Thoracic aortic aneurysm in an adolescent with intraoperative discovery of contained rupture: a case report

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**Background:** As surgical recommendations in adults based on size criteria of ascending aortic aneurysms become more refined, criteria for childhood/adolescence remains less clear. Multiple pathologic factors may predispose younger patients to thoracic aortic aortopathy and increase the risk of rupture. An evolving field of research is how to identify thoracic aortic dilation earlier in patients, risk stratify, and to obtain objective measures beyond size for proceeding with surgical intervention in order to prevent catastrophic thoracic aortic dissection.

**Case Description:** We report an adolescent case of dilated ascending aortic aneurysm with a functionally unicuspid/bicuspid aortic valve. This patient was taken to surgery electively, given the gradual increasing size of the ascending aorta. Intraoperatively, there was an unexpected intraoperative finding of a contained aortic rupture. The patient underwent an aortic root replacement with mechanical valve composite graft and coronary artery reimplantation (modified Bentall) with ascending hemiarch replacement. The patient did well with no post-operative complications. Aortic pathology and genetic analysis were performed. The patient was discovered to have a heterozygous variant in PTPN11 which is typically associated with Noonan syndrome; however, this is not known to be associated with aortopathy.

**Conclusions:** As criteria for surgical intervention in adult thoracic ascending aortic aneurysms continues to evolve, this case illustrates challenges when determining the optimal criteria for surgical intervention in adolescent patients.

**Keywords:** Adolescent ascending aortic aneurysm; case report; congenital thoracic aortic aneurysm disease (congenital TAA disease); congenital aortic root replacement

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Introduction

Background

In 2022, the American Heart Association updated aortic disease management guidelines. The most notable change was lowering the threshold for thoracic ascending aortic aneurysm size to 5.0 from 5.5 cm (1).

Rationale and knowledge gap

As recommendations in adults become more refined, criteria for surgical intervention in childhood/adolescence in association with inheritable syndromes remains less established (2). Complicating clinical decision-making is the increased evidence of genetic predisposition to aortic dissection at diameters <5.0 cm (3).

Objective

Our adolescent case presentation, based on an unexpected intraoperative finding of contained aortic rupture with a unicuspid aortic valve (AV), highlights the challenge of determining precisely when to proceed with aortic root replacement in this patient population. We present this case in accordance with the CARE reporting checklist (available at https://acr.amegroups.com/article/view/10.21037/acr-23-163/rc).

Case presentation

A 15-year-old, 57 kg male with known valvar aortic stenosis, unicuspid AV, dilated aortic root and thoracic aortic aneurysm (TAA) presented for routine follow-up. He had an episode of chest pain 1 year prior, which self-resolved. There was no relevant family history of congenital heart disease or connective tissue disease. Transthoracic echocardiogram (TTE) demonstrated a unicuspid AV with stable moderate stenosis (peak gradient, 67 mmHg), mild regurgitation, normal left ventricle (LV) size and function, and ascending aorta 39 mm (z-score, >6). The aortic dimensions increased over time: 3 mm/2 years. On computed tomographic angiography (CTA), the mid-ascending aorta measured 41 mm (z-score, >8), severely aneurysmal, with proximal ascending aorta 32 mm (Figure 1). Given the size and progression of the aorta, he was referred for surgical repair. After discussion of valve and root replacement options with the family, the decision was made to proceed with a mechanical valve.

The patient underwent an aortic root replacement using mechanical AV composite graft 21 mm with coronary artery reimplantation (modified Bentall) and ascending hemiarch replacement (24 mm graft) under deep hypothermic circulatory arrest (DHCA) with retrograde cerebral perfusion (Figure 2, Video 1). Routine monitoring was performed with an arterial/central line, foley, and electroencephalogram/somatosensory evoked potentials. Median sternotomy and central cannulation were performed. Cardioplegia was administered into the aortic root. Upon transection of the aorta, an old hematoma was visible along the medial, posterior aspect near the main pulmonary artery. There was a small intimal tear, suggesting a contained, remote perforation. The AV was functionally unicuspid/bicuspid and not repairable (Figure 3). Total bypass time was 154 minutes, cross-clamp time 117 minutes, with DHCA 13 minutes. The patient tolerated the procedure well. Post-bypass echo demonstrated good valve function with no perivalvar leak and excellent cardiac function.

Post-operatively, there were no major complications. He was started on heparin for bridging therapy to coumadin [international normalized ratio goal 2–3]. He was discharged on post-operative day 11. On 6-month follow-up, his mechanical valve is well functioning with mild regurgitation laterally, most likely due to washing jets. A representative slide of the aortic tissue demonstrates normal histology in terms of wall thickness and organization (Figure 4). There is no evidence of myxoid degeneration or elastic fiber degradation. Genetic analysis resulted in a heterozygous missense variant of uncertain significance in PTPN11. This

Highlight box

Key findings

- Adolescent thoracic aortic aneurysms may be prone to rupture prior to indications for surgery.

What is known and what is new?

- Criteria for adult aortic aneurysm surgical indications continues to evolve with decreasing size threshold. There are several known obstacles to establishing criteria in children and adolescents.
- This case provides a clinical example of why it is important to establish criteria for adolescent aortic aneurysm operative indications.

What is the implication, and what should change now?

- Definitive guidelines should be established beyond size criteria for adolescents with thoracic ascending aortic aneurysms in order to decrease the risk of rupture.
mutation is typically associated with Noonan syndrome and not known to be associated with aortopathy (4).

All procedures performed in this study were in accordance with the ethical standards of the institutional and/or national research committee(s) and with the Helsinki Declaration (as revised in 2013). Written informed consent was obtained from the patient’s parent for the publication of this case report and accompanying images. A copy of the written consent is available for review by the editorial office of this journal.

**Discussion**

**Key findings**

This case highlights multiple important factors when considering surgical repair for TAA in adolescents: atypical genetic syndromes, association with bicuspid AV (BAV), size <4.0 cm, and intraoperative contained rupture.

**Figure 1** Computed-tomography angiography of the chest. (A) Sagittal view demonstrating aneurysmal ascending aorta (black asterisks, **). (B) Volume-rendered 3-dimensional reconstruction demonstrating dilation of the ascending aorta (black asterisks, **).

**Figure 2** Dilated root, ascending aortic aneurysm and bicuspid aortic valve in a 15-year-old male.

**Video 1** Intraoperative aortic root replacement and hemiarch in an adolescent patient.
Strengths and limitations

As an isolated case report, there are limitations to generalizability of this specific case. And, given the unknown timeframe for when the patient experienced the contained aortic rupture, it is still unclear at exactly what size the aorta was at the time of rupture. We also did not have advanced imaging such as four-dimensional (4-D) flow magnetic resonance imaging (MRI). However, this intraoperative finding gives a unique clinical example of establishing improved criteria beyond size for evaluating adolescents with dilated ascending aortas.

Comparison with similar research

In evaluating BAV adult patients, a higher proportion require operative intervention at a younger age, presented with a smaller aorta (4.6 cm) and more rapid rate of growth (1.9 mm/year), compared to TAA without BAV (5). BAV is also associated with childhood TAA; it is unclear whether this predisposes to dissection at smaller diameters prior to adulthood (2). BAV genetic mutations could become a surrogate marker for early, progressive TAA. The intricate relationship between genetic mutations, aortic dissection, and aorta sizes continues to be explored (3).

Figure 3 Congenital aortic valve stenosis. (A) Intraoperative view from aortotomy in vivo. (B) Excised aortic valve demonstrating functionally unicuspid morphology.

Figure 4 Aortic tissue histology, using Movat's stain, demonstrating normal wall thickness and organization. (A) Specimen slide. (B) Specimen slide magnified 10x.
Specifically, in regards to younger male adult BAV patients, AV size and function may be a poor surrogate for intrinsic aortic properties (6). The loss of elastic fibers with aortic wall stiffness can be diagnosed irrespective of aortic wall dilation. This was observed in patients with no or mild AV impairment. Usage of serial echographic measurements may allow earlier identification and guide medical/surgical intervention of potentially modifiable risk factors. These observations continue to establish that only aortic size in younger patients may be an inadequate parameter to address what changes are occurring in the aortic wall in BAV patients that can predispose to rupture.

Although connective tissue disease such as Marfan syndrome and Loeys-Dietz syndrome are known to have an association with TAA leading patients to early dissection/rupture, this represents the minority of patients [i.e., Marfan disease only constitutes 5% of TAA patients (7)]. There is still much to learn about the true inheritability of aortopathy beyond connective tissue disorders. At the Yale Aortic Institute, Elefteriades (7), with mapping almost 500 family trees for TAA patients, discovered that at least 21% have at least one family member with a known TAA. These positive findings in familial relatives also demonstrated a higher growth rate and earlier age of presentation. It is likely that the pattern of inheritance is predominantly autosomal dominant, but more research will need to be done in order to elucidate the precise rate and method of inheritance.

Implications and actions needed

Our patient’s genetic results demonstrate a need for further testing in order to better understand the natural history and clinical implications for congenital aortic diseases. The complex relationship between genetic heritability, aortic pathology, aortic rupture/dissection, and aorta size continues to be studied with further research necessary (3,6).

As 4-D flow MRI imaging becomes more advanced, this allows us to increase surveillance and understanding of the aorta at local/regional levels of wall stress (8). Especially for pediatric patients, this non-invasive imaging that provides increased dimensionality for further understanding of the aorta is exciting. Larger patient volumes and studying different regional populations with ascending aortic disease will allow generalizability of these results.

Currently, indications for surgery in ascending aortic disease are limited by absolute size and rate of growth in adult patients. Some alternative ideas to categorize age-adjusted risk include z-scores, rate of growth, or a calculation of aortic root to descending aorta ratio. Ultimately, creating a mathematical model to predict a patient’s risk of rupture would be the ideal tool. Our intraoperative findings of a contained rupture suggest that offering surgery earlier in this patient would have been appropriate. In the future, a similar 15-year-old patient with aortic morphology should have improved guidelines with data to indicate an earlier indication for surgery in order to prevent rupture.

Conclusions

In summary, definitive guidance for optimal timing of surgery for TAA and BAV in childhood/adolescence remains deficient. Obstacles to identifying appropriate criteria and eligible patients include lower volume, lack of availability of genetic sequencing, and variability in clinical presentation. As guidelines develop to include more aortic parameters (8,9), biomarkers, tissue biorepositories, genetic mutations, and environmental factors for dissection (7) the treatment of TAA in childhood/adolescence will continue to improve.

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Footnote

Reporting Checklist: The authors have completed the CARE

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Conflicts of Interest: All authors have completed the ICMJE uniform disclosure form (available at https://acr.amegroups.com/article/view/10.21037/acr-23-163/coif). 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. All procedures performed in this study were in accordance with the ethical standards of the institutional and/or national research committee(s) and with the Helsinki Declaration (as revised in 2013). Written informed consent was obtained from the patient's parent (patient is a minor) for the publication of this case report and accompanying images. A copy of the written consent is available for review by the editorial office of this journal.

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