Gastric varices: Classification, endoscopic and ultrasonographic management

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INTRODUCTION

Gastroesophageal varices have been seen in approximately 50% of patients with cirrhosis of the liver. Their presence correlates with the severity of liver disease. While only 40% of Child A patients has varices, they are present in 85% of Child C patients. Variceal hemorrhage occurs at a yearly rate of 5-15%, and 6-week mortality after variceal hemorrhage is about 20%. In general, variceal bleeding ceases spontaneously in 40-50% of patients, but incidence of early rebleeding ranges between 30% and 40% within first 6 weeks, and about 40% of all rebleeding episodes occur within the first 5 days.

Gastric varices (GV) bleed less frequently than esophageal varices and are responsible for 10-30% of all variceal hemorrhages. However, gastric variceal bleeding tends to be more severe with higher mortality. In addition, a high proportion of patients, around 35-90%, rebleed after spontaneous hemostasis.

New endoscopic treatment options and interventional radiological procedures have broadened the therapeutic armamentarium for GV. This review provides an overview of the classification and pathophysiology of GV, which have direct consequences for management; an introduction to current endoscopic and interventional radiological management options for GV.

CLASSIFICATION OF GASTRIC VARICES

There are three types of classification commonly used for GV.
1. Sarin’s classification
2. Hashizome classification
3. Arakawa’s classification.

Most commonly used classification is Sarin’s classification of GV.

SARIN’S CLASSIFICATION

Gastric varices are categorized into four types based on the relationship with esophageal varices, as well as by their location in the stomach (Figure 1). Varieties include:

a. Gastroesophageal varix (GOV) type 1: Extension of esophageal varices along lesser
d. Gastroesophageal varix type 2: Extension of esophageal varices along great
c. Isolated gastric varix (IGV) type 1 and

Gastroesophageal varix type 1 is the most common type, accounting for 74% of all GV. However, the incidence of bleeding is highest with IGV type 1, followed by GOV type 2. Overall, the most important predictor of hemorrhage is the size of varices, with the highest risk of first hemorrhage (15%/year) occurring in patients

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with large varices. Other predictors of hemorrhage are decompensated cirrhosis (Child B or C) and the endoscopic presence of red wale marks.

HASHIZOME CLASSIFICATION

It is based on clinically significant endoscopic findings, and particularly from the viewpoint of findings associated with the lightly risk of rupture, as in the classification of esophageal varices [Figure 2]. Thus, endoscopic findings of GV were classified according to their form, location, and color. The form was classified into three types:

a. Tortuous (F1).
b. Nodular (F2).
c. Tumorous (F3).

The location was classified into five types and depends on hemodynamic factors:

a. Anterior (La).
b. Posterior (Lp).
c. Lesser curvature (Ll).
d. Greater curvature (Lg) of the cardia and.
e. Fundic area (Lf).

The colors can be classified in (a) white (Cw) or (b) red (Cr). The glossy, thin-walled focal redness on the varix was defined as red color spot (RC spot). The Hashizume group reported that the RC spot and larger forms were related to a significantly higher risk of gastric variceal bleeding. The form and location of Hashizume classification are shown in Figure 2.

Arakawa’s classification type I: Branches within the stomach wall are very few, and the supplying vessel, varix and draining vessel form a single continuous vein of a nearly unchanged caliber.

- Type Ia: A single supplying vessel forms a fundic varix.
- Type Ib: Plural supplying vessels join and form a varix that drains into a single draining vessel.

Vascular anatomy

To achieve best results of treatment and at the same time minimizing the complications it is very important to understand relevant vascular anatomy. This holds true for all interventional modalities of treatment likely endoscopic, endoscopic ultrasound (EUS) and radiological management of GV. GV drain into the systemic vein via the esophageal-parasophageal varices (gastroesophageal venous system), the inferior phrenic vein (IPV) (gastrophrenic venous system), or both. These drainage types generally correspond to the classification system of Sarin et al. GOV1 drains via esophageal and parasophageal varices, IGV1 drains via the left IPV, and GOV2 drains via both esophageal varices and the IPV. GV form at the hepatopetal collateral pathway that develops secondary to localized portal hypertension and drain via the gastric veins, thereby corresponding with IGV2 [Figure 3].

Gastroesophageal venous system

Esophageal venous plexus which normally lies at the lower end of esophagus Anastomoses with tributaries of the left gastric vein within and around the gastric wall. The esophageal vein drains via the gastroesophageal venous system into the superior vena cava (SVC). GOVs develop at this anastomosis between the left gastric vein and the azygos vein due to portal hypertension.

Gastrophrenic venous system

The gastric veins lie in the posterosuperior part of the gastric wall which have the potential to Anastomose with the IPV at the bare area of the stomach (gastrophrenic ligament). The majority of IGVs form in a large portosystemic venous shunt that develops based on this potential anastomosis between the gastric vein and the left IPV due to portal hypertension. The left IPV terminates either (a) inferiorly into the left renal vein (forming a gastrorenal shunt), often together with the left adrenal vein, or (b) transversely into the inferior vena cava (IVC) (forming a gastrocaval shunt. The proximal portion of the left IPV runs inferiorly to the
diaphragm. Some of the peripheral branches run superiorly to the diaphragm and supply the superior surface of the muscular diaphragm; one of these peripheral branches anastomoses with the left pericardiophrenic vein. The left IPV can also communicate with other peridiaphragmatic and retroperitoneal veins, including the subcostal and intercostal veins, small anastomotic veins to the right IPV, and adrenal vein or with the azygos venous system. Therefore, GV can also drain through these communications. Potential communication between these venous systems and the pulmonary vein has been demonstrated in anatomic studies and a small number of cases as one of the collateral pathways that develops as a result of obstruction of the SVC and portal hypertension.

Modern imaging by computed tomography scan
The following venous systems are important to understand the venous anatomy of GV on multi-detector computed tomography (CT) scan [Figure 4].

Gastroesophageal venous system
On oblique coronal reformatted multidetector CT images one can see that GOVs usually receive blood from the left gastric vein, which runs in the submucosal layer of the stomach along the lesser curvature and continues directly to esophageal varices beyond the esophagocardiac junction, finally joining the azygos and hemiazygos veins.

Gastrophrenic venous system
Gastrorenal shunt
A gastrorenal shunt is the most common drainage route of IGVs (80-85% of cases), running inferiorly and terminating in the left renal vein, often together with the left adrenal vein. The shunt frequently communicates with other phrenic or retroperitoneal drainage veins. Duplication or fenestration of gastrorenal shunts is rare.

Gastrocaval shunt
In 10-15% of cases gastrocaval shunt forms main drainage route for IGVS. It consists of the left IPV passing transversely in front of the esophageal hiatus below the diaphragm and terminating in the IVC (60% of cases) or the left hepatic vein (40%). The proximal portion of the shunt runs inferiorly to the diaphragm, whereas the peripheral branches run superiorly to the diaphragm. The peripheral branch of the left IPV communicates with other peridiaphragmatic veins; thus, gastrocaval shunts are always associated with multiple accessory drainage veins around the diaphragm.

Pericardiophrenic vein
The left pericardiophrenic vein is often seen with a gastrorenal or gastrocaval shunt at multidetector CT as an accessory drainage vein from GV. In a few cases (5%), it can serve as a main drainage route of GV.

The left pericardiophrenic vein anastomoses with the left IPV at the cardiac apex. It runs superiorly along the pericardium and in the left superior mediastinum then terminates into the left brachiocephalic vein. It may join the left ITV or left superior costal vein.

Other drainage routes
These small veins are often difficult to identify at multidetector CT due to their small Caliber. These veins are as follows:

1. The veins running in the thoracic wall, diaphragm, and retroperitoneum are often observed as accessory drainage routes of GV. The subcostal-intercostal veins communicate anteriorly with the ITV and posteriorly with the azygos vein.
2. There are small retroperitoneal and peridiaphragmatic veins, including an anastomotic vein between the bilateral IPVs, renal capsular vein, and unnamed retroperitoneal veins, that communicate with the gastrorenal shunt and the hemiazygos vein.
Combined drainage via gastroesophageal and gastrophrenic venous systems

In Sarin’s classification of GV, GOV 2 drains via both gastroesophageal and gastrophrenic venous systems and account for 15% of GV\(^[17]\) in this type of gastric varix, the anteromedial (cardiac) part of the varix is supplied by the left gastric vein and drains via the esophageal varices into the gastroesophageal venous system. The posterior (fundal) part of the varix is often supplied by the posterior gastric vein or short gastric vein and drains via the gastrophrenic venous system. In the balloon-occluded retrograde transvenous obliteration (BRTO) technique, sclerosant is injected via the gastrophrenic venous drainage system into the GV. Therefore, small parts of GV draining via the esophageal varices may remain after BRTO when the sclerosant does not fill or stagnate in them. This is called incomplete eradication and in such a scenario the high-pressure anteromedial part needs transhepatic embolization approach or endoscopic band ligation.

Management

a. Endoscopic treatment modalities for gastric variceal bleeding.
1. Gastric variceal sclerotherapy (GVS).
2. Gastric variceal obturation (GVO) with glue.
3. Gastric variceal band ligation (GVL) with or without detachable snares.
4. Thrombin injection (bovine or human).
5. Combined endoscopic therapy.

b. Radiologic intervention - transjugular intrahepatic portosystemic shunt (TIPS) and BRTO.

gastric variceal sclerotherapy

The endoscopic sclerotherapy has been less effective in the treatment of gastric variceal bleeding and eradication of GV as against esophageal varices where endoscopic sclerotherapy is one of the effective modes of treatment.\(^[12,18]\)

Because of the high volume of blood flow through GV compared with EV, resulting in rapid flushing away of the sclerosant in the bloodstream. GVS typically requires larger volumes of sclerosant than for EV\(^[7,19]\) and fundal varices (GOV2 and IGV1) because of the low rate of primary hemostasis, the low success rate for secondary variceal eradication, and the high rate of rebleeding and complications. Results of certain important trials on GVS are shown in Table 1.

Gastric variceal obturation

Tissue adhesive such as N-butyl-2-cyanoacrylate, which is a monomer that rapidly undergoes exothermic polymerization on contact with the hydroxyl ions present in water, has been used for Gastric variceal obturation.

METHODS

After puncturing the varix lumen with the needle, cyanoacrylate is injected in 1-1.5 mL aliquots by using normal saline or sterile water (about 0.8-1.0 mL, equal to the dead space) to flush the glue into the varix. As the needle is withdrawn from varix, a steady stream of the flush solution is aimed at the puncture site. Additional glue is injected until the varix is "hard" to palpation.\(^[25]\)

Weeks to months after the injection, the mucosa overlying the glue cast sloughs off and the plug is extruded into the stomach. Initial hemostasis rates of over 90% can be achieved.

Table 1: GVS in the management of gastric variceal bleeding: Comparison from different studies

<table>
<thead>
<tr>
<th>Authors</th>
<th>Agent (%)</th>
<th>n</th>
<th>Success (%)</th>
<th>Rebleeding (%)</th>
<th>Complications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gimson et al., 1991</td>
<td>EO/glue</td>
<td>41</td>
<td>40</td>
<td>16</td>
<td>Ulcer 29%, perforation</td>
</tr>
<tr>
<td>Oho et al., 1995</td>
<td>EO (5)</td>
<td>24</td>
<td>67</td>
<td>25</td>
<td></td>
</tr>
<tr>
<td>Chang et al., 1996</td>
<td>STD (1.5)</td>
<td>25</td>
<td>80</td>
<td>70</td>
<td>Ulcer 30%</td>
</tr>
<tr>
<td>Chang et al., 1996</td>
<td>GW (50)</td>
<td>26</td>
<td>92</td>
<td>30</td>
<td>Ulcer 30%</td>
</tr>
<tr>
<td>Sarin et al., 1997</td>
<td>AA (95)</td>
<td>18</td>
<td>67</td>
<td>34</td>
<td>Ulcer 100%</td>
</tr>
<tr>
<td>Ogawa et al., 1999</td>
<td>EO (5)</td>
<td>21</td>
<td>81</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>Sarin et al., 2002</td>
<td>AA (95)</td>
<td>8</td>
<td>62</td>
<td>25</td>
<td></td>
</tr>
</tbody>
</table>

EO = Ethanolamine oleate; STD = Sodium tetradecyl; GW = Glucose water; AA = Acetic acid; GVS = Gastric variceal sclerotherapy
Complications
Complications are mainly thrombotic in nature and include cerebral embolization and stroke, portal vein embolization, splenic infarction, coronary emboli, with a series demonstrating fatal and nonfatal pulmonary emboli in up to 5% of cases. Use of undiluted cyanoacrylate was shown to be safe and effective and also to be associated with fewer complications, in contrast to the diluted form.\(^{[26,27]}\)

- Several recent long-term studies also reported hemostasis rates of 90% and low rebleeding rates of 15% to 30% with cyanoacrylate injection, with 1-3 injections needed to achieve obliteration, with higher eradication rates for GOV1 and GOV 2 than IGV.\(^{[28,29]}\) Even in young infants, the use of cyanoacrylate glue is safe and effective for the treatment of gastric variceal bleed.\(^{[30]}\)

At present, it is clear that GVO using tissue adhesives has high efficacy and safety for the control of acute GV bleed and for the prevention of GV rebleeding and is the treatment of choice.\(^{[31,32]}\)

Results of certain trials on GOV are shown in Table 2.

Gastric variceal band ligation
Variceal band ligation (VBL) is an established treatment modality for the prevention of esophageal variceal bleeding control of active bleeding and rebleeding. However for GV GVL is not the 1st choice of treatment and evidence for the use of GVL for acute gastric variceal bleeding is mixed, and at best GVL is a second alternative therapy to tissue adhesives. Although initial hemostasis may be achieved with GVL, the main disadvantage has been a high rate of rebleeding, probably from feeding vessels. Repeat endoscopy and VBL is thus necessary at 1- to 2-weekly intervals until eradication of varices or until only small residual varices remain.

In five uncontrolled studies\(^{[33-35]}\) active GV bleed was controlled with 100% success with a very low rebleeding rate of 10-20%, and also the GV was obliterated in all cases.

Results of some trials of GVL are shown in Table 3.

Thrombin
Thrombin which is commercially available as a sterile lyophilized powder principally affects hemostasis by converting fibrinogen to a fibrin clot. A 5-mL solution of thrombin containing 1000 units/mL of thrombin will clot a liter of blood in <60 s. In a patient with bleeding GV, thrombin is reconstituted, and 1 mL aliquot is injected into the bleeding varix.\(^{[26,37]}\)

1. Yang et al.\(^{[38]}\) retrospectively evaluated the use of human thrombin in 12 patients with GV bleeding. Immediate hemostasis was achieved in all patients in whom there was active bleeding from GVs at the time of endoscopy \((n = 6)\). There were no immediate allergic reactions, thromboembolic complications or rebleeding. The results appeared promising, with immediate hemostasis achieved in 70% patients and with no patient having rebleed from GV over a follow-up period of 8 months.

2. Ramesh et al.\(^{[39]}\) showed that endoscopic treatment of bovine thrombin is effective in 92% of patients with

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### Table 2: GVO in the management of gastric variceal bleeding: Comparison from different studies

<table>
<thead>
<tr>
<th>Authors</th>
<th>Study design</th>
<th>n</th>
<th>Follow-up (month)</th>
<th>Hemostasis (%)</th>
<th>Rebleeding (%)</th>
<th>Mortality (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Seewald et al., 2008</td>
<td>Retro</td>
<td>131</td>
<td>60</td>
<td>100</td>
<td>17</td>
<td>47</td>
</tr>
<tr>
<td>Fry et al., 2008</td>
<td>Retro</td>
<td>33</td>
<td>9</td>
<td>88</td>
<td>15</td>
<td>18</td>
</tr>
<tr>
<td>Cheng et al., 2007</td>
<td>Retro</td>
<td>635</td>
<td>3-115</td>
<td>95</td>
<td>8</td>
<td>7</td>
</tr>
<tr>
<td>Joo et al., 2007</td>
<td>Retro</td>
<td>85</td>
<td>24</td>
<td>98</td>
<td>29</td>
<td>31</td>
</tr>
<tr>
<td>Kim et al., 2006</td>
<td>Pros</td>
<td>86</td>
<td>11</td>
<td>93</td>
<td>16</td>
<td>45</td>
</tr>
<tr>
<td>Noophun et al., 2005</td>
<td>Retro</td>
<td>24</td>
<td>8.3</td>
<td>71</td>
<td>10</td>
<td>6</td>
</tr>
<tr>
<td>Mahadeva et al., 2003</td>
<td>Retro</td>
<td>23</td>
<td>6</td>
<td>96</td>
<td>35</td>
<td>24</td>
</tr>
<tr>
<td>Greenwald et al., 2003</td>
<td>Pilot</td>
<td>44</td>
<td>12</td>
<td>95</td>
<td>20</td>
<td>23</td>
</tr>
<tr>
<td>Sarin et al., 2002</td>
<td>RCT</td>
<td>9</td>
<td>15.4</td>
<td>89</td>
<td>22</td>
<td>11</td>
</tr>
<tr>
<td>Dhiman et al., 2002</td>
<td>Retro</td>
<td>18</td>
<td>31.6</td>
<td>100</td>
<td>10.3</td>
<td>NA</td>
</tr>
<tr>
<td>Lo et al., 2001</td>
<td>RCT</td>
<td>31</td>
<td>14</td>
<td>87</td>
<td>31</td>
<td>9</td>
</tr>
<tr>
<td>Huang et al., 2000</td>
<td>Retro</td>
<td>90</td>
<td>13.2</td>
<td>100</td>
<td>23</td>
<td>39</td>
</tr>
</tbody>
</table>

RCT = Randomized controlled trial; Retro = Retrospective; Pros = Prospective; NA = Not available; GVO = Gastric variceal obluration

### Table 3: GVL in the management of gastric variceal bleeding: comparison from different studies

<table>
<thead>
<tr>
<th>Author</th>
<th>Therapy</th>
<th>n</th>
<th>Active bleed (%)</th>
<th>Success (%)</th>
<th>Rebleeding (%)</th>
<th>Obliteration (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yoshida et al., 1994</td>
<td>GVL-S</td>
<td>10</td>
<td>10</td>
<td>100</td>
<td>10</td>
<td>100</td>
</tr>
<tr>
<td>Cipolletta et al., 1998</td>
<td>GVL-S</td>
<td>7</td>
<td>100</td>
<td>100</td>
<td>0</td>
<td>–</td>
</tr>
<tr>
<td>Shiha and El-Sayed, 1999</td>
<td>GVL</td>
<td>27</td>
<td>67</td>
<td>89</td>
<td>18</td>
<td>100</td>
</tr>
<tr>
<td>Lee and Shih, 2008</td>
<td>GVL</td>
<td>22</td>
<td>100</td>
<td>100</td>
<td>18</td>
<td>–</td>
</tr>
</tbody>
</table>

GVL = Gastric variceal ligation
bleeding GV, without any rebleeding in a follow-up period of almost 2 years.

Thrombin seems to be a promising therapy, and has the benefits of achieving excellent initial hemostasis and being easy to use with a good safety profile. Thrombin appears to be successful even in patients with GOV2, and may have a role in this difficult group.

**Combined endoscopic therapy (endoscopic variceal ligation-injection sclerotherapy)**

Endoscopic variceal ligation-injection sclerotherapy (EVLIS) is safe and effective in achieving hemostasis and obliteration in all patients as shown by Chun and Hyun.[40] The combination of variceal ligation and cyanoacrylate injection in GV effectively controlled acute bleed in 89% of patients; however 33% rebleed on follow-up.[41] Combination with sclerotherapy is unlikely to be accepted for the management of acute bleeding in view of the increased risk of iatrogenic complications, and the need for greater technical skill and procedure time. For secondary prophylaxis, combined therapy should be compared with standard established treatments.

**Endoscopic ultrasound guided treatment**

This is a new modality for treatment of GV and this has emerged as a valuable tool for diagnosis, treatment planning, evaluation of treatment efficacy, estimation of recurrent bleeding potential and also helps visualize varices, perforating veins, collateral veins and allows predict varices at high risk.

Romero-Castro et al., in their small case series injected cyanoacrylate-lipiodol into GV at the level of perforating veins, under EUS guidance.[29,42] All the procedures were successful, without recurrent bleeding or other complications during followup. They postulated that targeting perforating veins would produce the maximal blood-flow blockage, with the lower amounts of cyanoacrylate needed, therefore reducing the rate of potential local and systemic complications.

Romero-Castro et al. reported in other small case series of patients with GV treated with EUS guided coil embolization. They inserted coils into the perforating veins in order to block the blood flow. The varices were eradicated in three out of four patients, and no complications occurred in the successfully treated patients during five months of follow-up.[34,45]

Deployment of coil (ECA) as shown in Romero et al., use of 19 gauge needle and standard steel synthetic fibered 0.035 inches in diameter with straight length of 50-150 mm, coiled diameter of 8-20 mm, 3.2-5.6 configurator loops. The size of the coil was calculated according to the diameter of the vessel measured on EUS and was chosen to be approximately 20% larger, the needle was forwarded into target vessel under EUS guidance and the stylet withdrawn. The larger coil (20 mm) where deployed by using, the stiffer part of a 0.035-inch guide wire and a pusher. Smaller coils <15 mm were deployed by using floppy part of respective guide wire. Placing the needle tip opposite wall of vessel was avoided to allow enough space for coil to curl. If EUS vision becomes blurred contrast injection under fluoroscopy was used. Thrombosis of vessel was confirmed by injection of contrast.

Binmoeller et al. in their study on thirty patients combined EUS-guided coiling and cyanoacrylate glue injection by transesophageal approach to the gastric fundal varices. They hypothesized that coils with attached synthetic fibers (“wool coils”) inserted in to the varices prior to cyanoacrylate injection would function as a “barrier” for cyanoacrylate to outflow into the larger veins causing embolic events. In their procedure, after positioning and endosonographic visualization of GV from the esophagus (through the diaphragmatic crus muscle), a 19-gauge fine needle aspiration (FNA) needle was inserted through the esophageal wall and diahragm muscle directly to the gastric varix, following coil delivery and 1 mL of cyanoacrylate with immediate repetition of the procedure as needed until varix obliteration.[44]

**Steps of endoscopic ultrasound guided treatment**

Active flow within gastroesophageal varices was confirmed by color Doppler before therapy.

1. Intraluminal water filling of the gastric fundus to improve acoustic coupling and visualization of GV.
2. Echoendoscope positioned in the distal esophagus to sonographically visualize the gastric fundus in an anterograde fashion (the diaphragmatic crus muscle was identified between the esophageal wall and the GFV).
3. Endoscopic ultrasound-FNA needle (19 g) puncture into the GFV by using a transesophageal-transcrural approach.
4. Embolization coil (12-20-mm diameter) delivered into varix through the FNA needle by using the stylet as a pusher.
5. Immediate injection of 1 mL of 2-octyl-CYA after coil deployment through the same needle over 30 s by using normal saline solution to flush the glue through the catheter.

Color Doppler confirmed the absence of flow in varix after treatment. If the persistent flow was identified, an additional 1.0 mL of CYA glue was delivered. If varix had persistent flow and appeared large enough to accommodate another coil, the varix was repunctured with a new FNA needle, and the technique described was repeated.[44]
Other methods used for gastric varices treatment

Radiologic intervention

Transjugular intrahepatic portosystemic shunt

When patients with GV bleeding are unresponsive to initial endoscopic treatment, a second endoscopic therapy should be attempted if possible. If a second attempt fails or the severity of bleeding precludes further endoscopic therapy, salvage therapy using surgical shunts or TIPSs should be considered for refractory GV bleeding. Most current studies of TIPS focus on treatment for refractory GV bleeding and prevention of GV rebleeding. A recent study showed that the primary hemostasis rate of TIPS for acute GV bleeding is 92.3%. Other studies showed initial hemostasis of TIPS for acute refractory GV bleeding is between 87% and 100%, 58-62 with an approximate rebleeding rate of 10-30%.

Frequent complications of TIPS are encephalopathy and shunt stenosis/occlusion, with post-TIPS encephalopathy occurring in 4-16% of patients. The shunt dysfunction could be reduced by using polytetrafluoroethylene (PTFE)-covered stent. Considering the currently available evidence, TIPS with PTFE-covered stent is the treatment of choice for patients who failed first-line medical and endoscopic therapy.

Balloon-occluded retrograde transvenous obliteration

Kanagawa et al. first introduced the BRTO procedure in 1996. GVs are associated with a gastro-renal shunt (GRS) in 60-85%. The GRS drains blood flow into systemic circulation and provides a pathway for radiologists to treat GV. The outflow of GRS was blocked by inflating the balloon, and 5% ethanolamine oleateiopamidol was injected in a retrograde manner [Figure 5].

The BRTO was used as either primary or secondary prophylaxis of GV in most series. Initial hemostasis of BRTO for acute GV bleeding ranges between 76.9% and 100%. It was noted that the re-bleeding rate was from 0% to 15.4%.BRTO had a similar initial hemostasis and lower rebleeding rate, compared with GVO or band ligation for acute GV bleeding. The BRTO had been shown to be as effective as TIPS for acute GV bleeding, without increasing hepatic encephalopathy. Portal pressure also increased significantly after BRTO, which caused worsening of esophageal variceal pressure. Other common complications of BRTO are hemoglobinuria, abdominal pain, pyrexia, and pleural effusion. Major complications are shock and atrial fibrillation.

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