Androgenic-Anabolic Steroid (Boldenone) Abuse as a Cause of Dilated Cardiomyopathy

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Case Presentation

A 34-year old, previously healthy male, presented to the ER with worsening dyspnea over a period of three weeks and the appearance of blood tinged sputum. Soon after presentation, the patient was hypoxic and required mechanical ventilation. Chest X-ray demonstrated bilateral infiltrates and he was admitted to the ICU with the tentative diagnosis of acute respiratory distress syndrome. Investigation by PICCO, Suggested the presence of heart failure were and the patient was transferred to the ICCU. On admission, the patient maintained normal blood pressure but still needed mechanical support with high FiO₂ values to maintain adequate oxygenation. Physical examination was remarkable for his muscular appearance. The heart sounds were regular. The lungs were clear to auscultation, although chest X-ray revealed signs of pulmonary edema. Blood count was remarkable for hemoglobin of 21 gr/dl with hematocrit of 65%. His blood chemistry showed creatinine level on 1.74 mg/dL, with no electrolytes imbalance. LFT were normal, total protein and albumin levels were normal, elevated CK with normal TnT. CRP 10 mg/L (normal level <10 mg/L). Blood cultures and panel of common respiratory infections were all negative. ECG showed normal sinus rhythm, with p pulmonale, signs of hypertrophy and inverted T waves in chest leads. Swan-ganz Cather measurements revealed cardiac output (CO) 3.9, capillary wedge pressure 34 mmHg, SVR 2070 and CI 2.2 consistent with cardiogenic shock. Transthoracic echocardiography demonstrated severe global dysfunction (left ventricular ejection fraction = 20%) with severely dilated left ventricle 5.1 cm (end-systolic diameter), asymmetric hypertrophy (Septum 14 mm, Posterior wall 12 mm). Minimal mitral and tricuspid regurgitation were also noted (Table 1 and Figure 1).

During his admission a coronary angiography was preformed, demonstrating normal coronaries arteries. Meticulous repeated history obtained from the patient’s family yielded that he had been a kickboxer and a bodybuilder. After asking for all the supplements the patient was taking to be brought to our attention, a vial of Boldenone, an androgenic anabolic steroid (AAS) usually used in veterinary medicine was found. It turned out that the patient was using this illicit drug for his bodybuilding needs. After treating his failing heart, the patient was discharged for outpatient follow-up. During 1 year of follow-up he was treated with a combination of beta-blockers, ACEi and aldactone antagonist according to heart failure guidelines recommendations, with no further admission for either cardiac or non-cardiac causes. Within few months, his cardiac function was partially recovered (EF-40%), also demonstrating improvement in left ventricular chamber measurements and withdrawal of his hypertrophy (Table 1), thus establishing the reversibility of the drug effect.

<table>
<thead>
<tr>
<th></th>
<th>At admission</th>
<th>1-year follow up</th>
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<tbody>
<tr>
<td>EF%</td>
<td>20%</td>
<td>40%</td>
</tr>
<tr>
<td>LA diameter (mm)</td>
<td>3.9</td>
<td>3.8</td>
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<tr>
<td>LA area (cm²)</td>
<td>14.0</td>
<td>20</td>
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<tr>
<td>LVEDD (cm)</td>
<td>6.1</td>
<td>5.6</td>
</tr>
<tr>
<td>LVESD (cm)</td>
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<td>IVS (mm)</td>
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<td>9.0</td>
</tr>
<tr>
<td>PW (mm)</td>
<td>12</td>
<td>9.0</td>
</tr>
<tr>
<td>Right atrium</td>
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<td>Normal</td>
</tr>
<tr>
<td>Rt. Ventricle</td>
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<td>Normal</td>
</tr>
<tr>
<td>Estimated Pulmonary Pres. (mmHg)</td>
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<tr>
<td>Aortic root</td>
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<td>2.9</td>
</tr>
<tr>
<td>Ascending Aorta</td>
<td>n/a</td>
<td>2.7</td>
</tr>
</tbody>
</table>

Table 1: Echocardiography measurements.

Panel A on admission, showing severe LV dysfunction
Panel B 1 year follow up with improvement of LV function
Discussion

The use of illicit performance-enhancing supplements is common among both professional athletes and high school students. Many of the supplements have no proven merits but are associated with serious adverse reactions [1]. As its name suggests, AAS has two distinct yet overlapping effects: anabolic, promoting cell growth, and androgenic-enhancing masculine characteristics. AAS anabolic effect increase cellular protein synthesis and this result in buildup of tissue especially in muscles. AAS exert their influence on the myocytes through androgen receptor leading to left ventricular hypertrophy and dilation, and impaired contraction and relaxation [3-5]. Echocardiographic studies have shown that supraphysiologic doses of AAS lead to both morphologic and functional changes of the heart. These include a tendency to produce myocardial hypertrophy, a possible increase of heart chamber diameters, unequivocal alterations of diastolic function and ventricular relaxation, and most likely a subclinically compromised left ventricular contractile function [6].

AAS abuse has previously been associated with various cardiovascular adverse events including acute myocardial infarction, pulmonary embolism, dilated cardiomyopathy, arrhythmia and sudden death [8-19].

Boldenone undecylate is a testosterone analogue used in veterinary medicine as an anabolic steroid. Trade names include Equipoise, Parenabol, Equigan and Ultragan [7]. It is used to improve the growth and food conversion in food-producing animals. In most countries worldwide, this anabolic steroid is forbidden for human uses and meat production as it had been developed for veterinary use. Due to its potential for abuse, AAS abuse has previously been associated with various cardiovascular adverse events including acute myocardial infarction, pulmonary embolism, dilated cardiomyopathy, arrhythmia and sudden death [8-19].

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In the case presented, infection was probably not the cause of his heart failure, his medical history in the days before his admission was not suggestive for infection, on admission there were no signs of infection, either in physical examination, laboratory exams and chest X-ray. Blood cultures and panel of common respiratory infections were all negative. Ischemic reason was excluded due to demonstration of normal coronary arteries on angiography. Also, the negative TnT levels in repeated exams during his first admission exclude the possibility of myocardial injury to either ischemic or viral infection. Other possible reasons as secondary metabolic or nutritional deficiency are less likely due to the age of the patient (34 year old on his admission), the fact that there were no pre-existing diseases as liver cirrhosis or chronic renal failure and last his nutritional status on admission were normal and no signs on malabsorption. Last, his remarkable recovery of in heart function does not follow the course of viral myocarditis or ischemic cause, but more the removal of the offending agent (Boldenone).

This case presentation emphasizes two important messages: the first is that major clues for the diagnosis, especially in obscure cases, can be drawn from meticulous history taking. The second message relates to the hazards of the increasing incidence of usage of illicit drugs, mainly anabolic steroids in young adults. We hope that awareness and education will prevent further cases.

References