Resolution of Complete Atrioventricular Block in a Patient with Severe Hypothyroidism

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Introduction

Causes for syncope are multifaceted. Hormonal etiology, specifically hypothyroidism, is associated with cardiac arrhythmias [1]. Sinus bradycardia, low voltage, nonspecific T-wave changes and dissociative atrioventricular (AV) abnormalities are some of descriptive electrocardiographic features [2]. In the majority of well know clinical presentations complete AV block requires the insertion of a permanent pacemaker. However hypothyroidism related bradycardia and consequential symptomatic AV blocks could be reversible with timely and proper management.

We report a case of severe hypothyroidism resulting in insertion of transient pacemaker with favorable clinical course and fully reversible complete AV block after accordant pharmacological hormonal substitution.

Case report

Fifty-two year-old female patient without significant past medical history presented to the emergency room accompanied by her family with syncope. The patient at this time was unable to provide detailed history. According to her family social history and family history were unremarkable. Her husband reported that during the breakfast she suddenly fell from the chair on the floor without preceding symptoms and lost her consciousness for about 2 minutes. He initially tried to perform the Heimlich maneuver because he thought her symptoms are related to chocking from the breakfast. After regaining her consciousness she couldn’t remember what happened. Apart from this occasion she reported feeling tired, generalized weakness, cold intolerance and noticed being increasingly depressive within the last 2-3 months. Similar syncope episodes were not observed.

Her vital signs revealed low heart rate of 22/min was noticed, normal blood pressure 123/72 mmHg and normal temperature 36.1°C. She had a puffy face and slow speech; examination of the neck revealed no struma. Cardiovascular exam demonstrated regular heart sounds without murmurs and bruits. The jugular venous pressure was normal. The electrocardiogram demonstrated complete AV block and bradycardia with minimum heart rate of 22/min (Figure 1). The chest x-ray showed mild pulmonary edema. Laboratory findings revealed severe hypothyroidism with a thyroid-stimulating hormone (TSH) of > 100 mlU/L, free T4 of 1.3 pmol/l (normal range 9-24), and mild anemia (hemoglobin of 10.1 g/dL). Anti-thyroid peroxidase antibodies (anti-TPO) and thyrotropin receptor antibodies (TSH receptor antibodies) were negative. Comprehensive metabolic panel was remarkable for elevated transaminases (Table 1).

Initial laboratory Work up

<table>
<thead>
<tr>
<th>White blood cell count</th>
<th>10.05 x 10^9 uL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemoglobin</td>
<td>10.1 g/dL</td>
</tr>
<tr>
<td>Hematocrit</td>
<td>33.4%</td>
</tr>
<tr>
<td>Platelet count</td>
<td>233 x 10^11 uL</td>
</tr>
<tr>
<td>Sodium</td>
<td>139 mmol/L</td>
</tr>
<tr>
<td>Potassium</td>
<td>4.1 mmol/L</td>
</tr>
<tr>
<td>Serum glucose</td>
<td>117 mg/dL</td>
</tr>
<tr>
<td>Bilirubin</td>
<td>2.3 mg/dL</td>
</tr>
<tr>
<td>AST</td>
<td>121 unit/L</td>
</tr>
<tr>
<td>ALT</td>
<td>134 unit/L</td>
</tr>
<tr>
<td>Alkaline phosphatase</td>
<td>98 unit/L</td>
</tr>
<tr>
<td>TSH</td>
<td>&gt; 100 mlU/L</td>
</tr>
<tr>
<td>Free T 4</td>
<td>1.3 pmol/L</td>
</tr>
</tbody>
</table>

Table 1: Laboratory workup at admission

A temporary pacemaker was inserted with a basal ventricular rate of 60 beats per minute (Figure 2). She was treated with intravenous levothyroxine with a loading dose of 100 mcg followed by orally daily dose of 25 mcg. Transthoracic echocardiography showed normal left ventricular systolic
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**Figure 1:** ECG at admission demonstrates bradycardic rhythm (heart rate of 22 beats per minute), without signs of ischemic changes

**Figure 2:** ECG after insertion of transient pacemaker

function with grade I diastolic dysfunction and mild mitral valve regurgitation. Coronary angiography was not significant for local lesions.

Four days after initiating the treatment the AV conduction was restored and the ECG showed sinus rhythm 47/ min, the external pacemaker was removed. She was discharged home on the seventh day. Five weeks after her initial presentation the ECG revealed full recovery (Figure 3), the TSH level decreased to 27.3 mIU/L and the T4 level was within normal limit.

**Discussion**

Hypothyroidism can lead to reduction in oxygen utilization in all major organs leading to cardiac oxygen mismatch. Decline in cardiac output as well as cardiac contraction, bradycardia, increased risk of atherosclerosis, and incremental development of cardiac resistance are just some of anticipated pathophysiologic cardiovascular changes [1]. Permanent symptomatic cardiovascular dysfunction is less common in
the setting of hypothyroidism. Common symptoms include shortness of breath, cold intolerance and generalized weakness.

Clinical presentation of hypothyroidism are associated with bradycardia, syncope, nonpitting edema or symptoms related to pleural and/or pericardial effusion [3]. Atrioventricular blocks and subsequently resulting ventricular tachycardia were observed in the setting of severe hypothyroidism [4]. The exact pathophysiologic cardiac conduction abnormality remains unclear. Histopathologic features of the myocardium may represent interstitial edema, fibrotic changes and mucinous vacuolization [5,6].

Following initiation of hormonal therapy (thyroxine) recessional electrocardiographic abnormalities are rapidly observed. Hypothyroidism is one of the rare bradycardiogenic etiologies, which can show complete recovery under pharmacological management. Few cases have been reported in the literature demonstrating favorable course with complete resolution of the atrioventricular conduction abnormality [7-10]. However extensive myocardial fibrotic changes can lead to irreversible conduction delay. Affected patients fail to improve despite proper thyroid hormone treatment [2].

About 4.6 percent of the U.S. population age 12 and older has hypothyroidism [11]. The clinical course in elderly population may be perfidiously.

The numerous and constant relationships and feedback mechanisms between the liver and all endocrine organs is reflected by the fact that an alteration of one oftentimes results in the malfunction of the other. Hypo and hyperthyroidism are frequently associated with hepatic alterations as in our patient. Typically these changes resolve with treating the underlying cause [12].

**Conclusion**

Hypothyroidism is often underestimated in elderly patients. Therefore accurate evaluation of the thyroid function is recommended before implanting a permanent pacemaker in all patients with delayed atrioventricular conduction, specifically in elderly patients. Hormonal replacement (thyroxine) in the setting of hypothyroidism should be the implemental approach. This case report demonstrates a full recovery of high degree AV-block to normal sinus rhythm in severe hypothyroidism following hormone replacement.

**References**

