TYPE B AORTIC DISSECTION MANAGEMENT GUIDELINES

The Society of Thoracic Surgeons/American Association for Thoracic Surgery Clinical Practice Guidelines on the Management of Type B Aortic Dissection

Thomas E. MacGillivray, MD, Thomas G. Gleason, MD, Himanshu J. Patel, MD, Gabriel S. Aldea, MD, Joseph E. Bavaria, MD, Thomas M. Beaver, MD, Edward P. Chen, MD, Martin Czerny, MD, Anthony L. Estrella, MD, Scott Firestone, MS, Michael P. Fischbein, MD, G. Chad Hughes, MD, Dawn S. Hui, MD, Kalie Kissoon, Jennifer S. Lawton, MD, Davide Pacini, MD, T. Brett Reece, MD, Eric E. Roselli, MD, and John Stulak, MD

Houston Methodist DeBakey Heart and Vascular Center, Houston, Texas; Division of Cardiac Surgery, University of Maryland School of Medicine, Baltimore, Maryland; Department of Cardiac Surgery, University of Michigan, Ann Arbor, Michigan; Division of Cardiothoracic Surgery, University of Washington School of Medicine, Seattle, Washington; Division of Cardiovascular Surgery, Hospital of the University of Pennsylvania, Philadelphia, Pennsylvania; Division of Thoracic and Cardiovascular Surgery, University of Florida, Gainesville, Florida; Division of Cardiovascular and Thoracic Surgery, Duke University School of Medicine, Durham, North Carolina; Department of Cardiovascular Surgery, University Heart Center Freiburg-Bad Krozingen, Freiburg, Germany; Department of Cardiothoracic and Vascular Surgery, The University of Texas Health Science Center at Houston and Memorial Hermann Hospital, Houston, Texas; The Society of Thoracic Surgeons, Chicago, Illinois; Department of Cardiothoracic Surgery, Stanford University, School of Medicine, Stanford, California; Department of Cardiothoracic Surgery, University of Texas Health Science Center at San Antonio, San Antonio, Texas; Division of Cardiovascular Surgery, Johns Hopkins University, Baltimore, Maryland; Department of Cardiac Surgery, University of Bologna, Bologna, Italy; Department of Cardiothoracic Surgery, University of Colorado School of Medicine, Aurora, Colorado; Department of Thoracic and Cardiovascular Surgery, Heart, Vascular, and Thoracic Institute, Cleveland Clinic, Cleveland, Ohio; and Department of Cardiovascular Surgery, Mayo Clinic, Rochester, Minnesota

EXECUTIVE SUMMARY

As a result of the rapidly evolving technology and techniques to manage patients with type B aortic dissection (TBAD), The Society of Thoracic Surgeons (STS) and the American Association for Thoracic Surgery (AATS) convened a panel of expert aortic surgeons to methodically review the current data to provide recommendations on the management of patients with TBAD.

MAJOR FINDINGS. Class I recommendations include using a stepwise approach to the evaluation and treatment of patients with TBAD, followed by close clinical surveillance. Optimal medical therapy (OMT) remains the recommended treatment for patients with uncomplicated TBAD. Thoracic endovascular aortic repair (TEVAR) is indicated for complicated hyperacute, acute, or subacute TBADs and favorable anatomy for TEVAR. Open surgical repair is reasonable over TEVAR as a more durable treatment in patients with connective tissue disorders who have TBAD with progression of disease despite OMT. Revascularization of the left subclavian artery (LSA) is recommended to decrease the risk of spinal cord ischemia (SCI) if TEVAR coverage obstructs antegrade LSA flow.
Additional strong recommendations include that open surgical repair should be considered for patients with chronic TBAD with indications for intervention unless comorbidities are prohibitive. TEVAR is reasonable for patients with chronic TBAD with indications for intervention with suitable anatomy but who are at high risk for complications of open repair due to comorbidities.

LIMITATIONS. Overall, the lack of high-quality data from randomized clinical trials limits the ability to make strong evidenced-based recommendations in several high-priority areas, including the optimal timing for intervention on TBAD patients with high-risk features or the routine use of TEVAR in uncomplicated TBAD to prevent long-term aortic-related complications and death. Several recommendations have been based on nonrandomized data or expert consensus.

INTRODUCTION

Aortic dissection is the most common catastrophic aortic event. In the years that followed the seminal report by Levinson and colleagues on 58 autopsy-proven aortic dissections in 1950, Dr Robert Shaw pioneered the concept of fenestration (performed transperitoneally) for an aortic dissection with lower extremity malperfusion. Over time the diagnosis and management of aortic dissection have evolved, resulting in improved outcomes with careful planning and appropriate intervention. Notably, an expectant approach yields essentially the same poor outcome that was seen in the 1950s during these early reports, thus justifying a thoughtful and careful approach that includes patient-specific interventional or open surgical therapies depending on clinical, pathologic, and anatomic features.

Globally, reported incidences of aortic dissection range from 3 to 10 cases per 100,000 patients in the United States to as high as 16 cases per 100,000 patients in Sweden. The true incidence of acute aortic dissection may be underrepresented by these population reports, given that many patients die before reaching a hospital, with the cause of death never proven.

The anatomic extent of the aortic dissection importantly impacts the prognosis and management of the patient. The DeBakey and the Stanford classification schemata are the most commonly used to codify the anatomic extent of disease. Patients with acute type A aortic dissection (TAAD), which comprise approximately two-thirds of all aortic dissections, have a high mortality if not managed with emergent surgery. The available evidence and guideline-directed treatment for patients with TAAD recommends surgery over medical therapy for most patients.

In contradistinction, OMT of uncomplicated (no evidence of rupture or malperfusion) TBAD has historically been the accepted standard, with open surgery reserved for complicated (rupture, malperfusion) cases. However, over the last 2 decades, endovascular therapies have been increasingly used to manage patients with complicated TBAD and those with “distinct features” associated with adverse sequelae. Although stent graft use for TEVAR of acute and chronic TBAD was first approved by the United States Food and Drug Administration in 2013, large clinical trials to guide the appropriate use criteria have to date been limited. Given the therapeutic options now available, this guideline put forth by a joint panel of experts from The Society of Thoracic Surgeons (STS) and the American Association for Thoracic Surgery (AATS) seeks to provide a current framework with which to approach patients with TBAD.

METHODOLOGY

The STS/AATS Guideline Steering Committee proposed the topic for this manuscript and provided the list of authors after a review of conflict of interest disclosures. The topic and authors were approved by the leadership of both societies in 2018. A systematic review to identify the topics of highest priority resulted in 11 questions using the Patient, Intervention, Comparison, Outcome (PICO) format. A search strategy using the PICO questions was performed using MEDLINE and Embase in September 2019 (Appendix 1) and resulted in 704 potentially relevant abstracts. A total of 50 manuscripts met the inclusion criteria and were reviewed for this manuscript, and reference lists were scanned manually for any relevant additional titles. Data were extracted into evidence tables (Appendix 2). Randomized trials and meta-analyses were assessed using a custom checklist for risk of bias (Appendix 3), while nonrandomized data was assessed using the Newcastle-Ottawa scale (Appendix 4).

The recommendations were developed and rated according to the-American College of Cardiology/American Heart Association classification system (Figure 1) using a modified Delphi method. The final manuscript was approved by a joint STS/AATS Guidelines Steering Committee, then independently by the STS Workforce on Evidence Based Surgery, Council Operating Board on Quality, Research, and Patient Safety, and Executive Committee, as well as the AATS Guidelines Committee and Executive Committee.

No commercial entity provided funding or influenced the manuscript in any way. A complete list of conflict of interest disclosures may be found in Appendix 5. STS and AATS are committed to updating this manuscript within 5 years of publication, at which point, this document should be considered expired.

PATHOPHYSIOLOGY OF TBAD

The etiology of aortic dissections is thought to be related to an underlying weakness in the aortic media
aggravated by an intimal tear, ruptured vaso vasorum (intramural hematoma), or a ruptured atherosclerotic plaque/ulcer. Location of the primary entry tear and the extent of aortic propagation determine the clinical course of acute aortic dissection. After an initial aortic wall injury at the intimal/medial level, propagation can occur proximally and/or distally. Although the primary entry tear can usually be allocated to an aortic segment, the proximal/distal propagation and extent of aortic involvement determine the disease classification (whether Stanford or DeBakey), clinical course, and management.

Traditionally, a primary entry tear in the descending aorta at or beyond the LSA, with propagation limited by the LSA proximally, has been classified as a Stanford type B or DeBakey type III aortic dissection. The suprarenal branches often act as an anatomic barrier limiting retrograde propagation. An entry tear located in the aortic arch proximal to the LSA (and may extend further retrograde into the aortic arch but not into the
ascending aorta) is designated a non-A-non-B dissection;16 if it extends further retrograde into the ascending aorta, it is a retrograde Stanford type A or retrograde DeBakey I dissection.13,17,18

The Society for Vascular Surgery and the STS have defined a new classification system that provides a nomenclature to facilitate the description and the reporting of aortic dissection.16 According to this classification system, type A describes any aortic dissection with an entry tear in zone 0, and type B includes any aortic dissection with an entry tear in zone 1 or a more distal aortic zone. In addition to being identified as type B, the dissection is further classified with 2 subscripts B(pd) describing the most proximal zone of involvement (p) and the most distal zone of involvement (d) (Figure 2). For example, a TBAD with an entry tear in zone 4 with retrograde extension to the midaortic arch and antegrade extension to just above the aortic bifurcation would be described as B.2,9 Furthermore, TBAD acuity is defined as (1) complicated, (2) high risk, or (3) uncomplicated (Figure 3).

Malperfusion occurs when a dissection compromises blood flow to an end-organ. An increased incidence of malperfusion has been reported when the primary entry tear originates in angulated aortic segments, such as the distal aortic arch or the proximal descending aorta.5,13-15 Distal propagation of the false lumen in the descending aorta may cause true lumen collapse, resulting in visceral, renal, spinal, and/or extremity malperfusion.14-16 A “malperfusion syndrome” refers to end-organ ischemia of a visceral, renal, lower extremity, brain, or spinal cord vascular bed manifesting with a clinically recognizable pathophysiologic change (eg, bowel ischemia with associated lactic acidosis), in contradistinction to “malperfusion,” which is defined as simply diminished blood flow to the arterial bed of a vital organ by clinical examination (eg, diminished/asymmetric limb pulse) or radiographic imaging.20 Whether a patient is suffering from malperfusion or malperfusion syndrome is a clinical decision based on a combination of clinical, biochemical, and imaging findings.21


FIGURE 3 Aortic dissection acuity.
Aneurysmal formation may occur secondary to presurization and degeneration of the false lumen over time. One hypothesis is that a mismatch between the blood flow into and out of the false lumen may cause increased pressurization of the false lumen. Morphologic features that can impact false lumen pressurization include location and size of the primary entry tear as well as the number and size of communications between lumina. High inflow (large intimal tear) and low outflow (small distal tear with few septal fenestrations) can result in high false lumen mean blood pressure and enlargement.\textsuperscript{22,23}

The fundamental principle of intervention is to exclude the primary entry tear and restore normal blood flow into the true lumen of the aorta and its major branches. Although coverage of the primary entry tear alone is often sufficient, a graft, stent, or stent graft extension may be required to resolve residual true lumen collapse when additional communications exist between lumina. In addition, stent graft-induced new entries (SINEs) may further prevent true lumen expansion. In the case of persistent true lumen collapse at the level of visceral arterial ostia, further true lumen stabilization with a noncovered stent may be helpful. Additional endovascular treatment of major branches is usually not needed once the aortic true lumen is expanded.\textsuperscript{24}

**ACUTE COMPLICATED TBAD**

- **TEVAR** is indicated for complicated hyperacute, acute, or subacute TBADs with rupture and/or malperfusion and favorable anatomy for TEVAR. (Class of Recommendation [COR] I, Level of Evidence [LOE] B-nonrandomized [NR])
- Open surgical repair for complicated hyperacute, acute, or subacute TBADs should be considered for those patients with unsuitable anatomy for TEVAR. (COR IIA, LOE B-NR)
- Fenestration may be considered for complicated hyperacute, acute, or subacute TBADs. (COR IIB, LOE C-limited data [LD])

The chronicity of dissection from the onset of symptoms may impact the risk of morbidity/mortality as well as the efficacy of endovascular therapies.\textsuperscript{25-27} Aortic dissections are now classified based on time from onset of symptoms as hyperacute (<24 hours), acute (1-14 days), subacute (15-90 days), and chronic (>90 days). TBADs are further categorized as “complicated,” “with high-risk features,” or “uncomplicated.”\textsuperscript{15} Complicated dissection is characterized by the presence of malperfusion syndrome or rupture. High-risk features may include refractory pain, refractory hypertension, bloody pleural effusion, aortic diameter >40 mm, imaging evidence of malperfusion, entry tear on lesser curvature, or false lumen >22 mm. Uncomplicated TBADs do not have evidence of rupture, malperfusion syndrome, or high-risk features.

Branch vessel obstruction causing malperfusion syndromes may be dynamic, static, or a combination of both. With dynamic obstruction, hemodynamic forces, such as increased false lumen pressurization, can shift the position of the dissection septum/flap during systole.
causing temporary obstruction, thus decreasing the amount of branch vessel true lumen flow (Figure 4). Dynamic branch vessel compromise can be transient or intermittent, and anti-impulse therapies can mitigate the septic shift and improve true lumen flow. Static obstruction of branch vessels results from a false lumen markedly compressing the true lumen at or near the ostium, with no distal fenestration/reentry tear, intussusception of the flap into the ostium, or false lumen-associated thrombus formation that creates static obstruction (Figure 3). These differing etiologies (static, dynamic, or both) of malperfusion have implications for successful treatment of end-organ ischemia. Among patients with malperfusion syndrome, 80% have dynamic obstruction. In these cases, TEVAR with coverage of the primary tear and any other major communications between the true lumen and false lumen may reverse the malperfusion by restoring sufficient blood flow into the true lumen.

Approximately 20% of patients with acute TBAD will have a malperfusion syndrome, with 5% to 7% having visceral ischemia. In the International Registry for Aortic Dissection (IRAD) data, visceral ischemia was strongly associated with in-hospital mortality of 30.8% vs 9.1% without ischemia (odds ratio [OR], 3.33; \( P < .0001 \)). TBAD with malperfusion is increasingly treated with endovascular therapies, from 35% in the early IRAD era 1996-2001 to 68% in the later IRAD era 2008-2013. Attendant with this shift, open surgery decreased from 47% to 18%. Mortality in the context of visceral ischemia was similar between open and endovascular repair (25.8% vs 25.5%, \( P \) nonsignificant).

TEVAR can expeditiously control a rupture or cover the primary entry tear to restore true lumen flow, resulting in reduced ischemic time and improvement of outcomes over medical management alone or combined with open surgery. Collectively, accrued data demonstrate improved outcomes with TEVAR for complicated TBAD compared with open surgery or medical therapy alone. Consequently, TEVAR has become the first-line treatment for complicated TBAD. However, the success of TEVAR alone to reestablish arterial end-organ flow is dependent on the underlying etiology of the malperfusion.

Several published series have described endovascular management of complicated TBAD, but few reports specifically detail the management or outcomes for the subset of patients with visceral malperfusion syndromes. Of those studies that detail malperfused regions, the proportion of visceral malperfusion ranges from 7.6% to 60%. Two series reported no need for adjunctive branch stenting or fenestration, but one reported a post-TEVAR colon resection, and the other reported a death at postoperative day 11 from persistent visceral ischemia.

A report from 2 high-volume European centers described 41 complicated TBAD patients, with an overall 41% branch vessel stenting rate. Nearly a quarter (\( n = 4 \) (23.5%)) of those with visceral malperfusion had branch stenting or fenestration. The 30-day mortality rate was 17.1%, with 2 deaths due to bowel infarction, and 3 patients underwent bowel resection without further visceral revascularization.

Three other series of complicated TBAD reported overall adjunctive branch stenting rates of 13.7% to 22%, including visceral branches and no cases of bowel resection or bowel-related mortality. In the most recent IRAD report, of the 51 acute TBAD patients with visceral ischemia, 63% underwent TEVAR, 31% underwent fenestration, and 33% underwent branch vessel stenting.

The University of Michigan group has reported their series of 182 patients using branch stenting and fenestration (without TEVAR) as the primary strategy for
acute TBAD complicated by malperfusion syndrome (ie, without rupture). The rationale is that this strategy can treat both dynamic and static obstruction while avoiding the risks of TEVAR, including retrograde type A dissection, neurologic complications of stroke and SCI, graft infection in the setting of necrotic tissue, and coverage of the LSA. This approach was first described to mitigate the high operative mortality of acute TAAD dissection with visceral ischemia. The mechanism by which this strategy works is to introduce a reentry tear into the distal aorta and stabilize the flap motion to prevent dynamic obstruction. Any residual static branch vessel obstruction is then treated by branch artery stenting. Over a 22-year period (1996-2018), the Michigan group reported a 7.7% mortality (no deaths in the last 8 years) and 0% paralysis.

By “converting” acute TBAD with malperfusion to anatomic features associated with uncomplicated TBAD, there remains a persistent risk for aortic rupture and growth, unlike what seen when using TEVAR, which can not only address the malperfusion but also treat the thoracic aorta. Indeed, reintervention rates for the fenestration and stenting approach have been reported as 21% at 5 years and 31% at 10 years. These concerns, as well as a lack of expertise with successful fenestration, have limited widespread adoption in many centers. Stent-assisted balloon-induced disruption and relamination in aortic dissection (STABILISE technique) has shown promising early results in achieving complete repair of the dissected aorta by inducing complete false lumen obliteration in several small series of patients.

UNCOMPlicated TBAD

- A stepwise approach to the evaluation and treatment of acute/subacute uncomplicated TBAD should be applied that includes identification of the primary entry tear site location, defining the proximity and distance of the dissection to the LSA, calibration of the maximum orthogonal aortic diameter, and confirmation of the lack of any organ malperfusion or other indications of complicated disease. (COR I, LOE B- NR)
- OMT is the recommended treatment for patients with uncomplicated TBAD. (COR I, LOE B- NR)
- Prophylactic TEVAR may be considered in patients with uncomplicated TBAD to reduce late aortic-related adverse events and aortic-related death. (COR IIB, LOE B-NR)
- Close clinical follow-up after hospital discharge is recommended for patients presenting with acute TBAD. (COR I, LOE B-NR)

TBAD has been regarded as having a more benign natural history compared with TAAD. OMT implies sustained anti-impulse therapy for control of both hypertension and heart rate while also limiting the maximum change in left ventricular pressure during early systole (ie, maximum $dP/dt$). Maintaining blood pressure $\leq 120/80$ mm Hg and heart rate $< 70$ beats/min are optimal, and $\alpha$- and $\beta$-blockers are useful primary agents. Once heart rate control is established, angiotensin-converting enzyme inhibitors, angiotensin receptor blockers, and/or dihydropyridine calcium channel blockers may also be useful.

However, close surveillance has shown that over time, a high percentage of patients with TBAD will experience subsequent sequelae resulting in death or requiring intervention. Single-center, clinical trial, and registry data have reported aneurysmal degeneration rates in excess of 70% and mortality rates approaching 25% to 30% at 3 to 5 years. Aneurysmal aortic degeneration is the prominent indication for intervention in the chronic phase of the disease. The presence of certain morphologic features (size and location of luminal tear or fenestrations) and compliance with OMT are associated with the development of complications requiring subsequent intervention.

TEVAR VS OMT FOR UNCOMPlicated TBAD. INSTEAd trial. The INvestigation of STEnt Grafts in Aortic Dissection (INSTEAD trial) prospectively compared prophylactic TEVAR plus OMT to OMT alone in patients with uncomplicated TBAD who were stable for the first 2 weeks from the onset of symptoms. Between 2 and 52 weeks (subacute and chronic phase), study patients were randomized to 1 of the 2 therapeutic cohorts. The primary end point was all cause mortality at 2 years. Secondary end points were aorta-related death and a composite of progressive aortic pathology and morphologic evidence of aortic remodeling (true lumen recovery or false lumen shrinkage and false lumen thrombosis). Although favorable aortic remodeling occurred in the TEVAR/OMT cohort (91.3%) compared with the OMT cohort (94.4%), there were no differences in the primary end point of all-cause mortality or aorta-related mortality at 2 years. The trial and its design were criticized for being underpowered, the measured outcome time too short, and the crossover rate from OMT to TEVAR/OMT too high (16.2%). Notwithstanding these criticisms, the patients were subsequently monitored out to 5 years (INSTEAD-XL). At 5 years, TEVAR/OMT was associated with improved aorta-specific survival and delayed disease progression, although these outcome measures were established post hoc.

ADSORB trial. The ADSORB trial (Acute Dissection Stent Grafting or Best Medical Treatment) compared OMT to OMT plus TEVAR in patients with acute, uncomplicated TBAD. The primary end point was a combination of incomplete/false lumen thrombosis, aortic dilatation, or aortic rupture at 1 year. The conclusion was that...
TABLE 1 Morphologic Features Posing High Risk of Late Sequelae

<table>
<thead>
<tr>
<th>Feature</th>
<th>Risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary entry tear at greater curve of distal arch</td>
<td>Proximal aortic dilatation, aortic arch remodeling, false lumen</td>
</tr>
<tr>
<td>Short proximity of entry tear to left subclavian artery ostium</td>
<td>compression, true lumen expansion, thrombosis, dissection propagation</td>
</tr>
<tr>
<td>Initial aortic diameter ≥20 mm</td>
<td>in the aortic arch</td>
</tr>
<tr>
<td>Initial false lumen diameter ≥22 mm</td>
<td></td>
</tr>
<tr>
<td>Number/size of fenestrations between true and false lumen</td>
<td></td>
</tr>
<tr>
<td>Stent graft-induced new entry</td>
<td></td>
</tr>
<tr>
<td>Partial false lumen thrombosis</td>
<td></td>
</tr>
</tbody>
</table>

remodeling with thrombosis of the false lumen and reduction of its diameter was induced by stent grafting, but long-term outcome comparisons are needed. These longer-term end points have not yet been reported.

Recognizing the significant methodologic limitations of these 2 randomized trials, other observational studies are also relevant. For example, Iannuzzi and colleagues compared 8717 OMT patients with 266 patients who underwent TEVAR and 182 patients who underwent open surgery. The 5-year survival was 59.8% in OMT patients, 66.7% for those undergoing open surgery, and 75.9% in TEVAR patients (TEVAR vs OMT hazard ratio, 0.68; 95% CI, 0.55-0.83; P < .01). Patients were not matched in this analysis, with TEVAR and surgery patients being significantly younger and healthier than OMT patients. Further, the median duration of follow-up for TEVAR patients was only 1.5 years.

Similarly, Qin and colleagues compared 154 OMT patients and 184 TEVAR patients and found a favorable 5-year survival estimate in TEVAR patients (89.2% vs 85.7%; log-rank P = .01). The study of 4706 patients by Shah and colleagues, focused on safety outcomes, showed no significant difference in 30-day mortality between groups, but TEVAR patients were at a 61% increased odds of stroke (8.1% vs 4.6%; OR, 1.61; 95% CI, 1.14-2.27; P = .0073).

Some caution is warranted in interpreting these studies, because treatment selection was not randomized, and potential confounding variables were not accounted for apart from age in the report by Shah and colleagues.

NATURAL HISTORY OF UNCOMPPLICATED TBAD. After the acute phase, the dissection flap stiffens, and the dissection transitions into its chronic phase. The stiff, scarred dissection flap renders the aorta less responsive to false lumen compression and true lumen expansion by subsequent TEVAR in the chronic phase, and thus, TEVAR may be less effective in imparting favorable remodeling, as has been observed in the acute and subacute phases. However, it is rare that complications such as malperfusion or retrograde propagation of the dissection occur in the aortic arch in the chronic phase. Progressive aortic dilatation is the most common sequela during the chronic phase. Proponents of TEVAR in the acute and subacute phases argue that earlier intervention may mitigate the incidence of aneurysmal progression during the chronic phase.

TREATMENT PARADIGM FOR ACUTE UNCOMPPLICATED TBAD. OMT is the current standard of care treatment for all patients with uncomplicated TBAD. A thorough morphologic analysis of the aortic imaging enables identification of patients with high-risk features who may benefit from early TEVAR in addition to OMT. The INSTEAD-XL trial provides the best available evidence supporting TEVAR plus OMT, as outlined previously. Thus, in stable TBAD with suitable anatomy and high-risk features, preemptive TEVAR may be considered to improve late outcome.

MORPHOLOGIC FEATURES POSING HIGHER RISK OF LATE SEQUELAE

A primary entry tear location at the greater curve of the distal arch may portend a higher risk of malperfusion and delayed aneurysmal dilatation. Proximity of the primary entry tear to the LSA ostium also appears to impact the development of complications during the initial 14 days after onset of symptoms, where a shorter distance to the LSA ostium is associated with higher risk.

An initial total aortic diameter of ≥40 mm or an initial false lumen diameter of ≥22 mm have each been identified as independent predictors of subsequent aneurysmal dilatation. The number and size of fenestrations (tears) between the true and false lumens in the thoracoabdominal aorta may be predictive of aneurysmal dilatation over time. Even after TEVAR, a large secondary or reentry tear distal to the TEVAR (eg, a distal SINE), can predict subsequent thoracoabdominal dilatation. Partial (incomplete) false lumen thrombosis has also been identified as an independent predictor of adverse outcome in TBAD (Table 1).

RISK OF RETROGRADE TAAD. In uncomplicated TBAD, prophylactic TEVAR aims to prevent anticipated late adverse aortic events. However, delayed retrograde TAAD after TEVAR for TBAD is a recognized potentially lethal complication with an incidence of 1.3% to 11%. Risk factors for retrograde TAAD after TEVAR include stent graft oversizing, use of a proximal bare-spring stent graft, aortic arch dilatation, a proximal tear site within the arch, notable “bird’s beaking” (failure of apposition of the proximal end of the stent graft at the lesser curve), and stent graft landing proximal to the LSA. The frozen elephant trunk procedure has been applied in patients with TBAD qualifying for treatment and at high risk for retrograde TAAD.
approach has been used for a carefully selected subgroup of patients at high risk for a proximal type I endoleak or retrograde TAAD.104

CHRONIC TBAD

- Open surgical repair should be considered for patients with chronic TBAD with indications for intervention, unless comorbidities are prohibitive or anatomy is not suitable for TEVAR. (COR IIA, LOE B-NR)
- TEVAR is reasonable for patients with chronic TBAD with an indication for intervention with suitable anatomy (adequate landing zone, absence of ascending or arch aneurysm) but who are at high risk for complications of open repair due to comorbidities. (COR IIA, LOE B-NR)
- TEVAR alone as sole therapy is not recommended in patients with chronic abdominal aortic aneurysm, an inadequate distal landing zone, and/or large distal reentry tears. (COR III: No benefit, LOE C-LD)

**INDICATIONS FOR INTERVENTION.** Indications for elective intervention in the chronic setting include aneurysmal dilatation (total ≥55-60 mm), increasing rate of diameter (>10 mm/y), and/or symptoms (pain, malperfusion).99-98 Acute redissection or rupture (ie, acute aortic syndrome) presentation should invoke intervention as appropriate for acute aortic dissection.

**TREATMENT: OPEN.** Patients with aneurysmal degeneration of a chronic TBAD typically have involvement extending through the visceral segment of the aorta (DeBakey type IIB) and often into the iliac vessels as well (zone 10). The portion of aorta affected by aneurysm may involve the entirety of the thoracoabdominal aorta or be limited to the more proximal descending aorta.

In open thoracoabdominal aortic operations, chronic dissection per se has not been shown to be a specific risk factor for stroke or paraplegia compared with non-dissected aneurysms.99 While a staged approach to repair has been shown to reduce neurologic complications by reducing ischemia to the spinal cord,100 performing a limited descending replacement for the aneurysm and leaving distal dissected aorta in the thoracoabdominal aorta leaves the patient vulnerable to further aneurysmal degeneration. The endovascular options for treating dissected aneurysms of the thoracoabdominal aorta are still quite limited by access and anatomy. When choosing an open approach, there are limited data to guide options about extent of repair. Pujara and colleagues101 showed reasonable early outcomes (8% mortality) and poor late outcomes (47%-51% event-free 5-year survival) after descending or thoracoabdominal repairs for chronic TBAD.

These operations are performed similarly to what has been described for aneurysm alone,102,103 with some additional considerations given the morphologic changes caused by aortic dissection. The operation is usually performed with cardiopulmonary bypass support (hypothermic circulatory arrest or partial left heart bypass) and cerebrospinal fluid (CSF) drainage.4 The decision about type and conduct of cardiopulmonary bypass support may be based on surgeon preference or the ability to cross-clamp the aorta proximal to the dissection, which most often originates at the LSA. Reimplantation of segmental intercostal branches may be difficult because the dissection can often involve or compromise the intercostal arterial ostia. Similarly, dissection that involves visceral branch vessels may make reconstruction more complex or require distal aortic or ostial branch vessel fenestration. Direct bypasses to the visceral and renal vessels may reduce the risk of late visceral patch dilatation.104

**THORACIC ENDOVASCULAR AORTIC REPAIR.** For the treatment of chronic TBAD with aneurysmal degeneration, TEVAR is less invasive and may potentially reduce periprocedural morbidity and mortality. However, the mid-and long-term fate of the aorta and need for aortic reintervention are of particular concern for chronic TBADs treated with TEVAR.105 The dissection flap, which is thin and dynamic in the acute setting, becomes fibrotic, thickened, and less mobile over time. These changes in flap characteristics may adversely affect the ability to fully expand a stent graft and consequently the true lumen, leading to persistent false lumen flow and reduced capacity for aortic remodeling (Figure 4).20 TEVAR for chronic dissection may facilitate subsequent, more limited thoracoabdominal repair as a staged completion with extension from the distal end of a TEVAR.106-108

There are no published randomized trials of open surgery compared with TEVAR for chronic TBAD. Two institutional studies retrospectively compared contemporaneous open and TEVAR cohorts for TBAD only. Of these, a propensity score-matched analysis found no difference in rates of spinal cord injury, renal replacement therapy, 30-day mortality, or 5-year survival. Treatment efficacy, defined as freedom from aortic rupture or reintervention, was superior with open therapy (96.7% vs 87.5%; hazard ratio, 4.6; $p = .025$).109 The second study found a higher incidence of spinal cord injury with open repair; otherwise, there were no differences in perioperative morbidity, survival at 1 or 5 years, or a composite outcome of freedom from aortic reintervention, rupture, or aortic-related death at 1 or 5 years.110
Late outcomes reported by TEVAR studies are widely heterogenous with regard to their chosen primary and composite end points and may have been impacted by selection bias resulting in dissimilar cohorts. When including only patients who had anatomy suitable for TEVAR, freedom from aortic reintervention appears superior with open therapy (99%-99% vs 53%-87% at 1 year and 79%-93% vs 73%-78% at 5 years), although direct comparisons are currently not available, and most TEVAR series did not report outcomes beyond 5 years. In a study by Pujara and colleagues, mortality was higher than in other open series, but this population also included 42% of patients undergoing urgent or emergency repair. Several TEVAR series report an evolution in technique over the study period, with changes in the extent of aortic coverage, sequence of stent implantation, endograft sizing, and use of tapered grafts that may have also impacted the results. Others report a consistently conservative approach of limited aortic coverage.

Additional endovascular approaches have been used as adjuncts to TEVAR to promote late reverse remodeling or to salvage late failures after the initial TEVAR. These adjunctive techniques, including ballooning a larger segment of the stent graft (ie, Knickerbocker technique), uncovered stent-assisted balloon dilatation (ie, Provisional Extension To Induce Complete Attachment (PETTICOAT) technique), false lumen embolization strategies, and the use of investigational or homemade branched endograft devices have been performed with reasonable success in single-center series with careful patient selection.

**TIMING OF INTERVENTION**

- In patients with acute uncomplicated TBAD with high-risk features, it may be reasonable to consider delaying treatment (beyond 24 hours up to 90 days) with TEVAR to reduce early adverse events and to improve late outcomes. (COR IIB, LOE C-LD)

Patients without frank or contained rupture, severe clinical or radiologic malperfusion, and/or other higher-risk features, who are stable enough to wait for intervention, may benefit from delayed TEVAR from beyond 24 hours up to as many as 90 days due to a lower risk of periprocedural complications. Miyai and colleagues showed that hyperacute patients had significantly higher in-hospital (14.9% vs 0% acute vs 2.8% subacute; \( P < .001 \)) and 30-day mortality than the acute or subacute groups (11.9% vs 0% vs 1.7%; \( P < .001 \)). Interpreting these studies collectively requires an understanding that different indications were used for intervention across the hyperacute and other groups. Regardless, these data underscore the value of interval surveillance imaging to identify impactful aortic changes. Change in aortic morphology (expanding diameter >4 mm, new onset of periaortic hematoma, and/or hemothorax) were found to be associated with poorer prognosis in the subacute phase. Optimal timing of TEVAR for patients with TBAD should be individualized according to the presenting or evolving clinical and/or radiologic features.

**CONNECTIVE TISSUE DISORDERS**

- **Open surgical repair over TEVAR is reasonable for more durable treatment in patients with connective tissue disorders and TBAD who have progression of disease despite OMT.** (COR I, LOE B-NR)

- **TEVAR is reasonable in patients with connective tissue disorders with acute complicated TBADs and anatomy favorable for TEVAR as a bridge to delayed open reconstruction.** (COR IIA, LOE C-LD)

Connective tissue disorders, including Marfan, Loeys-Dietz, and Ehlers-Danlos type IV syndromes, are characterized by genetic mutations resulting in inherent deficiencies in the strength of the connective tissues, including the aorta. Marfan syndrome is the most common connective tissue disease that affects the aorta, with an incidence of 1 in 5000 individuals. Patients afflicted with connective tissue disorders are more prone to develop and die from aneurysms and dissections of the aorta. Data to guide management recommendations for patients with connective tissue disorder who present with acute TBAD are generally limited to subgroup analyses from patient cohorts that are largely without connective tissue disorders.

According to IRAD, patients with Marfan syndrome represent <5% of all patients with acute aortic dissection, including 4% of acute TBAD. In another analysis of IRAD data, Marfan patients with acute TBAD were younger (40.3 ± 12.9 years vs 64.3 ± 13.7 years) relative to those without a connective tissue disorder. The 94 Marfan patients in the study presenting with acute TBAD were less likely to be treated with only OMT (50% vs 62.6%), more likely to be treated with open surgery (28.7% vs 9.7%), and equally likely to be treated with endovascular therapy (19.1% vs 25.3%) compared with a cohort of patients without Marfan syndrome. Freedom from reintervention was significantly worse in the Marfan patients (44.7% vs 81.5%, \( P < .001 \)).

Among the subgroups of patients with connective tissue disorders who present with TBAD, those with Ehlers-Danlos and Loeys-Dietz syndromes have the highest risk of morbidity and mortality both with and without interventional therapy. They experience rapid progression and high complication rates that justify extremely careful planning for operative or endovascular intervention.
Optimal medical treatment of patients with Marfan syndrome has been considered vital to decreasing the rate of aortic growth and the risk of dissection. Although β-blockers have been considered primarily indicated for patients with Marfan syndrome,126 angiotensin receptor blockers appear equally effective as β-blockers in young adults and children with Marfan syndrome.4,123-126 In Marfan patients with acute TBAD, OMT was equally effective in preventing in hospital mortality compared with non-Marfan patients (4.3% vs 7.8% \(P = .576\)).126

In one retrospective study, Marfan patients with acute type B dissections had significantly better survival and freedom from morbidity than non-Marfan patients treated with surgery.131 Of the 27 Marfan patients who were treated with open surgery, there were no deaths (0%) compared with a 17.6% mortality with open surgery in the non-Marfan patients (\(P = .011\)). Neurologic complications did not significantly differ between groups (7.7% vs 13.3%, \(P = .542\)).

Data regarding the use of TEVAR to treat connective tissue disorder patients with TBAD are limited to data registries and small series. Pacini and colleagues139 performed a comprehensive review of 5572 studies to assess the early and late results of TEVAR in Marfan patients with acute and chronic TBAD, revealing 12 citations with data on 54 patients (11 acute and 43 chronic). The procedural incidence of mortality, stroke, and paralysis were all 1.9%. The need for open surgical conversion was 5%. The overall rate of endoleak was 22% (type I, 16%; type II, 4%; type III, 2%), occurring in 12% of acute patients and 31% of chronic patients. At a median follow-up of 2.5 years, the mortality rate was 13% in this group of patients with an average age of 41 years.

The risk of retrograde TAAD during or after TEVAR for TBAD is reportedly higher in Marfan patients.140 Dong and colleagues140 reported outcomes of 443 patients with acute TBAD treated with TEVAR, and retrograde TAAD developed in 11 patients. Of the 6 Marfan patients in that series, a retrograde TAAD developed in 3 of 4 patients who did not have a previous ascending aortic graft. Two of the 3 patients died, and the remaining patient was lost to follow-up. In the European Registry on Endovascular Aortic Repair Complications, 83% of the reported retrograde TAADs occurred in patients treated for either acute or chronic TBAD, of which there was a relatively high proportion of Marfan patients.92 It has been suggested that use of endovascular therapy for TBAD in patients with Marfan syndrome is best considered only when previous graft replacement of the aorta allows for a safe proximal landing zone.129

The National Registry of Genetically Triggered Thoracic Aortic Aneurysm and Cardiovascular Conditions (GenTAC) reported outcomes of 22 TBAD patients treated with TEVAR, which included 10 patients with genetically triggered aortic disease (7 with Marfan and 1 each with Loeys-Dietz, Ehlers-Danlos, and mutation in the actin alpha 2, smooth muscle [ACTA2] gene).141 Retrograde TAAD occurred in 3 patients (25%), and 42% of the patients required reintervention at a median follow-up of 7 months.

Eid-Lidt and colleagues142 reported that among 10 Marfan patients treated with TEVAR for chronic TBAD, at a median follow-up of 59.6 months, endoleaks occurred in 44.4% and reintervention was required in 33.3%. One patient died of aortic rupture at 5 days and another died 9 months postprocedure. Ince and colleagues143 reported the use of TEVAR in 6 Marfan patients with TBAD, of whom 5 had undergone previous ascending aortic replacement. There was technical success in all 6 patients; however, remodeling and resolution of the dissection occurred in only 2 patients. Elective open surgical reconstruction was required in 2 patients and was being considered in a third. One patient died.

SINE tears are more common in Marfan patients treated with TEVAR. Dong and colleagues144 reported the incidence of SINE was 10 times higher in Marfan patients with type B dissection compared with non-Marfan patients (33.33% vs. 3.26%). Weng and colleagues145 reported that SINE developed in 5 of 8 patients (62.5%) with Marfan syndrome treated by TEVAR for TAAD or TBAD.

In Marfan patients with previous proximal aortic surgery, TEVAR has been successfully used to treat TBAD.146,147 Botta and colleagues146 reported 100% procedural success in 12 Marfan patients (acute, \(n = 5\); chronic, \(n = 7\)) with no deaths or strokes. At a median follow-up of 31 months, 1 patient underwent open surgery, and 2 other patients had distal extension of their dissection. Waterman and colleagues147 reported the results of 16 Marfan patients with previous open ascending and/or arch replacement who underwent TEVAR for descending aortic pathology. Primary treatment failure occurred in 7 patients (44%), comprising type I endoleak in 3 and 1 patient each with persistent false lumen flow, retrograde dissection, rupture, and type II endoleak.

The incidence of late conversion for open surgery after TEVAR is relatively high in patients with connective tissue disorders. Among a multcenter registry of 421 patients (Marfan, \(n = 15\)) with a median follow-up of 17 months, one-third of Marfan patients required stent graft explant.148 Similarly, Spiliotopoulos and colleagues149 reported 16 of 45 patients with complications after previous TEVAR had connective tissue disorders (Marfan, \(n = 14\); Loeys-Dietz, \(n = 2\)).

**SPINAL CORD PROTECTION ADJUNCTS TO TEVAR**

- Revascularization (open surgical or endovascular) of the LSA after TEVAR coverage that obstructs
antegrade LSA flow is recommended to decrease the risk of SCI. (COR I, LOE B-NR)
• It is reasonable to establish CSF drainage in type B dissection patients undergoing TEVAR if they are at increased risk for SCI (eg, coverage >20 cm or within 2 cm of the celiac artery origin or other risk factors) and time permits (ie, nonemergent circumstances). (COR IIA, LOE B-NR)
• It is reasonable to establish CSF drainage in type B dissection patients who develop symptoms of paraparesis/paraplegia. (COR IIA, LOE B-NR)

SPINAL CORD BLOOD SUPPLY. SCI represents one of the most devastating complications of thoracic aortic disease. SCI occurs in up to 5.8% of intervention patients in large TEVAR registries, including 4.1% in chronic aneurysmal TBAD, 5.3% in acute TBAD, and 5.8% in intramural hematoma.150-153 Increased stent graft coverage of the descending thoracic aorta (>200 mm) and distal coverage within 20 mm of the celiac artery have been implicated as risk factors for SCI.154 Other identified risk factors for SCI with TEVAR include age, chronic obstructive pulmonary disease, hypertension, emergency, heart and kidney disease, and prior aortic surgery.153,155 The spinal cord collateral network can be impaired by coverage of the LSA or hypogastric artery or by prior abdominal aortic surgery, increasing the risk of SCI.151,153,156 Permanent SCI has predictably devastating consequences on long-term mortality (as high as 75% at 1 year in TEVAR patients who show no neurologic improvement after SCI).153,157 The spinal cord blood supply consists of longitudinal arterial trunks as well as segmental arteries.158-162 In >90% of people, the anterior spinal artery diminishes in size as it descends caudally163 and requires additional arterial supply through the radiculomedullary branches of the segmental arteries. This accounts for the fact that the lower thoracic or thoracolumbar spinal cord is at risk during open descending thoracic and thoracoabdominal aortic repair. As such, the arterial supply to the spinal cord originates from only a few of the segmental arteries. Further, the number of segmental arteries supplying the paired posterior spinal arteries far exceeds those supplying the unpaired anterior spinal artery, which accounts for susceptibility of the anterior cord to ischemic insult with aortic repair.

The most important branch vessel feeding the anterior spinal artery is the great anterior radiculomedullary artery, also known as the artery of Adamkiewicz (AKA). This artery, which can be identified preoperatively using high-resolution computed tomography angiography,162,164 forms a characteristic “hairpin” loop when it reaches the anterior spinal artery. An important anatomic feature of the AKA is that it typically (85%) arises from the left side of the thoracoabdominal aorta and that the level of origin may be as high as T5 or as low as L2. Therefore, the AKA may arise along nearly the entire length of the aorta involved in an extent II thoracoabdominal repair. The origin is most commonly between T9 and L2 (75% of cases), T5 and T8 (15% of cases), and L1and L2 (10% of cases).161 Most people (74%) have only 1 AKA, but 26% of patients may have 2 or even 3 AKAs.165 Importantly, the LSA represents the primary source of collateral pathways to the AKA outside of the spinal column through the thoracodorsal and internal thoracic arteries, which provides basis for adjunctive LSA revascularization during TEVAR.166 Other important collateral sources include segmental arteries distal to the segmental artery feeding the AKA, the hypogastric arteries, and the left external iliac artery (Figure 6).10,167

LSA REvascularIZATION. As highlighted in a recent Cochrane review,168 there are currently no randomized...
controlled trials examining LSA revascularization after zone 2 TEVAR, although nonrandomized evidence exists to support LSA revascularization as a means to prevent SCI with mechanistic explanations of such benefit. A recent meta-analysis of 16 cohort studies with a total of 2591 zone 2 TEVAR patients found that LSA revascularization was associated with a significantly lower perioperative stroke rate (relative risk, 0.61; 95% CI, 0.45-0.82; \( I^2 = 20 \% \)) and SCI rate (relative risk, 0.59; 95% CI, 0.39-0.90; \( I^2 = 0 \% \)) and recommended consideration of revascularization for patients with LSA coverage during TEVAR. Another smaller meta-analysis of 5 observational studies and 1161 patients found an OR of 0.56 (\( P = .09 \)) for SCI after zone 2 TEVAR in patients with vs without revascularization. The data in that analysis did not support LSA revascularization for stroke prevention, and other large data sets, including the National Surgical Quality Improvement Program registry, and prior meta-analyses have found no benefit with regard to stroke prevention for LSA revascularization in patients undergoing zone 2 TEVAR, perhaps because strokes during zone 2 TEVAR are generally embolic in nature.

Currently available techniques for LSA revascularization in conjunction with TEVAR include surgical carotid-subclavian bypass, carotid-subclavian transposition, and carotid-axillary bypass, along with endovascular techniques, including chimney grafts, scallops, fenestrated grafts, and branched grafts. Surgical revascularization techniques are associated with not insignificant risks of phrenic and recurrent laryngeal nerve palsy, although these risks may be decreased with carotid-axillary bypass due to avoidance of any manipulation in the vicinity of the phrenic nerve. Carotid-subclavian transposition should be avoided in patients with a patent pedicled left internal mammary artery bypass graft due to the risk of myocardial ischemia during the mandatory period of proximal LSA clamp with this procedure. Although limited long-term data are available, late revascularization patency may be superior with carotid-subclavian transposition compared with the bypass techniques.

Recent data from the Society for Vascular Surgery Vascular Quality Initiative comparing open surgical and endovascular LSA revascularization techniques in 837 patients during zone 2 TEVAR found similar perioperative outcomes for open vs endovascular revascularization. Long-term comparative effectiveness data are currently lacking. A summary of current indications for LSA revascularization is found in Table 2.174,179,180

### TABLE 2 Indications for Left Subclavian Artery Revascularization Before Zone 2 Thoracic Endovascular Aortic Repair

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Presence of left internal thoracic artery bypass graft</td>
<td>In patients at risk for neurologic complications</td>
<td>Left vertebral artery originating directly from the arch</td>
</tr>
<tr>
<td>Termination of left vertebral artery at posterior inferior cerebellar artery or other discontinuity of vertebralbasilar collaterals</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Functioning arteriovenous dialysis fistula in left arm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prior infrarenal aortic repair with occlusion of lumbar and middle sacral arteries</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Planned long-segment (20 cm) coverage of the descending thoracic aorta where critical intercostal arteries originate</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypogastric artery occlusion</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Presence of early aneurysmal changes that may require subsequent therapy involving the distal thoracic aorta</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

CSF DRAINAGE IN TEVAR FOR TYPE B DISSECTION. In effort to maintain collateral flow to the spinal cord and prevent SCI, various adjuncts have been used during TEVAR deployment, including LSA revascularization, permissive hypertension, and CSF drainage. Spinal cord perfusion pressure (SCPP) is dependent on the systemic mean arterial pressure (MAP) minus the pressure within the spinal canal (SCPP = MAP – intracranial pressure). Drainage of CSF allows for optimizing SCPP because SCI can manifest immediately or days after an ischemic insult. Many patients with delayed presentation of paraparesis can be successfully recovered with reinsertion of CSF drainage and elevating MAP with vasopressors to improve SCPP.

CSF drainage protocols during TEVAR include preoperative intradural catheter placement in patients with identified SCI risk factors, such as large coverage extent (>200 mm, coverage within 20 mm of the celiac artery, prior abdominal aortic aneurysm repair, hypertension, older age, chronic kidney disease, and chronic obstructive pulmonary disease. CSF is intermittently drained as needed to maintain a spinal canal pressure of 10 mm Hg (or 14 cm H2O) for 24 hours or longer, and then clamped for an additional 24 hours before removal. Patients exhibiting signs of SCI can have additional CSF aspirated...
and/or the drain can be lowered with monitoring for improvement in symptoms. In symptomatic patients, the drain can remain in place for an additional 72 hours after stabilization of the neurologic examination.184

Data on CSF drainage specifically in TEVAR for TBAD are limited. However, a randomized trial found CSF drainage was beneficial during open repair of thoracoabdominal aortic aneurysms.186 Notably, there is much more lability in blood pressure during open surgery, and a recent protocol in TEVAR patients emphasizing revascularization to ensure LSA perfusion, permissive hypertension, motor evoked potential monitoring, and no CSF drainage resulted in no SCI in 223 patients.187

However, other centers have emphasized CSF drains, including Maier and colleagues,184 who used CSF drains for TEVAR patients felt to be at higher risk from coverage of 2 perfusion networks (eg, intercostal arteries and LSA) in 116 of 223 patients (52%). They found SCI in only 1 of 116 patients (0.8%) who had CSF drainage compared with 5 of 107 patients (4.7%) who did not have CSF drainage.184 The authors stated the number needed to treat 1 SCI was 26, and reported only 11 minor complications (10%). Based on their experience, they adopted universal CSF drainage for all patients except emergency procedures, patients on anticoagulation, or patients with intracranial disease.184

Acher and colleagues188 have also used an aggressive policy to use CSF drainage <8 mm Hg during TEVAR whenever >12 cm aorta or T8 to L2 are covered, along with a multimodal protocol of hypothermia (34 °C), MAP >100 mm Hg, and naloxone and steroids, reporting only 1 of 115 patients (0.6%) with temporary paresis. Mazzefi and colleagues185 reported using CSF drains in 102 patients, including 30 type B dissections defined as high risk (>150 mm coverage, prior TEVAR or EVAR, or poor pelvic perfusion) and found SCI in 4 patients, which resolved completely in 2 patients and partially in 1 patient treated with CSF drainage. There were 4 CSF drain complications, including headache, entrapment drain, bloody drainage, and a hematoma requiring laminectomy, but there were no permanent sequelae. Based on their overall experience, the authors also concluded that CSF drainage was warranted in high-risk patients.185

CSF drain complications have influenced practice. A meta-analysis of 34 studies of CSF drainage in 4714 patients with open and endovascular repair found a 6.5% complication rate (2.5% severe) and a CSF drain-related mortality event rate of 0.9%.189 The Mayo Clinic group reported moderate or severe complications in 17 of 187 patients (9%) receiving spinal drains before fenestrated-branched endovascular aortic repair, including spinal cord hematomas with transient paraparesis in 2 patients (1%), paraplegia in 2 (1%), intracranial hypotension in 12 (6%), and intracranial hemorrhage in 3 (2%).190 Accordingly, the Mayo group has limited their use of CSF drains with TEVAR for patients with extent I and II thoracoabdominal aortic aneurysms and selectively in extent III thoracoabdominal aortic aneurysms.190

Mousa and colleagues191 developed a formal scoring system with data from the Vascular Quality Initiative Registry based on anatomic (coverage), procedural (hemodynamic stability and time), and clinical variables (age, renal, emergency, prior aortic surgery) to stratify risk for SCI after TEVAR to help guide CSF drain placement. Nonetheless, a comprehensive review of 43 TEVAR studies concluded that selective CSF drainage was indeed warranted in high-risk patients undergoing extensive coverage along with avoidance of hypotension.183 The role of neurophysiologic monitoring with somatosensory evoked potentials and motor evoked potentials is controversial but may be helpful during thoracic aortic and thoracoabdominal aortic surgery to predict neurologic injury.192

MANAGEMENT OF TBAD WITH ARCH INVOLVEMENT

- Optimal medical therapy is reasonable in patients with uncomplicated TBAD and retrograde extension of dissection from a tear at or distal to the LSA, as long as retrograde extension is limited to the arch (zones 1 and 2). (COR IIA, LOE C-LD)

The short- and long-term natural history of retrograde extension into the arch of a TBAD is variable. IRAD data suggest that arch involvement of TBAD does not change the behavior in the short- or long-term relative to those without retrograde extension. Nauta and colleagues193 identified 67 (of 404 total type B dissections in IRAD from 1996-2014) TBAD patients with retrograde extension. They identified no differences between these patients and more distal dissections with regard to complicated presentation, treatment, and 5-year survival.193 This did not include data regarding intermediate- or long-term interventions.

Another retrospective report suggests non-A, non-B dissections with arch involvement may have a more malignant short- and long-term course compared with TBAD. Among 20 non-A, non-B dissections, Valentine and colleagues194 found 2 patients with acute retrograde extension, and compared with 79 other TBADs, the arch-involved patients required more early interventions and higher rates of stroke. Neither of these investigators suggest that these dissections should be treated differently in the absence of a complicated presentation, rather they espouse that complications should be treated when they occur. TEVAR with debranching, fenestrated/branch graft TEVAR, and open arch repair, with or without frozen elephant trunk, approaches have each been successfully used.88,195-202
SUMMARY

In summary, information and recommendations regarding the incidence, diagnosis, medical therapies, and interventional strategies to best manage patients with TBAD continue to evolve and accumulate. To outline evidence-based practice recommendations to manage patients with TBAD, the STS/AATS writing group has performed a comprehensive and methodical review and assigned treatment recommendations based upon supportive evidence. There remain gaps in evidence where expert consensus recommendations have been provided in lieu of clinical trials to guide patient management, such as:

- The ideal timing of TEVAR in the acute phase of TBAD
- Comparisons of open surgery vs TEVAR for chronic TBAD
- Appropriate size indications in chronic TBAD
- The role of acute TEVAR to prevent chronic sequelae in uncomplicated TBAD

Both associations recognize that the medical evidence will grow and treatment strategies will evolve such that these clinical practice guidelines will require continued revision as more data become available.

REFERENCES

2. Levinson DC, Edmaeudes DT, Griffith GC. Dissecting aneurysm of the aorta; its clinical, electrocardiographic and laboratory features; a report of 58 autopsied cases. Circulation. 1950;1:360-387.


