidomide, have shown positive results in case reports and small clinical trials, but have not shown sufficient efficacy to function as alternatives to endoscopic modalities. In addition, certain agents, such as estrogen/progesterone, corticosteroids, tranexamic acid, and cyproheptadine may result in unnecessary side effects. As a result, these medical therapies are still considered experimental and are generally not used as standards of care for GAVE-induced hemorrhage [52] (Table 2).

**Conclusion**

GAVE is uncommon, but encountered by most endoscopists as a cause of severe upper gastrointestinal bleeding. There is no consensus for the optimal therapeutic approach. Data reviewed here favor the use of endoscopic ablation over medical treatments, given their more rapid effect and reported success. However, larger and controlled trials are lacking comparing endoscopic to...
### Table 1  Endoscopic treatments.

| References                  | Treatment modality   | n   | Power settings | Mean # sessions | Response rate | Mean Hgb increase | Follow-up duration (months) | Complications                           |
|-----------------------------|----------------------|-----|----------------|-----------------|---------------|-------------------|------------------------------|----------------------------------------|
| Mathou et al. [43]          | Nd:YAG laser         | 24  | 20 – 30 W      | Median 2        | 20/24 (83 %)  | N/A               | Range 9 – 127                | Gastric perforation – 1; Pyloric stenosis – 2 |
| Petrini and Johnston [46]   | Heater probe         | 12  | 4              | 10/12 (83 %)    | 20            | N/A               |                              |                                        |
| Komiyama et al. [47]        | Heater probe         | 1   | 80 W           | 2               | 1/1 (case report) | N/A               |                              | None                                   |
| Sebastian et al. [21]       | Argon plasma coagulation | 12  | 1.5 L/min, 40 W | Median 2 (1 – 5) | 12/12 showed improvement; 2 recurrences at 4 and 9 months | +4.07 | Range 6 – 30 | None                                   |
| Naga et al. [24]            | Argon plasma coagulation | 29  | 2.8 – 4.1 L/min, 60 – 80 W | (1 – 3) | 22/25 CR | +2.7 | 3 | N/A                                   |
| Roman et al. [29]           | Argon plasma coagulation | 21  | 0.81 L/min; 50 – 80 W | 2.81 (1 – 5) | 6/21; 2 recurrences, 11 unrelated deaths, 2 lost to follow-up. | +2.23 | Mean 14.9 (1 – 60.6) | Hematemesis – 2; septicemia – 1 |
| Fuccio et al. [31]          | Argon plasma coagulation | 20  | Median 3 (1 – 10)  | 14/20 CR; 6 recurrences | Mean 28 | Hyperplastic polyps – 3 |
| Wells et al. [40]           | Endoscopic band ligation | 9   | N/A            | 1.9             | 5/9 (56%)    | +2.8             | 10.1 months | Post-procedure nausea/vomiting – 1 |
| Sato et al. [41]            | Endoscopic band ligation | 12  | 3 (range 2 – 4) | 11/12 (91.7 %) | 14.6 months | None                        |
| Gross et al. [37]           | Radiofrequency ablation | 6   | N/A            | 1.7             | 5/6 (83 %) | 1.6              | 2 months | None                                   |
| McCorisk et al. [38]        | Radiofrequency ablation | 21  | N/A            | 4               | 18/21 (86 %) | 2.4              | 6 months | None                                   |

### Table 2  Medical therapy.

| Reference                  | Type of therapy   | n   | Response rate | Duration of follow-up, months | Complications                           |
|-----------------------------|-------------------|-----|---------------|-------------------------------|----------------------------------------|
| Shulz et al., 2009 [53]     | IV Cyclophosphamide | 3   | 3/3 (100 %) CR | 8 – 36                        | N/A                                   |
| Soykan et al., 2003 [54]    | Cyproheptadine    | 1   | 1/1 (100 %)   | 8                             | Delirium in elderly                   |
| Ge et al., 2011 [55]        | Thalidomide       | 78  | 20 (7.4 %); GAVE patients were not distinguished | 8 – 52                        | Leukopenia – 1; somnolence – 1, peripheral edema – 4, bitter taste – 2, thrombopenia – 1, bradycardia – 1, headache – 1, tremor – 1, rash – 1, tinnitus – 1, blurred vision – 1, herpes zoster – 1, pruritis – 1 |
| Nardone et al., 2001 [56]   | Octreotide        | 17; 3 GAVE patients | 1/3 (33 %) CR; 2/3 (66 %) PR | 36 – 48                        | None                                   |
| Barbara et al., 1998 [57]   | Octreotide        | 1   | 0/1 (0 %)     | 24                            | Continuous melena                    |
| Calam et al., 1980 [58]     | Prednisolone      | 1   | 1/1 (100 %)   | 6                             | None                                   |
| Jabbari et al. [2]          | Prednisolone      | 1   | 1/1 (100 %); PR | N/A                           |
| Tran et al., 1999 [59]      | Estrogen/progesterone | 6   | 1/6 (16.7 %) no response | 3 – 12                        | Gynecomastia – 2, metrorrhagia – 1    |
| Moss et al., 1992 [60]      | Estrogen/progesterone | 1   | 1/1 (100 %)   | 12                            | Cyclical uterine bleeding            |
Competing interests: None

References

1 Rider JA, Klotsz AP, Kirsner JB. Gastritis with veno-capillary ectasia as a source of massive gastric hemorrhage. Gastroenterology 1953; 24: 118 – 123

2 Jabbabi M, Cherry R, Lough JO et al. Gastric antral vascular ectasia: the watermelon stomach. Gastro Endoscopy 1984; 87: 1165 – 1170

3 Stotzer PO, Willen R, Kilander AF. Watermelon stomach: not only an antral disease. Gastroenterology Endosc 2002; 55: 897 – 900

4 Gilliam JH, Geisinger KR, Wu WC et al. Endoscopic biopsy is diagnostic in gastric antral vascular ectasia. The “watermelon stomach.” Dig Dis Sci 1989; 34: 885 – 888

5 Poyen JL, Cails P, Voigt JF et al. Severe portal hypertensive gastroopathy and antral vascular ectasia are distinct entities in patients with cirrhosis. gastroenterology 1995; 108: 138 – 144

6 Westerhoff M, Tretiakova M, Hovan L et al. CD61, CD31, and CD34 improve diagnostic accuracy in gastric antral vascular ectasia and portal hypertensive gastropathy: An immunohistochemical and digital morphometric study. Am J Surg Pathol 2010; 34: 494 – 501

7 Barak KW, Lee SS, Beck PL. Portal hypertensive gastropathy and gastric antral vascular ectasia (GAVE) syndrome. Gut 2001; 49: 866 – 872

8 Soperas E, Perez Ayuso RM, Poca E et al. Increased gastric PGE2 biosynthesis in cirrhotic patients with gastric vascular ectasia. Am J Gastroenterol 1990; 85: 138 – 144

9 Hung EW, Mayes MD, Sharif R et al. Gastric antral vascular ectasia and its clinical correlates in patients with early diffuse systemic sclerosis in the SCOT trial. J Rheumatol 2013; 40: 455 – 460

10 Ceribelli A, Cavazzana I, Airò P et al. Anti-RNA polymerase III antibodies as a risk marker for early gastric antral vascular ectasia (GAVE) in systemic sclerosis. J Rheumatol 2010; 37: 1544

11 Valdez BC, Henning D, Busch RK et al. A nucleolar RNA helicase recognized by autoimmune antibodies from a patient with watermelon stomach disease. Nucleic Acids Res 1996; 24: 1220 – 1224

12 Garcia MC, Zhou J, Henning D et al. Unique epitopes in RNA helicase II/Gu protein recognized by serum from a watermelon stomach patient. Mol Immunol 2000; 37: 351 – 359

13 Sollam H, McNeary TA, Chen JD. Systematic review: pathophysiology and management of gastrointestinal dysmotility in systemic sclerosis (scleroderma). Aliment Pharmacol Ther 2006; 23: 691 – 712

14 Lowes JR, Rode J. Neuroendocrine cell proliferations in gastric antral vascular ectasia. Gastroenterology 1989; 97: 207 – 212

15 Ripoll C, Garcia-Tsao G. The management of portal hypertensive gastropathy and gastric antral vascular ectasia. Dig Liver Dis 2011; 43: 345 – 351

16 Yew BS, Ng KY, Ang DS et al. Gastric antral vascular ectasia successfully controlled by argon plasma coagulation. Ann Acad Med Singapore 2007; 36: 702 – 703

17 Yusoff I, Brennan F, Ormonde D et al. Argon plasma coagulation for treatment of watermelon stomach. Endoscopy 2002; 34: 407 – 410

18 Herrera S, Borradori M, Llach J et al. The beneficial effects of argon plasma coagulation in the management of different types of gastric vascular ectasia lesions in patients admitted for GI hemorrhage. Gastrointest Endosc 2008; 68: 440 – 446

19 Kwan V, Bourke MJ, Williams SJ et al. Argon plasma coagulation in the management of symptomatic gastrointestinal vascular lesions: experience in 100 consecutive patients with long-term follow-up. Am J Gastroenterol 2006; 101: 58 – 63

20 Chiu CY, Lu LS, Wu KL et al. Comparison of argon plasma coagulation in managing of upper gastrointestinal angiodyplasia and gastric antral vascular ectasia hemorrhage. BMC Gastroenterol 2012; 12: 67

21 Sebastian S, McLaughlin R, Qasim A et al. Endoscopic argon plasma coagulation for the treatment of gastric antral vascular ectasia (watermelon stomach): long-term results. Dig Liver Dis 2004; 36: 212 – 217

22 Bhatti MA, Khan AA, Alom A et al. Efficacy of argon plasma coagulation in gastric vascular ectasia in patients with liver cirrhosis. J Coll Physicians Surg Pak 2009; 19: 219 – 222

23 Leclere S, Ben-Soussan E, Antonietti M et al. Bleeding gastric vascular ectasia treated by argon plasma coagulation: a comparison between patients with and without cirrhosis. Gastrointest Endosc 2008; 67: 219 – 225

24 Naga M, Esmae S, Nagui M et al. Long-term effect of argon plasma coagulation (APC) in the treatment of gastric antral vascular ectasia (GAVE). Arab J Gastroenterol 2011; 12: 40 – 43

25 Probst A, Scheuvel R, Wienbeck M. Treatment of watermelon stomach (GAVE syndrome) by means of endoscopic argon plasma coagulation (APC): long-term outcome. Z Gastroenterol 2001; 39: 447 – 452

26 Alves DM, Sakai P, Oliveira CV et al. Watermelon stomach: clinical aspects and treatment with argon plasma coagulation. Arq Gastroenterol 2000; 36: 191 – 195

27 Nakamura S, Mitsuanga A, Kontshi H et al. Long-term follow up of gastric antral vascular ectasia treated by argon plasma coagulation. Dig Endosc 2006; 18: 128 – 133

28 Shibukawa G, Irisawa A, Sakamoto N et al. Gastric antral vascular ectasia (GAVE) associated with systemic sclerosis: relapse after endoscopic treatment by argon plasma coagulation. Intern Med 2007; 46: 279 – 283

29 Raman S, Saurin JC, Dumortier J et al. Tolerance and efficacy of argon plasma coagulation for controlling bleeding in patients with typical and atypical manifestations of watermelon stomach. Endoscopy 2003; 35: 1024 – 1028

30 Farooq FT, Wong RC, Yang P et al. Gastric outlet obstruction as a complication of argon plasma coagulation for watermelon stomach. Gastrointest Endosc 2007; 65: 1090 – 1092

31 Fuccio L, Zaggari RM, Serrantini M et al. Endoscopic argon plasma coagulation for the treatment of gastric antral vascular ectasia-related bleeding in patients with liver cirrhosis. Digestion 2009; 79: 143 – 150

32 Baudet JS, Salata H, Soler M et al. Hyperplastic gastric polyps after argon plasma coagulation treatment of gastric antral vascular ectasia (GAVE). Endoscopy 2007; 39: E320

33 Rosenfeld G, Enns R. Argon photocoagulation in the treatment of gastric antral vascular ectasia and radiation prophitis. Can J Gastroenterol 2009; 23: 801 – 804

34 Maguruma N, Okamoto K, Kimura T et al. Endoscopic ablation therapy for gastrointestinal superficial neoplasia. Dig Endosc 2012; 24: 139 – 149

35 Zhou C, Adler DC, Becker L et al. Effective treatment of chronic radiation prophitis using radiofrequency ablation. Therap Adv Gastroenterol 2009; 2: 149 – 156

36 Rustrup T, Meshimo H. Endoscopic management of chronic radiation prophitis. World J Gastroenterol 2011; 17: 4554 – 4562

37 Grass SA, Al-Haddad M, Gill KR et al. Endoscopic mucosal ablation for the treatment of gastric antral vascular ectasia with the HALO90 system: a pilot study. Gastrointest Endosc 2008; 67: 324 – 327

38 McGovern T, Krishnam K, Keefer L et al. Radiofrequency ablation for refractory gastric antral vascular ectasia (with video). Gastrointest Endosc 2013; 78: 584 – 588

39 Dray X, Repici A, Gonzalez P et al. 1040 Radiofrequency Ablation Treatment of Gastric Antral Vascular Ectasia: Results From an International Collaborative Study. Gastrointest Endosc 2013; 77: AB180

40 Wells CD, Harrison ME, Gurudu SR et al. Treatment of gastric antral vascular ectasia (watermelon stomach) with endoscopic band ligation. Gastrointest Endosc 2008; 68: 231 – 236
41 Sato T, Yamazaki K, Akaike J. Endoscopic band ligation versus argon plasma coagulation for gastric antral vascular ectasia associated with liver diseases. Dig Endosc 2012; 24: 237 – 242
42 Rodriguez SA, Douglas G. American Society for Gastrointestinal Endoscopy Technology Committee. Mucosal ablation devices. Gastrointest Endosc 2008; 68: 1031 – 1042
43 Mathoung NG, Lovat LB, Thorpe SM et al. Nd:YAG laser induces long-term remission in transfusion-dependent patients with watermelon stomach. Lasers Med Sci 2004; 18: 213 – 218
44 Kantsevoy SV, Cruz-Correa MR, Vaughan CA et al. Endoscopic cryotherapy for the treatment of bleeding mucosal vascular lesions of the GI tract: a pilot study. Gastrointest Endosc 2003; 57: 403 – 406
45 Cho S, Zanati S, Yong E et al. Endoscopic cryotherapy for the management of gastric antral vascular ectasia. Gastrointest Endosc 2008; 68: 895 – 902
46 Petrini JL Jr, Johnston JH. Heat probe treatment for antral vascular ectasia. Gastrointest Endosc 1989; 35: 324 – 328
47 Komiyama M, Fu K, Morimoto T et al. A novel endoscopic ablation of gastric antral vascular ectasia. World J Gastrointest Endosc 2010; 2: 298 – 300
48 Chong VH. Snare coagulation for gastric antral vascular ectasia ablation. Gastrointest Endosc 2009; 69: 1195
49 Okamoto T, Okayama Y, Hiroi M et al. Gastric vascular ectasia treated by endoscopic mucosal resection. Dig Endosc 2002; 14: 9 – 11
50 Sherman V, Klasseen DR, Feldman LS et al. Laparoscopic antrectomy: a novel approach to treating watermelon stomach. J Am Coll Surg 2003; 197: 864 – 867
51 Spahr I, Villeneuve JP, Dufresne MP et al. Gastric antral vascular ectasia in cirrhotic patients: absence of relation with portal hypertension. Gut 1999; 44: 739 – 742
52 Fuccio L, Mussetto A, Laterza L et al. Diagnosis and management of gastric antral vascular ectasia. World J Gastrointest Endosc 2013; 51: 6 – 13
53 Schulz SW, O’Brien M, Maqsood M et al. Improvement of severe systemic sclerosis-associated gastric antral vascular ectasia following immunosuppressive treatment with intravenous cyclophosphamide. J Rheumatol 2009; 36: 1653 – 1656
54 Soykan I, Toruner M, Idilman R et al. Reversal of iron deficiency anemia in a patient with gastric antral vascular ectasia treated with cyproheptadine. J Clin Gastroenterol 2003; 36: 183 – 184
55 Ge ZZ, Chen HM, Gao YJ et al. Efficacy of thalidomide for refractory gastrointestinal bleeding from vascular malformation. Gastroenterology 2011; 141: 1629 – 1637 e1 – e4
56 Nardone G, Rocco A, Balzano T et al. The efficacy of octreotide therapy in chronic bleeding due to vascular abnormalities of the gastrointestinal tract. Aliment Pharmacol Ther 1999; 13: 1429 – 1436
57 Barbera G, De Giorgio R, Salvio LB et al. Unsuccessful octreotide treatment of the watermelon stomach. J Clin Gastroenterol 1998; 26: 345 – 346
58 Calam J, Walker RJ. Antral vascular lesion, achlorhydria, and chronic gastrointestinal blood loss: response to steroids. Dig Dis Sci 1980; 25: 236 – 239
59 Tran A, Villeneuve JP, Bilodeau M et al. Treatment of chronic bleeding from gastric antral vascular ectasia (GAVE) with estrogen-progestrone in cirrhotic patients: an open pilot study. Am J Gastroenterol 1999; 94: 2909 – 2911
60 Moss SF, Ghosh P, Thomas DM et al. Gastric antral vascular ectasia: maintenance treatment with oestrogen-progesterone. Gut 1992; 33: 715 – 717