Aortopulmonary Window: Classification, Associated Cardiac Anomalies, Treatment Options, and Clinical Outcome

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Objective: Aortopulmonary window (APW) is an uncommon congenital cardiac abnormality marked by a septation defect between the ascending aorta and pulmonary artery. This study aimed to define the clinical characteristics, diagnostic features, treatment strategies, and follow-up outcomes of pediatric patients diagnosed with APW.

Material and Methods: We retrospectively reviewed children diagnosed with APW from 2010 to 2023. Morphological APW typing of our patients was based on the classification that is settled by the Society of Thoracic Surgeons (STS) Congenital Heart Surgery Database Committee. The patients’ demographic data, symptoms at admission, transthoracic echocardiography, cardiac computed tomography, management modalities, and follow-up data were evaluated.

Results: Twenty-five children were diagnosed with APW over the study period. Thirteen patients were male (52%), and the median age at presentation of the patients was three months (8 days-7.5 years). Two patients were diagnosed with coronary fistula by echocardiography at the first admission and were diagnosed with APW after catheterization. APW was detected in one patient while being operated on for large ventricular septal defect. According to the STS classification, 32% (n=8) of the patients were type III, 32% (n=8) were type I, 16% (n=4) were intermediate type, 12% (n=3) were type II, and 4% (n=1) were APW with aortic interruption. Associated cardiovascular malformations were in 76% (n=19) of the patients. Fifteen patients (60%) underwent surgery. Transcatheter closure of APW was performed in four patients (16%).

Conclusion: Detection of the APW requires careful and systematic investigation. Transcatheter closure can be performed in selected cases where the defect is suitable. Although rare, this defect, which can cause severe left-right shunting, should be kept in mind as a cause of pulmonary hypertension and unexplained cardiac dilation and should be investigated in patients whose cause cannot be determined.

Keywords: Aortopulmonary window, echocardiography, pulmonary hypertension, surgical treatment, transcatheter treatment

ABSTRACT

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Introduction

Aortopulmonary window (APW) is an infrequent congenital cardiac defect characterized by incomplete partitioning of the aortic and pulmonary artery walls, leading to communication between these two structures. The presence of two separate semilunar valves helps distinguish APW from truncus arteriosus (1). This cardiac anomaly is a rare occurrence, accounting for approximately 0.2-0.6% of all congenital heart defects (2). Previous studies have reported a higher incidence of this cardiac anomaly among males (1,2,3). However, findings have been observed, with some reports indicating a female predominance (4,5), while others suggest a comparable occurrence rate between males and females (6). APW is frequently accompanied by other congenital heart defects. These associated anomalies may include ventricular septal defect (VSD), tetralogy of Fallot, transposition of the great arteries, double-outlet right ventricle, aortic interruption, aortic atresia, and in rare instances, coronary abnormalities (4,6,7).

In APW, intercommunication between the aorta and pulmonary artery causes a left-to-right (systemic-to-pulmonary) shunt, with the degree of shunting dictating the clinical course of the disease.

Shunting from the aorta to the pulmonary artery considerably increases during the neonatal period when pulmonary vascular resistance decreases. This causes congestive heart failure, which manifests as tachypnea, tachycardia, irritability, exhaustion, sweating, poor feeding or failure to thrive. If left untreated, pulmonary vascular obstructive disease may ensue at an early age, which is irreversible. Hence, early closure of large APWs is imperative to avert heart failure or pulmonary vascular disease. While surgery remains the standard treatment of choice (6,8), transcatheter APW closure can be a suitable alternative for patients with adequate rims (9,10).

The main aim of this study was to delineate the clinical attributes, diagnostic features, therapeutic approaches, and follow-up of pediatric patients diagnosed with APW.

Material and Methods

This retrospective analysis examined 25 patients who received an APW diagnosis at our clinic between 2010 and 2023, ranging from 8 days to 7.5 years. Before conducting the investigation, all ethical standards were met in accordance with the Declaration of Helsinki, with approval obtained from the University of Health Sciences Turkey, Istanbul Mehmet Akif Ersoy Thoracic and Cardiovascular Surgery Training and Research Hospital Ethics Committee (March 2022/2022.03.19). Patient data, including demographic information, admission symptoms, transthoracic echocardiography, cardiac computed tomography (CT), management approaches, and follow-up information, were retrospectively evaluated.

Our patient cohort’s morphological APW typing was determined based on the Society of Thoracic Surgeons (STS) Congenital Heart Surgery Database Committee classification (11) modified version of Mori et al.’s (3) classification. This classification system includes five subtypes:

1. APW with interrupted aortic arch,
2. Type I (proximal) APW, located above the sinus of Valsalva with a small inferior rim,
3. Type II (distal) APW, located at the highest part of the ascending aorta with a well-formed inferior rim and a small superior rim,
4. Type III (complete defect of the aortopulmonary septum) APW involves most of the ascending aorta with minimal superior and inferior rims,
5. Intermediate-type APW exhibits adequate superior and inferior rims and is more amenable to transcatheter closure.

According to the occurrence of concomitant cardiovascular anomalies, the study population was divided into two groups: the isolated APW group (n=6) and the complex APW group (n=19). Management strategies for the patients included surgical or transcatheter closure of the defect or medical treatment for pulmonary hypertension (PH) in patients with negative vasoreactivity. The regular outpatient follow-up was conducted at 3-6 months intervals. Follow-up was defined as the period from presentation to the last admission or death. Of the total patient cohort, three patients were lost to follow-up, one died after the operation, and the remaining 21 completed the follow-up period.

Statistical Analysis

The statistical analyzes were carried out using the SPSS 25 program (SPSS Inc., Chicago, IL, USA). For categorical data, frequencies and percentages are shown. The median value and interquartile range for variables with non-normal distributions were used, whereas the mean and standard deviation were used for variables with normal distributions.

Results

Demographic and Clinical Characteristics

From 2010 to 2023, 25 patients were diagnosed with APW in our clinic. The median age at presentation was three
months (8 days-7.5 years) and the median weight was 4.25 kg (2.1-21 kg). Of the patients, 13 were male (52%) and 12 were female (48%), resulting in a male-to-female ratio of 1.08 to 1. Congestive heart failure was the most common presenting symptom observed in 12 patients (50%), while 11 patients (44%) had heart murmurs. Two patients with significant PH and a negative vasoreactivity test were asymptomatic at admission. Diagnostic evaluations included echocardiography, cardiac catheterization (54.2%), and cardiac CT (48%) (Figure 1). Table 1 provides further details of the patients with APW.

We could visualize the AP window by 2D echocardiography in 22 (88%) patients. Two patients were initially diagnosed with a coronary fistula but were later diagnosed with APW after catheterization. In one patient, APW was detected during surgery for large VSD. In 21 patients with APW, there was significant left-to-right shunting that led to left atrial and ventricular dilatation. In most cases (83.3%), except for four, severe PH was observed, with pulmonary pressure almost equivalent to systemic pressures. Pulsed-wave and color flow Doppler demonstrated negative diastolic flow in the descending aorta due to diastolic left-to-right shunting through the APW.

One patient (4%) had APW with aortic interruption, while type III APW was noted in 32% (n=8) of the patients, type I in 32% (n=8), intermediate type in 16% (n=4), and type II in 12% (n=3), as per the STS classification. In addition, a patient had a 15 mm wide APW between the descending aorta and the main pulmonary artery at the level of the left subclavian artery.

In the cohort under investigation, a substantial proportion (76%, n=19) of the patients presented with associated cardiovascular malformations. Within the subset of patients with complex APW, the most frequent co-occurring cardiovascular malformations included topsy-turvy heart in 4 cases (16%), atrial septal defect in 4 cases (16%), and VSD in 4 cases (16%). An interrupted aortic arch type A and patent ductus arteriosus was in one patient (4%) with complex APW. Notably, one male patient (4%) with complex APW type I presented with abnormal coronary anatomy, specifically an anomalous origin of right coronary artery originating from the pulmonary artery (ARCAPA). Three patients (12%) displayed a right aortic arch; of these, two had an aberrant left subclavian artery (ALSA). Finally, a 7.5-year-old male patient within the complex APW cohort had tetralogy of Fallot as an associated cardiac defect.

Patients who were diagnosed with a topsy-turvy heart had a recognizable rotational abnormality affecting their heart and major arteries. Atrioventricular and ventriculoarterial connections on the left-sided heart showed concordance, and atrial configurations showed the typical superior-inferior

![Figure 1. Echocardiography and computed tomography images of the patients with aortopulmonary window.](image-url)

(a) Transthoracic echocardiography from apical four-chamber view shows enlarged left atrium (LA), and left ventricle (LV). (b) Transthoracic echocardiography from a parasternal short-axis view shows a wide defect between the ascending aorta (Ao) and the main pulmonary artery (PA) (white arrow). Computerized tomography shows a large aortopulmonary window (white arrow) in axial (c) and coronal (d) views.

RA: Right atrium, RV: Right ventricle
Table 1. Details of patients with aortopulmonary window

<table>
<thead>
<tr>
<th>Patient</th>
<th>Age (month)</th>
<th>Weight (kg)</th>
<th>Symptom</th>
<th>Gender</th>
<th>Aortopulmonary window type</th>
<th>Concomitant cardiovascular anomalies</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>3.5</td>
<td>4.5</td>
<td>Heart failure symptoms</td>
<td>M</td>
<td>III</td>
<td>Topsy-turvy heart</td>
<td>Surgery</td>
</tr>
<tr>
<td>2</td>
<td>4</td>
<td>3.8</td>
<td>Murmur</td>
<td>M</td>
<td>Intermediate type</td>
<td>Bicuspid aorta, right aortic arch, aberrant left subclavian artery</td>
<td>Transcatheter occlusion (ADO-I)</td>
</tr>
<tr>
<td>3</td>
<td>12</td>
<td>11</td>
<td>Murmur</td>
<td>F</td>
<td>I</td>
<td>Bicuspid aorta, aortic stenosis</td>
<td>Negative vasoreactivity</td>
</tr>
<tr>
<td>4</td>
<td>4</td>
<td>5</td>
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<td>M</td>
<td>I</td>
<td>Left pulmonary artery stenosis</td>
<td>Surgery</td>
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<tr>
<td>5</td>
<td>3.5</td>
<td>21</td>
<td>None</td>
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<td>Between the descending aorta and main pulmonary artery</td>
<td>Topsy-turvy heart</td>
<td>Negative vasoreactivity</td>
</tr>
<tr>
<td>6</td>
<td>15</td>
<td>6.5</td>
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<td>F</td>
<td>I</td>
<td>-</td>
<td>The decision of surgery was made. The patient lost to follow-up</td>
</tr>
<tr>
<td>7</td>
<td>6</td>
<td>5.3</td>
<td>Murmur</td>
<td>F</td>
<td>Intermediate type</td>
<td>Ventricular septal defect</td>
<td>Transcatheter occlusion (ADO-I)</td>
</tr>
<tr>
<td>8</td>
<td>72</td>
<td>18</td>
<td>Murmur</td>
<td>M</td>
<td>I</td>
<td>ARCAPA</td>
<td>Surgery</td>
</tr>
<tr>
<td>9</td>
<td>4</td>
<td>6.7</td>
<td>Heart failure symptoms</td>
<td>M</td>
<td>I</td>
<td>Ventricular septal defect</td>
<td>The decision of surgery was made. The patient lost to follow-up</td>
</tr>
<tr>
<td>10</td>
<td>5</td>
<td>5.5</td>
<td>Murmur</td>
<td>F</td>
<td>I</td>
<td>Right-sided aortic arch, aberrant left subclavian artery</td>
<td>Surgery</td>
</tr>
<tr>
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<td>10 days</td>
<td>2.1</td>
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<td>F</td>
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<td>Atrial septal defect</td>
<td>Transcatheter occlusion (ADO-II)</td>
</tr>
<tr>
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<td>Murmur</td>
<td>M</td>
<td>III</td>
<td>Atrial septal defect, a right-sided aortic arch, left persistent superior vena cava</td>
<td>Surgery</td>
</tr>
<tr>
<td>13</td>
<td>5</td>
<td>4</td>
<td>Heart failure symptoms</td>
<td>M</td>
<td>I</td>
<td>-</td>
<td>Surgery</td>
</tr>
<tr>
<td>14</td>
<td>2.5</td>
<td>3.7</td>
<td>Heart failure symptoms</td>
<td>M</td>
<td>II</td>
<td>-</td>
<td>Surgery</td>
</tr>
<tr>
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<td>19 days</td>
<td>3</td>
<td>Heart failure symptoms</td>
<td>M</td>
<td>III</td>
<td>Topsy-turvy heart</td>
<td>Surgery</td>
</tr>
<tr>
<td>16</td>
<td>2.5</td>
<td>2.7</td>
<td>Murmur</td>
<td>M</td>
<td>I</td>
<td>Ventricular septal defect</td>
<td>Surgery</td>
</tr>
<tr>
<td>17</td>
<td>2</td>
<td>2.8</td>
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<td>F</td>
<td>III</td>
<td>Atrial septal defect</td>
<td>Surgery</td>
</tr>
<tr>
<td>18</td>
<td>2</td>
<td>2.7</td>
<td>Heart failure symptoms</td>
<td>F</td>
<td>III</td>
<td>-</td>
<td>Surgery</td>
</tr>
<tr>
<td>19</td>
<td>20 days</td>
<td>3</td>
<td>Heart failure symptoms</td>
<td>F</td>
<td>III</td>
<td>Topsy-turvy heart</td>
<td>Surgery</td>
</tr>
<tr>
<td>20</td>
<td>8</td>
<td>8.3</td>
<td>Murmur</td>
<td>F</td>
<td>Intermediate type</td>
<td>-</td>
<td>Transcatheter occlusion (ADO-I)</td>
</tr>
<tr>
<td>21</td>
<td>90</td>
<td>20</td>
<td>None</td>
<td>M</td>
<td>II</td>
<td>Tetralogy of Fallot, absent left pulmonary artery</td>
<td>Negative vasoreactivity</td>
</tr>
</tbody>
</table>
relationship of cardiac chambers. In addition, a significant connection was observed between the aorta and pulmonary artery (APW), which was associated with systemic pulmonary artery hypertension. Consistent with previous literature, all topsy-turvy heart cases were born to consanguineous parents, with two cases presenting as siblings. Among the patients analyzed, 23 out of 25 cases (92%) displayed no extracardial anomalies, while 8% of patients (two cases) demonstrated associated non-cardiac abnormalities, including Cornelia de Lange syndrome and tracheoesophageal fistula.

**Management and Follow-up**

Fifteen patients (60%) underwent surgery (patch repair of APW) at a median of 2.2 months (22 day-6 years) when they had a median weight of 3.6 kg (range, 2.7-18 kg). Among them, one male patient with abnormal coronary anatomy underwent reimplantation of ARCAPA during APW repair, and one patient with an interrupted aortic arch underwent patch augmentation of the aortic arch. A patient with esophageal compression resulting from an ALSA was successfully treated with ALSA transaction. Following surgical intervention, a patient diagnosed with a topsy-turvy heart manifested acute respiratory distress and left lung hyperinflation, leading to a rapid clinical deterioration. On the tenth postoperative day, venoarterial extracorporeal membrane oxygenation (ECMO) was initiated due to persistent respiratory failure. The patient of multiorgan failure and died on the 16th postoperative day, despite receiving six-day ECMO treatment (12). Transcatheter closure of APW was performed in four patients (16%) with aortopulmonary septal defects with adequate septal rims located at a safe distance from the aortic and pulmonary valves, coronary arteries, and pulmonary artery bifurcation at a median of 7.5 months (range: 4 months to 1.4 years) (Figure 2). No patient died during the intervention. Following a mean follow-up period of 33.6±28.3 months (4 months to 79 months), all patients remained asymptomatic without requiring medication. In three patients with large defects, cardiac catheterization indicated high pulmonary vascular resistance, and the pulmonary vasoreactivity test with nitric oxide and 100% oxygen inhalation was non-reactive. Consequently, medical PH treatment was initiated. However, three patients required surgical intervention but were not followed up.

**Discussion**

This study provides an analysis of the clinical and diagnostic features, treatment approaches, and consequences of 25 pediatric patients with APW from a tertiary cardiac center. The APW is characterized by a deficient septum between the pulmonary artery and ascending aorta. Previous research has linked this anomaly with a diverse range of other cardiac malformations, including interrupted aortic arch, VSDs, topsy-turvy heart, tetralogy of Fallot, transposition of the great arteries, and coronary anomalies (4,7,13,14). More than half of our patients demonstrated associated cardiovascular abnormalities in our study. Among patients with complex APW, topsy-turvy heart, atrial septal defect, and VSD were the most frequently observed associated conditions (50%). In the case reports we reviewed, rare cardiovascular abnormalities were identified in association with APW, including a right aortic arch with an aberrant origin of the left subclavian artery (15), an isolated origin of the left subclavian artery from the left pulmonary artery (16), total abnormal pulmonary venous return (17), a right pulmonary artery originating from the ascending aorta (8), and crisscross pulmonary arteries (18). We have not encountered such rare abnormalities in our patients. Aortic stenosis, bicuspid aortic valve, bilateral superior vena cava, partially aberrant pulmonary venous return, and ARCAPA are a few unique coexisting cardiovascular problems that we did notice.

### Table 1. Continued

<table>
<thead>
<tr>
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<th>Weight (kg)</th>
<th>Symptom</th>
<th>Gender</th>
<th>Aortopulmonary window type</th>
<th>Concomitant cardiovascular anomalies</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>22</td>
<td>20 days</td>
<td>3.5</td>
<td>Heart failure symptoms</td>
<td>M</td>
<td>III</td>
<td>Partial anomalous pulmonary venous return</td>
<td>The decision of surgery was made. The patient lost to follow-up</td>
</tr>
<tr>
<td>23</td>
<td>1</td>
<td>5</td>
<td>Heart failure symptoms</td>
<td>M</td>
<td>III</td>
<td>Ventricular septal defect</td>
<td>Surgery</td>
</tr>
<tr>
<td>24</td>
<td>1</td>
<td>3.4</td>
<td>Heart failure symptoms</td>
<td>F</td>
<td>APW with aortic interruption type A</td>
<td>Aortic interruption</td>
<td>Surgery</td>
</tr>
<tr>
<td>25</td>
<td>14 days</td>
<td>3</td>
<td>Murmur</td>
<td>F</td>
<td>I</td>
<td></td>
<td>Surgery</td>
</tr>
</tbody>
</table>

ADO: Amplatzer duct occluder, APW: Aortopulmonary window, ARCAPA: Anomalous origin of right coronary artery originating from the pulmonary artery, F: Female, M: Male
The clinical diagnosis of APW can be challenging in some cases and may require repeat echocardiography, diagnostic cardiac catheterization, or intraoperative assessment. Our study found that some patients were initially misdiagnosed but were later accurately diagnosed through further evaluation. One plausible explanation for the initial misdiagnosis of APW in a patient diagnosed during VSD surgery could be equalizing aortic and pulmonary pressure due to PH. Similarly, in two patients with a coronary fistula diagnosed during echocardiography, the small defect was mistaken for a fistula, and APW was only diagnosed during cardiac catheterization. In Kiran et al.’s (2) study, four patients were initially misdiagnosed with APW, and the factors contributing to this were analyzed. When there is unexplained heart failure, left heart dilatation due to a significant left-to-right shunt, and PH, an APW should be suspected. However, when the patient’s clinical status and echocardiographic findings appear compatible, complex variants may be challenging to diagnose and easily overlooked. In addition to transthoracic echocardiography, this pathology can be demonstrated with the help of catheter or CT angiography and magnetic resonance imaging. Failure to diagnose APW patients can lead to irreversible obstructive changes in the pulmonary vascular bed, making the patient inoperable (19). In our cases, the late presentation of three patients with large defects (1 year, 3.5 years, and 7.5 years) highlights the importance of early detection and prompt intervention to avoid irreversible pulmonary obstructive changes, as all three had significant PH and negative vasoreactivity tests.

Important clinical factors in managing patients with APW include the size of the defect, the degree of left-to-right shunting into the pulmonary trunk, the presence of concurrent cardiovascular malformations, and the emergence of PH. Early closure is imperative given the potential for rapid onset of congestive heart failure owing to high pulmonary blood flow. The conventional approach to treating APW is surgical intervention, and several studies (4,5,6,8,13,20) have examined surgical outcomes in this context. However, transcatheter closure may serve as a viable alternative in cases with no associated cardiovascular lesions and adequate rims. In fact, several studies (9,10,21) have documented successful experiences with transcatheter occlusion of APW. It is of utmost importance to carefully assess the location and size of the defect as well as the amount of superior and inferior rims present to minimize the risk of device-related complications such as embolization, coronary artery blockage, and damage to great vessels and valves. Additionally, the type of defect plays a critical role in determining the feasibility of transcatheter closure, with intermediate-type defects being more amenable to this approach.

**Study Limitations**

We acknowledge that our study has several limitations. Retrospective reviews of patients with APW were performed at a single center, which limits the generalizability of our findings. Additionally, our study was limited by the middle follow-up period.

**Conclusion**

In patients with AP windows, the current preferred method of diagnosis is echocardiography. Accurate diagnosis necessitates a meticulous and systematic evaluation. Small defects associated cardiac anomalies, or PH may present a diagnostic challenge, and the misdiagnosis is possible. Transcatheter closure can be considered in appropriate cases with sufficient septal rims. Although rare, this defect can
cause severe left-right shunt, leading to PH and unexplained cardiac dilation. Therefore, it should be considered a possible cause and investigated in patients whose underlying causes cannot be determined.

**Ethics**

**Ethics Committee Approval:** All ethical standards were met in accordance with the Declaration of Helsinki, with approval obtained from the University of Health Sciences Turkey, Istanbul Mehmet Akif Ersoy Thoracic and Cardiovascular Surgery Training and Research Hospital Ethics Committee (March 2022/2022.03.19).

**REFERENCES**


