Analysis of Outcomes of Endovascular Embolisation: A Cross-Sectional Multicenter Study on 46 Visceral Artery Pseudoaneurysms

Mohammad Koriem Mahmoud Omar
Assiut University Hospital

Moustafa H. M. Othman
Assiut University Hospital

Robert A. Morgan
St George’s University of London Division of Cardiac and Vascular Sciences: St George’s University of London Molecular and Clinical Sciences Research Institute

Abdelkarem Hasan Abdallah
Assiut University Hospital

Hany M. A. Seif
Assiut University Hospital

Mohamed Zidan
Assiut University Hospital

Mahmoud K. A. Khairallah (m.khairallah@aun.edu.eg)
Assiut University Hospital

Reham Abd El-Aleem
Assiut University Hospital

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Abstract

Purpose
Visceral artery aneurysms are subdivided into true aneurysms and pseudoaneurysms. Visceral artery pseudoaneurysms (VAPAs) are uncommon in clinical practice but may have serious clinical outcomes up to death. Endovascular management is a safe effective alternative option to traditional surgical procedures. This study assesses the outcome of different embolic materials and techniques used in the endovascular management of visceral artery pseudoaneurysms.

Materials and methods
This is a multicentric prospective analysis of endovascular embolisation of 46 VAPAs with a mean pseudoaneurysm size of 13 ± 11.35 mm. Management using coils only was done in 28/46 patients (60.87%), NBCA glue only in 16/46 patients (34.78%), combined coils and NBCA glue in 1/46 patient (2.17%), and Amplatzer plugs only in 1 patient (2.17%). The management techniques were sac packing in 9/46 patients (19.57%), inow occlusion in 28/46 patients (60.87%) and trapping in 9/46 patients (19.57%).

Results
The overall clinical success rate was 93.48%, the overall perioperative complication rate was 15.22% and 30-day mortality was zero. For the coil subgroup (n = 28), the clinical success was 92.86%, while the subgroup of NBCA glue (n = 16) showed clinical success of 93.75%. There was no significant statistical difference between clinical success among coil, and NBCA glue subgroups (P > 0.05). The technical success rate was 100%. Effectiveness of the procedures during the follow-up was 97.83%. Target lesion re-intervention rate was 2.17%.

Conclusion
Transarterial embolisation can provide high technical and clinical success rates with low perioperative complication and re-intervention rates, as well as satisfactory procedure effectiveness in the management of VAPAs.

Background:
Visceral artery aneurysms (VAAs) typically occur within celiac trunk and its branches, superior or inferior mesenteric arteries and renal arteries. Aneurysms are subdivided into true aneurysms and pseudoaneurysms. Generally, true aneurysms are asymptomatic and occur secondary to underlying arterial diseases while pseudoaneurysms are a sequence of direct trauma, or inflammation of the vessel (1). Despite the rarity of visceral artery aneurysms, pseudoaneurysms are more frequently encountered in specialized centers dealing with acute trauma patients or high volumes of abdominal interventions than true aneurysms that are often incidentally discovered (2).

The imaging appearance of visceral artery pseudoaneurysms (VAPAs) is similar to that of true aneurysms, but typically exhibit more irregular margins, and the pseudoaneurysm is typically surrounded by a hematoma (2). Up to 70% of pseudoaneurysms and 20% of true aneurysms are liable to rupture and mortality occurs in 25–100% (3). Hyperdynamic circulation e.g. pregnancy, portal hypertension and infections are risk factors for rupture. Eighty per cent of the aneurysms of the hepatic artery are liable to rupture, followed by aneurysms of SMA and pancreaticoduodenal arcades (4–6).

In general, asymptomatic true visceral artery aneurysms that are less than 2 cm can be followed up without further management (2, 7). On the other hand, pseudoaneurysms must be managed regardless their presentation, size and location owing to their high possibility of rupture (7).

Endovascular management of VAPAs has been widely used as a safe and effective alternative treatment to the more invasive surgical procedures with higher mortality rate reaching 5% and mortality increases substantially if emergency surgery is required for aneurysm rupture repair (8–11). The current study was designed to assess the outcome of different embolic materials and techniques used in the endovascular management of visceral artery pseudoaneurysms. True visceral artery aneurysms are uncovered in this article.

Methods:
This is a multicentric prospective analysis of 46 patients with 46 visceral artery pseudoaneurysms of any size who were admitted to our hospitals and had endovascular management between July 2018 to March 2020. Informed consent was obtained from the patients. Those patients were presenting with either abdominal pain, or intrabdominal hemorrhage, or gastrointestinal (GIT) bleeding and/ or hemobilia or hematuria. A full medical history of co-morbidities and risk factors was taken for each patient. Clinical assessment and abdominal ultrasonography were done to all patients. Hemodynamically unstable patients received urgent medical support before further assessment.

Computed tomography angiography (CTA) was done to diagnose and confirm VAPAs in all patients prior to catheter angiography. CTA was performed either with a 64-slice multidetector helical CT, the Siemens SOMATOM Sensation 64 or 128-slice multidetector helical CT, the Siemens SOMATOMS Definition 128 (Siemens, Erlangen, Germany).
The following data were recorded: age, sex, associated co-morbidities along with risk factors, presentation, size as well as shape of pseudoaneurysm, affected artery, and location of the lesion within the artery (proximal, middle, or distal).

**Endovascular embolisation technique:**

Under local anesthesia, the procedures were performed by experienced (>10 years) interventional radiologists in dedicated interventional radiology suites on Artis Zee flat-type monoplane or Artis Q biplane digital subtraction angiography machines (Axiom-Artis; Siemens, Erlangen, Germany). Right transfemoral artery approach was performed in all cases.

Arterial access to the lesions was achieved by using 4 or 5 Fr standard angiographic catheters (Cobra, C1 angiographic catheter; Cook; Bloomington, IN), or (Sidewinder Simmons, Sim 1 Cordis; Johnson & Johnsons, Miami, FL) and 2.4 or 2.7 Fr coaxial microcatheter (Progreat Terumo Corporation, Tokyo, Japan) with different guide wires. The decision to use different types of embolic materials or even a combination was based on the arterial anatomy and on the decision of the interventional radiologist. Embolisation using coils only was done in 28/46 patients, while N-butylcyanoacrylate (NBCA) glue only was used in 16/46 patients. Combined coils and NBCA glue were used in 1/46 patient, and Amplatz vascular plugs were used in 1 patient.

When embolisation was performed using metallic detachable or pushable coils (MReye (Cook) or Interlock (Boston Scientific]) of variable diameters and lengths; the coils were oversized by ~20% compared with the target artery diameter.

When NBCA glue (Histoacryl Blue®; B. Braun, Melgungen, Germany) was used, the tip of the microcatheter was placed inside the aneurysm sac or as close as possible to the neck of the aneurysm. However, if the catheter tip could not be properly placed at the neck of the aneurysm because of the small caliber or tortuosity of the artery, it was wedged into the inlet of the arteries to be embolised to limit retrograde pericatheter reflux of the glue.

According to the desired rate of polymerization, NBCA was diluted manually with ethiodized oil (Lipiodol Ultra-Fluid®; Guerbet, Roissy-Charles-de-Gaulle, France, Switzerland), a polymerization-retardant. Specifically, when embolising a vessel of high-rate blood flow, or when the catheter was intralesional, we required quick in vivo polymerization and a ratio of 1:1 oil to NBCA was used. To delay glue polymerization, in situations where the microcatheter tip was positioned distant from the desired site of polymerization, a greater volume of ethiodized oil (ie, 2:1, 3:1 dilutions) was added.

The lumen of the microcatheter was flushed with 5% dextrose before injection of the NBCA mixture, thus preventing polymerization before reaching the arterial segments. Using a 1-mL syringe and under careful fluoroscopic monitoring, NBCA mixture was injected. In order to prevent adherence of the catheter tip to the vessel wall, the microcatheter was removed immediately after injection. Then, the guiding catheter was aspirated to clear its inner lumen, and post-embolic angiography was performed.

Amplatzer Vascular Plugs (St Jude Medical, St Paul, MN, USA) were used in a selected case (figure 1) where there was a pseudoaneurysm in a high-flow gastroduodenal artery (GDA) in order to reduce the risk of migration and systemic embolisation of traditional occlusion devices.

The embolisation techniques used in our study are illustrated in table 1. Figures 2 & 3 show the use of different embolic materials and techniques in the management of different visceral artery pseudoaneurysms.

**Table 1: Endovascular embolisation techniques used in our study:**

| Parent vessel flow preservation | Sac packing | Only the aneurysmal sac is filled with the embolic material |
|---------------------------------|-------------|------------------------------------------------------------|
| No parent vessel flow preservation | Trapping (sandwich, isolation, and front-to-back-door techniques): with or without sac packing | Embolic materials (coils or plugs) are deployed distally and proximally to the aneurysmal neck done to isolate the lesion and to prevent retrograde filling from the collaterals. The outflow artery ‘the back door’ is closed first, followed by inflow artery ‘the front door’.
| Inflow occlusion | Occlusion proximal to the aneurysmal neck |

**Follow up:**

All patients were followed up after discharge for 12 months on an outpatient basis. The follow up protocol of VAPA patients after endovascular treatment consisted of clinical assessment and duplex ultrasound examination at 1, 3, 6, and 12 months. CT was the basic tool of assessment in case of clinical suspicion of complications or symptoms recurrence.

**Study outcomes and definitions:**

- **Clinical success according to SIR guidelines** (12): is referred to as the 30-day clinical outcome based on clinical or imaging data or both per established guidelines. Resolution of signs and symptoms that prompted the endovascular procedure along with the absence of unexpected procedure-related complications within 30 days of the endovascular management is considered clinical success.
- **Perioperative complications were classified according to CIRSE classification system** (13).
- **Technical success according to SIR guidelines** (12): is defined as successful deployment of the embolic material within the intended artery with immediate complete aneurysm exclusion in the final angiographic control without evidence of contrast media extravasation.
- Perioperative procedure-related 30-day mortality rate.
- Effectiveness of the procedure: depends on complete exclusion of the aneurysm from the circulation without emergence of new symptoms and signs requiring aneurysmal re-intervention during the follow up (9).
- Target lesion re-intervention rate: is defined as requiring an additional procedure (open surgical or percutaneous or endovascular) due to target lesion recurrence or re-bleeding (14).

Statistical analysis:

Data was collected and analyzed using SPSS (Statistical Package for the Social Science, version 20, IBM and Armonk, New York). Continuous data were expressed in the form of mean ± SD and range while nominal data were expressed in the form of frequency (percentage). Chi square test was used to compare the clinical success between coil, and NBCA glue subgroups. P < 0.05 was considered the threshold of statistical significance.

Results:

Demographics and characteristics of aneurysms among enrolled patients are described in Table 2.
Table 2
Patients’ demographics and characteristics of the pseudoaneurysms.

| Demographics n = 46 |  |
|--------------------|---|
| Age (Years)        | 58.09 ± 22.66 |
| Sex                |  |
| - Male             | 34 (73.9%) |
| - Female           | 12 (26.1%) |
| Risk factors of the vascular lesions: | |
| - History of previous intervention (either endoscopy, percutaneous needle biopsy or surgery) | 26 (56.52%) |
| - Penetrating duodenal ulcers | 7 (15.21%) |
| - Intrabdominal infection and/ or inflammatory process | 4 (8.7%) |
| - Underlying vascular disease: (Vasculitis) | 1 (2.17%) |
| - Major trauma     | 1 (2.17%) |
| - Bleeding colonic diverticula | |
| Presentations      |  |
| - GIT hemorrhage and/or haemobilia | 16 (34.78%) |
| - Intra-abdominal hemorrhage | 14 (30.43%) |
| - Hematuria        | 11 (23.91%) |
| - Abdominal pain   | 5 (10.87%) |
| Characteristics of aneurysms among enrolled patients | |
| Shape of the aneurysm: Saccular | 46 (100%) |
| Mean size of the aneurysm (mm) | 13 ± 11.35 |
| Artery affected:   |  |
| - Renal artery     | 16 (34.78%) |
| - Gastroduodenal artery | 10 (21.74%) |
| - Superior mesenteric artery | 7 (15.22%) |
| - Hepatic artery   | 3 (6.52%) |
| - Pancreaticoduodenal arcades | 3 (6.52%) |
| - Inferior mesenteric artery | 2 (4.35%) |
| - Splenic artery   | 2 (4.35%) |
| - Cystic artery    | |
| Location of the aneurysm in relation to the segment of the affected artery: | |
| - Proximal segment | 4 (8.7%) |
| - Middle segment   | 9 (19.57%) |
| - Distal segment   | 33 (71.74%) |

Endovascular management among enrolled patients:

Tables 3, 4 & 5 show detailed endovascular management of VAPAs among enrolled patients. Overall clinical success was achieved in 43/46 patients (93.48%). For the subgroup of coils (n = 28), clinical success was achieved in 26/28 patients (92.86%). On the other hand, the subgroup of NBCA glue (n = 16) showed 93.75% (15/16) clinical success. We reported no significant statistical difference regarding clinical success among coil, and NBCA glue subgroups (P > 0.05). In lesions managed through sac packing technique (n = 9), clinical success was achieved in 7/9 patients (77.78%), while in lesions managed through inflow occlusion (n = 28) and trapping techniques (n = 9), clinical success was achieved in 27/28 (96.43%), and 9/9 (100%) of the patients, respectively.

Perioperative complications were reported in 7/46 patients (15.22%). Grade-2 complication was reported in 4 patients (8.7%) representing mild post embolisation syndrome (transient pain requiring only oral analgesia with no prolongation of hospital stay). Grade-3 complication was reported in 1 patient (2.17%) that had cystic artery pseudoaneurysm and was complicated by aneurysmal sac rupture and re-bleeding after being managed by coils through sac packing technique. That was successfully managed by inflow occlusion of the parent artery using NBCA glue. Grade-4 complication (permanent mild sequelae) was reported in two patients (4.35%). One patient with cystic artery pseudoaneurysm after being embolised by NBCA glue through sac packing technique developed ischemia of the gall bladder with subsequent necrosis and abscess formation that required further percutaneous tubal drainage and
cholecystectomy later-on. The other patient had pseudoaneurysm in the jejunal branch of SMA and was complicated by focal jejunal loop ischemia after being managed by coils through inflow occlusion technique. That was successfully managed by laparotomy and resection anastomosis surgery of the ischemic jejunal loop. The 3 patients who had grade 3 & 4 complications were responsible for the small percentage of the overall clinical failure in the study.

Technical success was achieved in 100% of the patients with no reported 30-day mortality in our study. Procedure effectiveness was achieved in 45/46 patients (97.83%). Only one patient required re-intervention that had cystic artery pseudoaneurysm with successful clinical outcome later-on.
| N | Anatomy | Morphology | Co-Morbidities & risk factors/ Presentation | Urgent or elective management | Embolisation technique | Embolic material | Technical success | Complications | Clinical success | Effectiveness of the procedure |
|---|---------|------------|---------------------------------------------|-------------------------------|------------------------|-----------------|-----------------|---------------|----------------|--------------------------------|
| 1 | SA      | 23 mm saccular aneurysm | Pancreatitis/ Abdominal pain | Urgent | Sac packing | NBCA Glue | Yes | Grade 2 (Mild post embolisation syndrome) | Yes | Yes | ![](https://via.placeholder.com/15) |
| 2 | SMA     | 45 mm saccular aneurysm | Vasculitis/ Intrabdominal hemorrhage | Urgent | Sac packing | NBCA Glue | Yes | No | Yes | YES | ![](https://via.placeholder.com/15) |
| 3 | RT RA   | 20 mm saccular aneurysm | Iatrogenic/ Hematuria | Urgent | Sac packing | NBCA Glue | Yes | No | Yes | YES | ![](https://via.placeholder.com/15) |
| 4 | RT HA   | 13 mm saccular aneurysm | Iatrogenic/ Intrabdominal hemorrhage | Urgent | Sac packing | NBCA Glue | Yes | No | Yes | YES | ![](https://via.placeholder.com/15) |
| 5 | LT RA   | 5.5 mm saccular aneurysm | Iatrogenic/ Hematuria | Urgent | Sac packing | NBCA Glue | Yes | No | Yes | YES | ![](https://via.placeholder.com/15) |
| 6 | Cystic a | 20 mm saccular aneurysm | Acute cholecystitis/ GIT bleeding and haemobilia | Urgent | Sac packing | NBCA Glue | Yes | Grade 3 (Ischemia of the GB with subsequent necrosis & abscess formation) | No | Yes | ![](https://via.placeholder.com/15) |
| 7 | GDA     | 8 mm saccular aneurysm | Penetrating duodenal ulcer/ GIT bleeding | Urgent | Inflow occlusion | NBCA Glue | Yes | No | Yes | YES | ![](https://via.placeholder.com/15) |
| 8 | RT RA   | 11 mm saccular aneurysm | Iatrogenic/ Hematuria | Urgent | Inflow occlusion | NBCA Glue | Yes | Grade 2 (Mild post embolisation syndrome) | Yes | Yes | ![](https://via.placeholder.com/15) |
| 9 | LT RA   | 2 mm saccular aneurysm | Vasculitis/ Intrabdominal hemorrhage | Urgent | Inflow occlusion | NBCA Glue | Yes | No | Yes | YES | ![](https://via.placeholder.com/15) |
| 10| GDA     | 2.5 mm | Vasculitis/ GIT bleeding | Urgent | Inflow occlusion | NBCA Glue | Yes | No | Yes | YES | ![](https://via.placeholder.com/15) |
| 11| RT RA   | 4 mm saccular aneurysm | Iatrogenic/ Hematuria | Urgent | Inflow occlusion | NBCA Glue | Yes | No | Yes | YES | ![](https://via.placeholder.com/15) |
| 12| LT RA   | 9 mm saccular aneurysm | Iatrogenic/ Hematuria | Urgent | Inflow occlusion | NBCA Glue | Yes | No | Yes | YES | ![](https://via.placeholder.com/15) |
| 13| LT RA   | 5.3 mm saccular aneurysm | Iatrogenic/ Intrabdominal hemorrhage | Urgent | Inflow occlusion | NBCA Glue | Yes | No | Yes | YES | ![](https://via.placeholder.com/15) |
| 14| RT HA   | 13 mm saccular aneurysm | Iatrogenic/ Intrabdominal hemorrhage | Urgent | Inflow occlusion | NBCA Glue | Yes | No | Yes | YES | ![](https://via.placeholder.com/15) |
| 15| RT RA   | 24 mm saccular aneurysm | Iatrogenic/ Hematuria | Urgent | Inflow occlusion | NBCA Glue | Yes | No | Yes | YES | ![](https://via.placeholder.com/15) |
| 16| GDA     | 13 mm saccular aneurysm | Penetrating duodenal ulcer/ GIT bleeding | Urgent | Inflow occlusion | NBCA Glue | Yes | No | Yes | YES | ![](https://via.placeholder.com/15) |

SA: splenic artery, SMA: superior mesenteric artery, RA: renal artery, HA: hepatic artery, GDA: gastroduodenal artery, NBCA: N-butylcyanoacrylate
Table 4

Pathophysiological criteria of the pseudoaneurysms treated with coils and their management techniques and outcomes.

| No | Anatomy      | Morphology            | Co-Morbidities & risk factors/ Presentation | Urgent or elective management | Embolisation technique | Embolic material                  | Technical success | Complications | Clinical success | CoM | E | C |
|----|--------------|-----------------------|--------------------------------------------|-------------------------------|--------------------------|------------------------|----------------------------------|------------------|---------------|------------------|-----|---|---|
| 1  | GDA          | 44.5 mm Saccular aneurysm | iatrogenic/ Abdominal pain                 | Urgent                        | Trapping                 | 3 detachable micro coils  | Yes                | No             | Yes              |     | Y |   |
| 2  | GDA          | 2.5 mm saccular aneurysm  | Penetrating duodenal ulcer/ GIT bleeding   | Urgent                        | Sac packing              | 2 pushable coils         | Yes                | No             | Yes              |     | Y |   |
| 3  | GDA          | 10.5 mm saccular aneurysm | Pancreatitis/ Intrabdominal hemorrhage     | Urgent                        | Trapping with sac packing | 3 detachable micro coils  | Yes                | No             | Yes              |     | Y |   |
| 4  | GDA          | 26.5 mm saccular aneurysm | Penetrating duodenal ulcer/ GIT bleeding   | Urgent                        | Trapping with occlusion of the collaterals | 3 detachable micro coils  | Yes                | No             | Yes              |     | Y |   |
| 5  | IMA          | 14.5 mm saccular aneurysm | iatrogenic/ GIT bleeding                  | Urgent                        | Trapping                 | 2 detachable micro coils  | Yes                | No             | Yes              |     | Y |   |
| 6  | GDA          | 15 mm saccular aneurysm  | Penetrating duodenal ulcer/ GIT bleeding   | Urgent                        | Trapping                 | 2 detachable micro coils  | Yes                | No             | Yes              |     | Y |   |
| 7  | GDA          | 15 mm saccular aneurysm  | Pancreatitis/ Abdominal pain              | Urgent                        | Trapping                 | 3 detachable micro coils  | Yes                | No             | Yes              |     | Y |   |
| 8  | Pancreaticoduodenal a | 3 mm saccular aneurysm  | iatrogenic/ Intrabdominal hemorrhage      | Urgent                        | Trapping                 | 3 detachable micro coils  | Yes                | No             | Yes              |     | Y |   |
| 9  | IMA          | 4 mm saccular aneurysm   | iatrogenic/ GIT bleeding                  | Urgent                        | Trapping with sac packing | 3 detachable micro coils  | Yes                | No             | Yes              |     | Y |   |
| 10 | Cystic a     | 7 mm saccular aneurysm   | iatrogenic/ GIT bleeding and haemobilia   | Urgent                        | Sac packing              | Single detachable micro coil | Yes                | Grade 3 (Re-bleeding required re-intervention) | No |    |    |
| 11 | RT RA        | 3 mm saccular aneurysm   | iatrogenic/ Hematuria                     | Urgent                        | Inflow occlusion         | Single pushable coil     | Yes                | No             | Yes              |     | Y |   |
| 12 | LT RA        | 20 mm saccular aneurysm  | iatrogenic/ Hematuria                     | Urgent                        | Inflow occlusion         | 2 pushable coils         | Yes                | Grade 2 (Mild post embolisation syndrome) | Yes |     |   |
| 13 | Pancreaticoduodenal a | 27 mm saccular aneurysm  | Penetrating duodenal ulcer/ Intrabdominal hemorrhage | Urgent                        | Inflow occlusion         | 2 detachable micro coils  | Yes                | No             | Yes              |     | Y |   |
| 14 | SMA          | 34 mm saccular aneurysm  | Vasculitis/ Intrabdominal hemorrhage      | Urgent                        | Inflow occlusion         | 3 detachable micro coils  | Yes                | No             | Yes              |     | Y |   |
| 15 | Pancreaticoduodenal a | 19.75 mm saccular aneurysm | Penetrating duodenal ulcer/ Intrabdominal hemorrhage | Urgent                        | Inflow occlusion         | 2 detachable micro coils  | Yes                | No             | Yes              |     | Y |   |
| 16 | RT RA        | 5.5 mm saccular aneurysm  | iatrogenic/ Hematuria                     | Urgent                        | Inflow occlusion         | 2 pushable coils         | Yes                | No             | Yes              |     | Y |   |
| 17 | RT RA        | 5 mm saccular aneurysm    | iatrogenic/ Hematuria                     | Urgent                        | Inflow occlusion         | Single pushable coil     | Yes                | No             | Yes              |     | Y |   |

GDA: gastroduodenal artery, IMA: inferior mesenteric artery, RA: renal artery, SMA: superior mesenteric artery, SA: splenic artery
| N | Anatomy | Morphology | Co-Morbidities & Risk factors/ Presentation | Urgent or elective management | Embolisation technique | Embolic material | Technical success | Complications | Clinical success | Effectiveness of the procedure |
|---|---|---|---|---|---|---|---|---|---|---|
| 18 | LT RA | 14.1 mm saccular aneurysm | Iatrogenic/ Intrabdominal hemorrhage | Urgent | Inflow occlusion | 2 pushable coils | Yes | No | Yes | Y |
| 19 | RT RA | 3 mm saccular aneurysm | Iatrogenic/ Hematuria | Urgent | Inflow occlusion | 2 pushable coils | Yes | No | Yes | Y |
| 20 | RT RA | 3.5 mm saccular aneurysm | Iatrogenic/ Intrabdominal hemorrhage | Urgent | Inflow occlusion | 2 detachable micro coils | Yes | No | Yes | Y |
| 21 | LT RA | 33.5 mm saccular aneurysm | Septic emboli/ Abdominal pain | Urgent | Inflow occlusion | 3 detachable micro coils | Yes | No | Yes | Y |
| 22 | SA | 3.5 mm saccular aneurysm | Trauma/ Intrabdominal hemorrhage | Urgent | Inflow occlusion | 2 pushable micro coils | Yes | No | Yes | Y |
| 23 | SMA | 2 mm saccular aneurysm | Iatrogenic/ GIT bleeding | Urgent | Inflow occlusion | 2 detachable micro coils | Yes | Grade 4 (Bowel loop ischemia) | No | Y |
| 24 | SMA | 5 mm saccular aneurysm | Iatrogenic/ GIT bleeding | Urgent | Inflow occlusion | 3 detachable micro coils | Yes | Grade 2 (Mild post embolisation syndrome) | Yes | Y |
| 25 | SMA | 2.2 mm saccular aneurysm | Iatrogenic/ GIT bleeding | Urgent | Inflow occlusion | 2 detachable micro coils | Yes | No | Yes | Y |
| 26 | SMA | 3.2 mm saccular aneurysm | Iatrogenic/ GIT bleeding | Urgent | Inflow occlusion | 2 detachable micro coils | Yes | No | Yes | Y |
| 27 | SMA | 5.1 mm saccular aneurysm | Iatrogenic/ GIT bleeding | Urgent | Inflow occlusion | 2 detachable micro coils | Yes | No | Yes | Y |
| 28 | IMA | 5 mm saccular aneurysm | Diverticula / GIT bleeding | Urgent | Inflow occlusion | 2 detachable micro coils | Yes | No | Yes | Y |

GDA: gastroduodenal artery, IMA: inferior mesenteric artery, RA: renal artery, SMA: superior mesenteric artery, SA: splenic artery

Table 5
Pathophysiological criteria of the pseudoaneurysms treated with Amplatzer vascular plugs or mixed NBCA & coils and their management techniques and clinical success

Discussion:
It is essential to mention that the clinical response of endovascular embolisation of VAPA depends on the type of the embolic agent and adequacy of the embolisation process. When choosing an embolic agent, many factors should be taken into consideration. These factors include site, and size of the lesion, as
well as the flow pattern of vessels to be occluded, the availability of embolic agents, the experience and knowledge of the radiologist who will perform the procedure, the speed and reliability of delivery, the duration of the occlusive effect, and the avoidance of non-target embolisation (15). In our study, we used mainly permanent occlusive agents to avoid recanalization of the lesion and recurrence of presenting symptoms would be expected to be less. Coils were the most frequent materials used in the management either alone (60.87%) or with NBCA glue (2.17%). Techniques of embolisation used in the study were sac packing, inflow occlusion and trapping in 19.57%, 60.87% and 19.57% of the patients, respectively.

In our study, the overall clinical success rate was 93.48% with zero 30-day mortality rate. These results were comparable to those of Venturini et al who achieved 83% clinical success with a 7% 30-day mortality rate (10). For the subgroup of coils (n = 28), the clinical success was 92.86%, while the subgroup of NBCA glue (n = 16) showed clinical success of 93.75%. There was no significant statistical difference regarding clinical success between the coil and NBCA glue subgroups as the embolic material of management (P > 0.05). These results were similar to Alwarraky et al who reported a non-significant statistical difference between coils and NBCA glue as permanent embolic materials in the endovascular management of acute renal bleeding (16).

In the current study, 7/46 patients (15.22%) developed perioperative complications. This is comparable with the range of complication rates for endovascular techniques generally reported in other series (0–50%) (17–19).

Our technical success rate was 100% (46 out of 46 patients) and this was comparable to the most of other similar studies in the literature (20–23). Procedure effectiveness in the current study was 97.83% with complete aneurysmal sac exclusion without the emergence of new symptoms and signs requiring aneurysmal re-intervention. Only one patient with a cystic artery aneurysm showed revascularization of the aneurysmal sac on follow-up imaging. In line with this result, Spiliopoulos et al that showed a long-term efficacy of endovascular management with only 6.1% target lesion re-intervention rate during a mean period of follow-up of 19.1 ± 21.4 months (14). In this series, the target lesion re-intervention rate was 2.17% (the 1 patient who had a cystic artery pseudoaneurysm). The pseudoaneurysm was initially embolised by 30 cm x 6 mm detachable micro coil. However, it was complicated after 1 week of the procedure by rupture of the aneurysmal sac and migration of the coil into the CBD down the duodenum (Fig. 5); hence, re-embolisation was done using NBCA in two different sessions; in the first session, sac packing was done with complete aneurysm exclusion from the final angiographic image. Again, it was complicated by sac rupture 1 month later. In the second session, parent artery embolisation (inflow occlusion) was done successfully. Target lesion re-intervention rate in previous studies ranged between 6.7–15% (3, 10, 14).

The satisfactory results of endovascular embolisation could be due to the continuous advances in embolic materials and catheter designs used in interventional catheter-based techniques; the development of microcatheter technology has enabled selective catheterization of even small-caliber vessels and the use of micro coils and different polymerization rates of NBCA glue has allowed a more targeted embolisation (10).

The main limitations in our study were 1) the mid-term evaluation and so, knowledge of the durability of embolisation is limited to 1 year only, 2) the small number of patients in each subgroup for comparison, and 3) the non-randomization of the studied subgroups. In the future, a randomized study to compare efficacy of each embolic agent and each embolisation technique is desirable.

Conclusion

Transarterial embolisation of visceral artery pseudoaneurysms can provide high technical and clinical success rates with low perioperative complication and re-intervention rates, as well as satisfactory procedure effectiveness in the management of visceral artery pseudoaneurysms. There is no significant statistical difference between clinical success among coil, and NBCA glue in the embolisation of visceral artery pseudoaneurysms.

Abbreviations

VAA visceral artery aneurysm, VAPA: visceral artery pseudoaneurysm, GIT: gastrointestinal, CT: computed tomography, NBCA: N-butylcyanoacrylate, SIR: Society of Interventional Radiology, CIRSE: Cardiovascular and Interventional Radiological Society of Europe, SMA: superior mesenteric artery, IMA: inferior mesenteric artery, CHA: common hepatic artery, HA: hepatic artery, RA: renal artery, SA: splenic artery, GDA: gastroduodenal artery.

Declarations

Ethics approval: All procedures performed in the study were in accordance with the ethical standards of the institutional research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards. This study was approved by the Research Ethics Committee of faculty of medicine Assiut University in Egypt IRB number 17200220.

Consent to participate: Informed consent was obtained from all individual participants included in the study.

Consent for publication: Consent for publication was obtained for every individual person’s data included in the study.

Availability of data and materials: The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Conflicts of interests: All authors declare that they have no conflicts of interest which include financial or personal relationships that inappropriately influence their actions.

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Figure 1

Embolisation of gastroduodenal artery pseudoaneurysm. A) CT angiogram showing gastroduodenal artery pseudoaneurysm surrounded by hematoma. B) Selective angiogram of the gastroduodenal artery demonstrating the lesion. C) Embolisation of the gastroduodenal artery pseudoaneurysm by trapping technique with 6.5mm and 5mm diameter microvascular plugs distally & 7mm diameter Amplatzer IV plug proximally. D) Final angiogram showing complete exclusion of the pseudoaneurysm from the circulation.

Figure 3

Embolisation of gastroduodenal artery pseudoaneurysm by trapping technique with multiple micro coils. A) Selective angiogram of the celiac axis and gastroduodenal artery demonstrate pseudoaneurysm arising from the gastroduodenal artery with associated replaced right hepatic artery arising from the
gastroduodenal artery at the neck of the pseudoaneurysm. B) Embolisation of the front and back doors of the pseudoaneurysm as well as the replaced right hepatic artery using 4, 5 and 6 mm detachable 0.018 coils.

Figure 4
Emboliisation of CHA pseudoaneurysm by sac packing technique with multiple coils and NBCA/ Lipiodol mixture.

Figure 5
Emboliisation of cystic artery pseudoaneurysm. A) Coiling of cystic artery pseudoaneurysm using sac packing technique. B) Embolisation of the re-filled pseudoaneurysm using NBCA/ Lipiodol mixture ‘sac packing technique’. C) Embolisation of right hepatic artery proximal to the stump of cystic artery after 2nd
time re-filling of the pseudoaneurysm using NBCA/ Lipiodol mixture ‘inflow occlusion’.