Endovascular treatment for aortic arch pathologies: chimney, on-the-table fenestration, and in-situ fenestration techniques

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Background: Revascularization of the supra-aortic major branches in thoracic endovascular aortic repair (TEVAR) is challenging owing to the complex anatomic configuration of aortic arch pathologies. This study aims to evaluate the feasibility, effectiveness, and safety of three major techniques—chimney, fenestrated, and in-situ fenestration—for patients with aortic arch pathologies.

Methods: A retrospective analysis was performed involving 234 patients with aortic arch lesions, who underwent TEVAR with adaptations in technique (chimney, fenestrated, or in-situ fenestration) between January 2016 and December 2017.

Results: One hundred and twenty-six patients underwent the chimney technique (98 single chimneys, 24 double chimneys, and four triple chimneys); one hundred and two patients (102/234) were treated with on-the-table fenestration technique (92 single fenestrations, nine double fenestrations, and one double fenestration plus innominate artery chimney); and the remaining six patients underwent in-situ needle fenestration technique. Overall, indications included aortic dissections (99/234), aortic arch aneurysms (60/234), penetrating aortic ulcers (72/234), and re-interventions (3/234). The technical success rates were 99.6%. There were five cases of early all-cause mortality. The patency rates of overall branches were 99.6%. There were 15 cases with type Ia endoleak—14 in the chimney group (11.1%) and one in the on-the-table fenestration group (1%). Five patients underwent re-interventions. The median follow-up time for all patients was 28 (range, 16–41) months.

Conclusions: Our experience suggests that chimney, on-the-table fenestration, and in-situ needle fenestration techniques are feasible, effective, and safe treatment options for aortic arch pathologies with encouraging mid-term results. Long-term durability concerns require further evaluation.

Keywords: Endovascular procedure; aortic diseases; aneurysm; dissecting
Introduction

Treating aortic arch pathologies such as aortic dissections and aneurysms is challenging for vascular surgeons. Although conventional surgical repair remains the therapeutic gold standard, surgical repair with cardiopulmonary bypass and deep hypothermic circulatory arrest is associated with a high risk of mortality (1). The hybrid technique is the refined alternative to open surgery; however, it is still associated with high mortality (2). High-risk patients with comorbidities are deemed unsuitable to receive surgical repairs.

Thoracic endovascular aortic repair (TEVAR) has seen rapid development and application in the past two decades. Owing to the development of stent-grafts and endovascular techniques, the indications for TEVAR have expanded, and it is currently used to treat aortic arch pathologies. Despite favourable long-term results, the applicability of custom-made stent-grafts is limited because of technical difficulties and long manufacturing delays, especially in urgent cases with severe acute end-organ ischemia or impending rupture (3). Moreover, custom-made stent-grafts are not commercially available at all times in many institutes.

The chimney (C-TEVAR), on-the-table fenestration (F-TEVAR), and in-situ fenestration (I-TEVAR) are TEVAR-assistive techniques, that could be accomplished with readily available devices. C-TEVAR has earned worldwide popularity owing to the procedure being less complicated, feasible across various anatomic features, and being applicable in a bail-out situation (4). F-TEVAR, also known as physician-modified fenestrated graft technique, is a quick procedure for an off-the-shelf stent-graft by the surgeon in the operating room (5-7). I-TEVAR, including in-situ laser fenestration and in-situ needle fenestration, is also a potential technique for revascularization of supra-aortic branches (8,9). Because each of these techniques has strengths and limitations, endovascular treatment strategies for aortic arch diseases are still controversial. Herein, we provided the outcomes of 234 consecutive patients who received C-TEVAR, F-TEVAR, or I-TEVAR for aortic arch pathologies, aimed to assess the feasibility, effectiveness, and safety of these techniques and probably provide good alternatives for open repair in selected patients.

Methods

The patient cohort was studied retrospectively from January 1, 2016 to December 31, 2017. The institutional review boards of both Fuwai Hospital, National Centre for Cardiovascular diseases and the Second Xiangya Hospital approved the study, and the need for informed consent was waived (ID: D171100002917004, Dated: January 1, 2016).

Patient eligibility

An interdisciplinary board composed of endovascular and cardiovascular surgeons, radiologists, and anaesthesiologists was responsible for comprehensively evaluating each patient to decide treatment options. Patients who met the following criteria were considered suitable candidates for undergoing assistive techniques in TEVAR: those with (I) aortic arch pathologies involving single or multiple supra-aortic branches; (II) aortic arch pathologies including aortic dissection (AD), penetrating aortic ulcer (PAU), and pseudoaneurysm of aortic arch (PAA); and (III) patients who were aged with comorbidities and hence unsuitable for the risks of open or hybrid surgery.

The exclusion criteria for endovascular treatment were as follows: patients with (I) proximal intimal tear of the AD involving the ascending aorta; (II) concomitant pathologies such as cardiac tamponade and severe aortic regurgitation which necessitated surgical treatment; and (III) other anatomical formations not suitable for TEVAR, such as severe tortuosity in the access route arteries, angulation or compression, or very large diameter of the maximum trans-aortic diameter severely limiting device use.

The most suited TEVAR-assistive technique was selected based on the following principles: F-TEVAR was the first option when aortic arch pathologies were located merely at the inner curve of the aortic arch, because a large-sized fenestration could be made to efficiently align the fenestration with the vascular branches. When pathologies involved the outer curve of the aortic arch and the diameters of the branch artery were large, F-TEVAR with a small-sized fenestration was used. Varieties of fenestrations are shown in Figure 1. When the angulation between the aortic arch and the descending aorta was hostile for orientation of the fenestration (Figure 2), and when branch artery dissection was present, C-TEVAR was employed. When the proximal intimal tear in the AD was located at the outer curve and very close to the branch arteries, the risk of an endoleak was considerable with the previous two techniques. In such cases, I-TEVAR was used.
Pre-procedure preparation
All patients were administered general anaesthesia in our hybrid operating suite. The common femoral artery was exposed surgically or percutaneously using Perclose ProGlide suture device (Abbott Laboratories Co., Ltd, USA).

Selection for stent-graft
In the C-TEVAR group, Valiant (Medtronic, Inc, Minneapolis, MN, USA) and Hercules (MicroPort Medical Co., Ltd, Shanghai, China) were selected as the main thoracic aortic stent-grafts, because these grafts have better flexibility than Ankura (Lifetech, Scientific Co., Ltd, Shenzhen, China) for stent-graft apposition with branch endografts. In contrast, Ankura stent-graft was used in F-TEVAR, because the unique marker in an Ankura graft could facilitate stent-graft orientation. The proximal part of the Ankura stent-graft has two differently shaped radiopaque markers—an “8”-shaped marker on one side and an O-shaped marker on the opposite side. Moreover, Ankura also contains a spiral support strut, which is aligned with the “8”-shaped marker. The linear alignment of the “8”-shaped marker and the spiral strut of Ankura stent-graft indicated the exact position of the fenestration, which helped surgeons to accurately orient the fenestration to the orifice of the aortic branches. In contrast, Valiant stent-graft has four almost identical markers that need further modification to distinguish one marker for the orientation procedure.

Two types of covered endografts, Fluency (C.R. Bard, Inc, Murray Hill, NJ, USA) and Viabahn (Gore & Associates, Flagstaff, AZ, USA), were mostly used as grafts.
for branch arteries. A bare endograft (Biotronik Medical Co., Ltd, Berlin, Germany) was also used in the F-TEVAR group.

**Chimney technique procedure**
Our standardized C-TEVAR procedure has been previously described (10,11). Basically, the chimney graft for the supra-aortic branches was inserted via brachial/carotid access and the aortic stent-graft was introduced through the common femoral access. Then, the chimney grafts were released parallel to the main stent-graft with the proximal segment protruding at least 2.0 cm into the aortic lumen and the distal end remaining in the branch artery, while ensuring that at least 1.0 cm of overlapping segment remained between the two grafts. The aortic stent-grafts were selected with a 0–20% oversizing, and the chimney graft was selected with a 0–5% oversizing. Routine moulding of the chimney graft was performed via a balloon.

**On-the-table fenestration procedure**
Adequate understanding of the anatomical structure via the pre-operative CTA was essential for the success of orientation. Digital subtraction angiography (DSA) was performed and the diameters of vascular branches were measured to design a fenestration with comparable size to the target vessels. The proximal part of the aortic stent-graft

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**Figure 2** Morphology of the aortic arch. (A,B) The vertical and horizontal 3D fusion image of a patient with aortic arch aneurysm. Red lines showed an angle of 54 degrees between the aortic arch and the descending aorta. For this hostile aortic arch angulation, chimney technique would be selected. (C,D) The vertical and horizontal 3D fusion image of another patient with aortic arch aneurysm. Red lines showed an angle of 22 degrees. This less angulated aortic arch allows planning of on-the-table fenestration technique.
was unsheathed in vitro, measured, and marked by a sterile pen. Then, the fenestration was fashioned by a scalpel alongside the spiral support strut beside the eight-shaped marker (Figure 1A). The edges of the fenestration were kept neat, and all fragments were carefully removed (Figure 1B).

Then, the fenestrated stent-graft was carefully reassembled. Guided by an extra-stiff guide wire, the stent-graft was inserted into the descending aorta via femoral access. After confirming that the “8”-shaped marker and the spiral strut of the stent-graft were aligned with the target vessel, the stent-graft was carefully released. Then, angiography was performed to check the results (Figure 3A, B).

The indications for stenting of aortic branches, already preserved by F-TEVAR, were as follows: (I) the IA and LCCA, being crucial brain supplying arteries, should always be stented by endografts, and (II) the following evaluation results showed that the LSA was not well preserved and required stenting: (i) intraoperative surveillance of bilateral blood pressure showed a significant blood pressure difference between bilateral brachial arteries; (ii) completion DSA showed poor blood flow through the orifice of the LSA. Either antegrade femoral access (Figure 3C) or retrograde brachial/carotid access (Figure 3D) could be used for delivery of endografts into the aortic branches. Femoral
access required no extra brachial or carotid incisions, and was generally tried first.

**In-situ needle fenestration procedure**
First, the aortic stent-graft was introduced to the aortic arch and deployed at the planned landing zone. Then, a balloon expandable puncture needle (Lifetech, Scientific Co., Ltd, Shenzhen, China; Figure 4A,B) was introduced through the Fustar steerable vascular sheath (Lifetech) from the brachial access and advanced to the orifice of the LSA. The fenestration was created by inflating the balloon to activate the puncture needle (Figure 4C). Then, the guidewire was passed through the newly created fenestration to reach the aortic lumen. After balloon expansion of the fenestration site (Figure 4D), a covered stent-graft was deployed to ensure patency of the aortic branch. Finally, the stent-graft for the aortic branch was again dilated via a balloon, and completion DSA was performed to check the result.

**Post-operative management**
Each patient was managed postoperatively by oral medication including antiplatelet drugs, antihypertensive drugs, and beta-blockers. The follow-up for each patient was scheduled at 1, 3, 6, and 12 months, and annually thereafter.

**Statistical analysis**
Continuous data are expressed as the mean (± SD) or
median (range). Cumulative survival was analysed using Kaplan–Meier survival curves. Analyses were conducted using SPSS 23.0 software (IBM Corporation, Somers, NY, USA). GraphPad Prism 8 (GraphPad Software, San Diego, CA, USA) was used for graphical representation of data.

Results

From January 1, 2016 to December 31, 2017, a total of 859 patients underwent surgeries for aortic arch pathologies in both Fuwai Hospital and Second Xiangya Hospital. We excluded 424 patients undergoing open arch replacement and 201 patients receiving the hybrid technique. Finally, 234 patients who received TEVAR combined with assistive techniques were included. Of these, 126 (53.8%) in the C-TEVAR group underwent the chimney technique, 102 (43.6%) in the F-TEVAR group underwent on-the-table fenestration technique, and 6 (2.6%) in the I-TEVAR group underwent the in-situ needle fenestration technique. Baseline characteristics and patient demographics of each group are detailed in Table 1.

In the C-TEVAR group, 95, 28, and three cases of single-chimney, double-chimney, and triple-chimney techniques were performed. A total of 158 chimney grafts (CGs) were utilised, including 100 Fluency grafts (LSA, n=60; LCCA, n=31; IA, n=9) and 58 Viabahn grafts (LCCA, n=14; LSA, n=43; ARSA, n=1). In the F-TEVAR group, 91 single fenestrations (90 for the LSA, 1 for the ARSA), nine double fenestrations (7 for LSA and LCCA, 2 for LSA and aberrant left vertebral artery), and one double fenestration for LCCA and LSA combined with chimney for IA were performed. Thirteen branch arteries were stented (IA, n=1; LCCA, n=2, LSA, n=10) using 12 covered endografts (Fluency, n=1; Viabahn, n=11) and 1 bare endograft (Biotronik, n=1). Three (23.1%) LSA stent-grafts were deployed by femoral access, while the remaining 10 stent-grafts were deployed from brachial or carotid access. In the I-TEVAR group, all six patients underwent the in-situ fenestration technique for LSA. More details are listed in Table 2.

Periprocedural outcomes

Technical success was defined as isolation of pathologies while ensuring patency of the supra-aortic branches in the completion DSA. This was achieved in 233 of 234 patients (99.6%). Alignment of fenestration to the LSA failed in one patient in the F-TEVAR group (1/102, 1.0%) because of the lack of experience at the start of our practice. The occluded LSA in this case was revascularized by stenting the bail-out chimney graft and remained patent during the follow-up.

In all, 5 of 234 (2.1%) patients died within 30 days. One patient who received C-TEVAR for a huge aortic arch pseudoaneurysm died of sudden cardiac arrest 15 days postoperatively. Another patient with AD developed renal ischemic complications and died of renal failure. The third patient with AD died of bowel ischemia necrosis because of severely compressed distal aortic true lumen, despite emergent measures taken to expand the true lumen. In the F-TEVAR group, one patient died of ischemic stroke and the other died from bowel ischemia necrosis.

Early type Ia endoleak, detected exclusively by the post-operative CTA on discharge, was found in 15 patients (6.4%)—14 (11.1%) in the C-TEVAR group and one (1.0%) in the F-TEVAR group. The median ICU stay duration was 1 (range, 0–8) day in the C-TEVAR group, 1 (range, 0–7) day in the F-TEVAR group, and 1 (range, 1–2) day in the I-TEVAR group, and the respective median hospital stays were 6 (range, 1–22) days, 6 (range, 5–19) days, and 6 (range, 5–8) days.

Mid-term follow-up outcomes

The LSA of one patient in the F-TEVAR group was found occluded at the 12-month follow-up. The LSA was not stented because intraoperative evaluation of the LSA after deployment of fenestrated stent-graft showed favourable results. The CTA showed that the occlusion was near the orifice of the LSA, which was not caused by stent-graft migration.

Post-operative outcomes are listed in Table 3, and Kaplan–Meier estimates of mortality are shown in Figure 5. Ischemic stroke was diagnosed based on symptoms and images. Overall, 4 (1.7%) patients developed stroke—2 (1.6%) in the C-TEVAR group and 2 (2.0%) in the F-TEVAR group. No patient had paraplegia. The median follow-up time for all patients was 28 (range, 16–41) months.

The 15 patients who developed type Ia endoleaks were managed by modifications to oral medication, i.e., by stopping antiplatelet drugs and increasing antihypertensive drugs. Two patients with endoleak in the C-TEVAR group who had refractory chest pain and increasing aortic diameters received re-interventions. The first patient, who underwent the chimney technique (Medtronic 34-34-200 mm and Fluency 8–80 mm for a 30-mm diameter landing zone and a 7-mm diameter LSA), received angioplasty for the chimney endograft one year later, which significantly
reduced the endoleak. The other patient with endoleak received a chimney endograft (Fluency 100–60 mm) poorly attached to the vessel wall of the LSA (10 mm) because of insufficient oversizing. One year later, another endograft was placed distal to the previous chimney endograft, and the endoleak stopped after balloon moulding for both grafts.

Three other patients, including two with retrograde type A aortic dissection (RAAD) and one with compressed IA endograft, received re-interventions. One patient with AD and intramural hematoma involving the aortic arch presented with severe acute renal ischemia, and thus underwent emergency C-TEVAR with Medtronic 38-38-

| Table 1 | Baseline characteristics of patients |
|---------|-------------------------------------|
| Characters | On-the-table fenestration (n=102) | Chimney technique (n=126) | In situ fenestration (n=6) |
| Male | 83 (81.4) | 105 (83.3) | 4 (60.0) |
| Age, mean (SD) | 59.1 (10.9) | 58.0 (13.5) | 55 (4.9) |
| Timing of treatment | | | |
| Elective | 60 (58.8) | 56 (44.4) | 3 (50.0) |
| Emergency | 42 (41.2) | 70 (55.6) | 3 (50.0) |
| Rupture | 3 (7.1) | 5 (7.1) | 0 (0) |
| Visceral ischemia | 15 (35.7) | 24 (34.3) | 1 (33.3) |
| Limb ischemia | 11 (26.2) | 21 (30.0) | 2 (66.7) |
| Renal ischemia | 10 (23.8) | 16 (22.9) | 0 (0) |
| Spinal cord ischemia | 3 (7.1) | 4 (5.7) | 0 (0) |
| Multiple organ ischemia* | 12 (28.6) | 28 (40.0) | 1 (33.3) |
| Indication for treatment | | | |
| Aortic dissection | 37 (36.3) | 56 (44.4) | 6 (100.0) |
| Chronic aortic dissection | 3 (2.9) | 3 (2.4) | 0 (0) |
| Thoracic aortic aneurysm | 14 (13.7) | 21 (16.7) | 0 (0) |
| Penetrating aortic ulcer | 42 (41.2) | 30 (23.8) | 0 (0) |
| Aortic pseudoaneurysm | 8 (7.8) | 17 (13.5) | 0 (0) |
| Type Ia endoleak | 1 (1.0) | 2 (1.6) | 0 (0) |
| Intramural hematoma | 18 (17.6) | 16 (12.7) | 0 (0) |
| Comorbidities | | | |
| Coronary heart disease | 24 (23.5) | 26 (20.6) | 1 (20.0) |
| Diabetes mellitus | 13 (12.7) | 19 (15.1) | 0 (0) |
| Hypertension | 83 (81.4) | 107 (85.7) | 6 (100.0) |
| Hyperlipidemia | 33 (32.4) | 48 (38.1) | 3 (50.0) |
| Heart failure | 5 (4.9) | 7 (5.6) | 0 (0) |
| COPD | 9 (8.8) | 9 (7.1) | 0 (0) |
| Previous stroke | 9 (8.8) | 12 (9.5) | 0 (0) |
| Traumatic aortic injury | 2 (2.0) | 3 (2.4) | 0 (0) |
| Marfan syndrome | 1 (1.0) | 2 (1.6) | 0 (0) |

Values are n (%) unless age are mean (SD). COPD, chronic obstructive pulmonary diseases. *Multiple organ ischemia: ischemia of two or more vital organ systems.
200 mm and Fluency 8–80 mm. The other patient who developed refractory chest pains was found to have RAAD at the 3-month follow-up and received open surgery after 19 months. Another patient was diagnosed with intramural hematoma that progressed to a PAU with refractory pain after 2 weeks. Therefore, the patient was treated by C-TEVAR, with Ankura 36–32–160 mm and Fluency 6–60 mm; however, CTA showed the presence of RAAD one month later. This patient underwent open heart surgery after 7 months. The patient with a compressed endograft of the IA received balloon moulding 1 year later. The above five patients recovered uneventfully after re-interventions.

**Discussion**

Our experience with TEVAR—chimney, on-the-table fenestration, and in-situ fenestration techniques—for aortic arch pathologies showed encouraging outcomes with low mortality and complication rates, indicating that these endovascular techniques were safe and feasible options. Patients recovered well and quickly and benefited from less-invasive treatments.
In our practice, each technique has its strengths and limitations. C-TEVAR is a simple procedure that could adapt to the complex aortic arch anatomy, such as severe tortuosity of the aortic arch and angulation of the supra-aortic branches. Moreover, the chimney technique could be used in bail-out situations like unintentional coverage of branch vessels (12). However, the main limitation of this approach is the susceptibility of endoleaks, because of the so-called ‘gutter’ between stent-grafts. In the present study, 11% patients in the C-TEVAR group developed endoleaks, which was consistent with recent publications that reported 9.9–14.3% and 13.0–19.7% endoleak rates in the single and double chimney techniques, respectively (10,11,13-15). Moreover, chimney endografts are permanently compressed by the thoracic stent-graft which presents a potential threat to the long-term stability and patency of the graft.

### Table 3 Postoperative outcomes

| Variables                              | On-the-table fenestration (n=102) | Chimney technique (n=126) | In situ fenestration (n=6) |
|----------------------------------------|----------------------------------|--------------------------|---------------------------|
| All-cause mortality in F/U            | 5 (4.9)                          | 10 (7.9)                 | 0 (0)                     |
| Cardiac events                        | 2 (2.0)                          | 4 (3.2)                  | –                         |
| Cerebral events                       | 1 (1.0)                          | 2 (1.6)                  | –                         |
| Renal failure                         | 0 (0.0)                          | 2 (1.6)                  | –                         |
| Bowl ischemia                         | 1 (1.0)                          | 1 (0.8)                  | –                         |
| Trauma                                | 1 (1.0)                          | 1 (0.8)                  | –                         |
| Major complications in F/U           | 7 (6.8)                          | 11 (8.7)                 | 0 (0)                     |
| Cardiac events                        | 2 (2.0)                          | 4 (3.2)                  | –                         |
| Stroke                                | 2 (2.0)                          | 2 (1.6)                  | –                         |
| Renal failure                         | 2 (2.0)                          | 2 (1.6)                  | –                         |
| Bowl ischemia                         | 1 (1.0)                          | 1 (0.8)                  | –                         |
| RAAD                                  | 0 (0.0)                          | 2 (1.6)                  | –                         |
| Paraplegia                            | 0 (0.0)                          | 0 (0.0)                  | –                         |
| Total EL-Ia                           | 1 (1.0)                          | 14 (11.1)                | 0 (0)                     |
| Total EL-Ia remained in F/U           | 1 (100.0)                        | 9 (64.3)                 | –                         |
| EL-Ia disappeared after OMT           | 0 (0.0)                          | 3 (21.4)                 | –                         |
| EI-Ia disappeared after redo reintervention | 0 (0.0)                        | 2 (14.3)                 | –                         |
| Diminished EL-Ia after OMT            | 0 (0.0)                          | 3 (21.4)                 | –                         |
| Type II endoleak                      | 0 (0.0)                          | 1 (0.8)                  | 0 (0)                     |
| Type III endoleak                     | 0 (0.0)                          | 0 (0.0)                  | 0 (0)                     |
| Patency of reconstructed branches     | 101 (99.0)                       | 126 (100.0)              | 6 (100.0)                 |
| Re-intervention in F/U                | 0 (0.0)                          | 5 (4.0)                  | 0 (0)                     |
| Surgery for RAAD                      | –                                | 2 (1.6)                  | –                         |
| Management of EL-Ia                   | –                                | 2 (1.6)                  | –                         |
| Management of compressed CG           | –                                | 1 (0.8)                  | –                         |
| F/U time                              | 28 [20–41]                       | 27 [19–39]               | 23 [16–26]                |

Values are n (%) except that time are median months [range]. F/U, follow-up; RAAD, retrograde type A aortic dissection; EL-Ia, type Ia endoleak; OMT, oral medicine treatment; CG, chimney graft.
F-TEVAR is a promising and more judicious technique used to repair the aorta without altering the anatomic structures. In our practice, we took advantage of the characteristics of Ankura stent-grafts, thus making the design, modification, and orientation procedures much more efficient and effective. In the present study, only one of 102 (1.0%) patients developed an endoleak, which was consistent with recent publications that reported endoleak rates of 0–4.2% (5-7). These results might be attributed to our strategy of choosing F-TEVAR more often in cases of aortic arch pathologies that involved only the inner curve of the aorta, as noted in the 41.2% patients treated for penetrating ulcers.

I-TEVAR has shown great potential in endovascular treatment for aortic arch pathologies (9,16). Both C-TEVAR and F-TEVAR are susceptible to endoleaks when the proximal intimal tear of the AD is very close to the branch arteries. I-TEVAR, with a relatively smaller fenestration, may prevent endoleak occurrence. Although the sample size of the I-TEVAR group (n=6) was rather limited, all six patients experienced technical success without endoleaks. However, the I-TEVAR approach had several limitations. First, I-TEVAR required specific anatomic conditions, namely a vertical angle of the target branch to the aortic arch. Tortuosity and sharp angle of the branch vessels could greatly increase the procedural difficulty (9). Second, temporary carotid arteries coverage raises concerns for ischemic complications; hence, I-TEVAR was not the first choice for revascularization of the LCA and IA. Third, I-TEVAR was associated with greater procedural and fluoroscopy times and volume of contrast owing to more procedural steps and technical difficulties than C-TEVAR and F-TEVAR (17).

Therefore, based on the individual strengths and limitations and encouraging outcomes of these three techniques in the early and mid-term period, each of these three techniques has application in endovascular surgery. However, long-term results remain uncertain and need further validation. In future, we hope to report another study of these techniques in a larger cohort and with a longer follow-up period.

**Conclusions**

Total endovascular treatments for patients with aortic arch pathologies are challenging because of the diverse vascular anatomies of both the aorta and the aortic branches. The chimney, fenestrated, and in-situ needle fenestration techniques in TEVAR provided encouraging mid-term results in our selected patient cohort and are hence feasible, effective, and offer a safe alternative to open surgery. Additional long-term studies are required to evaluate the robustness and applicability of these techniques.

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**Footnote**

Conflicts of Interest: The authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The institutional review boards of both Fuwai Hospital, National Centre for Cardiovascular diseases and the Second Xiangya Hospital approved the study, and the need for informed consent was waived (ID: D171100002917004, Dated: January 1, 2016).

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