Idiopathic Dilatation of Pulmonary Artery with Effort Angina

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Abstract: Idiopathic pulmonary artery dilatation is a rare vascular abnormality with an incidence of 0.007% in autopsy. Diagnosis is established by excluding diseases that induce pulmonary arterial enlargement. Dilatation of the pulmonary artery is complicated with mild pulmonic valve stenosis. Here, we present a rare case of idiopathic dilatation of the pulmonary artery with mild valvular pulmonic stenosis which presented with effort angina and was diagnosed using echocardiography and cardiac catheterization.

(key words—idiopathic pulmonary artery dilatation with pulmonary stenosis, idiopathic pulmonary artery dilatation with effort angina)

I. Case summary

36 years old female patient born of non consanguineous marriage from low socioeconomic status having no significant past history of medical illness presented with exertional chest pain of 15 days duration, located in midline, heaviness type, moderate in intensity, non radiating, lasting more than 5-10 minutes, subsiding with taking rest. There was also history of exertional breathlessness (NYHA II) of 15 days duration. No history of cough, fever, palpitation, giddiness, breathlessness in lying down position at night.

Pulse rate 84/min, respiratory rate 15/min, BP 126/80 mmHg, saturation 98% in room air. There was no pallor, icterus, cyanosis, pedal edema, JVP was not elevated. Apical impulse was located in 5th intercostal space medial to mid clavicular line, normal character. Precordial pulsations in left second intercostal space, left 3rd-4th intercostals spaces. Left second space was dull to percuss. Ejection click with ejection systolic murmur was heard in the same area. Ejection click was louder during expiration. ESM was of grade 2/6, high pitched, crescendo decrescendo in nature. Respiratory system, gastro intestinal system, central nervous system examination was normal.

Patient’s investigations were; Hemoglobin 13gm/dl, total count 10200/mcL, neutrophils 62%, lymphocytes 30%, eosinophil 4%, monocytes 5%, basophils 1%. ESR 12mm. Natriuretic peptide (BNP) and antineutrophil cytoplasmic autoantibody (ANCA) levels were within normal limit, and there was no evidence of infectious diseases such as tuberculosis or syphilis.

X-Ray chest revealed mild cardiomegaly of right ventricular type, mild right atrial dilatation, massive enlargement of main pulmonary artery with normal pulmonary vascular markings. Echocardiographic findings were; enlarged enlarged main pulmonary artery (3.8 cms), right pulmonary artery (1.9 cms), left pulmonary artery (1.8 cms) in parasternal short axis (ULN is 2.1 cm for MPA and 1.4 cm for LPA and RPA). Right ventricle dimension was 2.1 cm. Right ventricle free wall thickness was 5mm. There was no intracardiac and extracardiac shunt. Left ventricular function was normal. The right descending pulmonary artery measured 17 mm on the chest radiograph; the upper limit of normal is 16 mm.

Cardiac catheterization was done through right femoral vein/right femoral artery approach. RV angiogram showed trabeculated right ventricle with normal pulmonary valve and massively dilated main pulmonary artery. Hemodynamic data were as follow:

<table>
<thead>
<tr>
<th>Site</th>
<th>pressure</th>
<th>saturation</th>
</tr>
</thead>
<tbody>
<tr>
<td>SVC</td>
<td>Mean- 4mmHg</td>
<td>74%</td>
</tr>
<tr>
<td>RA</td>
<td>Mean- 4mmHg</td>
<td>78%</td>
</tr>
<tr>
<td>RV</td>
<td>48/7mmHg</td>
<td>82%</td>
</tr>
<tr>
<td>PA</td>
<td>30/8(18) mmHg</td>
<td>79%</td>
</tr>
<tr>
<td>LV</td>
<td>120/12 mmHg</td>
<td>98%</td>
</tr>
<tr>
<td>AO</td>
<td>120/84(108)mmHg</td>
<td>98%</td>
</tr>
</tbody>
</table>

There was no significant step up in saturation suggestive of intra cardiac as well as extra cardiac left to right shunts. Coronary angiogram revealed normal coronaries. Hence, the diagnosis of Idiopathic pulmonary artery dilatation was made.
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II. Discussion.

Idiopathic dilatation of the pulmonary artery is a rare disease first described by Wessler and Jaches [1]. The incidence of this disease is 6 cases per 1000 cases of congenital cardiac disorders [2]. Greene et al. in 1949 [3] proposed the following diagnostic criteria for this condition: (1) simple dilatation of the pulmonary trunk; (2) absence of intraorextra-cardiac shunts; (3) absence of chronic cardiopulmonary disease; (4) absence of arterial disease such as syphilis, atherosclerosis, or arteritis. Deshmukh et al. in 1960 [4] added the fifth criterion, i.e. normal pressure in the right ventricle and pulmonary artery. Only 5 cases of idiopathic pulmonary aneurysms with mild pulmonary valve stenosis have been reported to date (5). The pressure gradients of all five patients were lower than 20 mmHg, as in our case. Van Buchem et al. [6] reported that the normal diameter of the pulmonary artery ranged from 22 to 33 mm. Post-stenotic dilatation was observed when the pressure gradient was more than 25 mmHg. In contrast, patients with idiopathic pulmonary aneurysms showed remarkably dilated pulmonary arteries in the absence of pressure gradients (≤ 5 mmHg). Although a mild pressure gradient might increase pulmonary artery enlargement, it was not considered as one of the main reasons for the marked pulmonary artery dilatation. In 1987, Futagami et al. [7] reported 60 Japanese cases of idiopathic dilatation of the pulmonary artery. Pressure data of 39 of these patients were analyzed. A pressure gradient of ≤ 5 mmHg, considered as no or trivial pressure gradient was observed in 64% of the 39 patients, but 36% of patients showed mild pressure gradients (6—22 mmHg) between the right ventricle and the pulmonary artery. These data suggest that a mild pressure gradient might often be seen in patients with idiopathic pulmonary aneurysms. CMR may be used in the diagnosis of pulmonary artery aneurysms in case of doubt in echocardiography as a non-invasive modality with advantages of measuring the precise diameter of the aneurysm, use of contrast agents, evaluation of the right ventricular outflow tract, no radiation exposure, and quantification of pressure gradient of pulmonary artery stenosis. (5). It is the preferred option for combined use with echocardiography in the diagnosis and follow-up of patients with IDPA.

Characteristic pattern is found on analysis of the phonocardiogram; it consists of a normal first heart sound, a systolic click, a faint or absent pulmonic systolic murmur, wide splitting of the second heart sound which is usually fixed, and occasionally a diastolic murmur at the pulmonary area. This distinctive pattern may prove to be of value in the diagnosis of idiopathic dilatation of the pulmonary artery (8). Pulmonary insufficiency observed in 80% of cases, which was not seen in our case. Exertional chest pain observed in our case could be due to massive dilatation of pulmonary artery with increased wall stress which is pronounced during effort.

No data are available for the long-term prognosis of patients with idiopathic dilatation of the pulmonary artery with mild pulmonary artery stenosis. More so with symptomatic cases, although there are case reports of asymptomatic pulmonary dilatation detected on routine examination in sixth decade (10). Idiopathic pulmonary artery dilatation is clearly a benign and non-progressive condition in contrast to our case. Any rapid progression of symptoms with apparent idiopathic dilatation of pulmonary artery may have an arteritis or connective tissue disorder. Eventhough asymptomatic cases need to be followed up for the development of connective tissue diseases.

III. Conclusions:

Idiopathic pulmonary artery dilatation is a rare disease after exclusion of conditions causing pulmonary artery dilatation. Any patient with apparent diagnosis of idiopathic pulmonary hypertension needed to be followed up for long time to know benign course. Any rapid course of symptoms should be evaluated for possible infectious and inflammatory diseases, although occasionally the manifestation of idiopathic pulmonary artery dilatation.

References