INTRODUCTION

Electrical injury can cause cardiac arrhythmias, myocardial/valvular rupture, structural changes in coronary arteries, pericardial effusion, and various electrocardiographic changes. In this report, we present a patient who developed sick sinus syndrome (SSS) as a result of electrical injury.

CASE REPORT

A 20-yr-old male patient suffering from dizziness and syncope attacks was referred to our hospital. He had a history of high-voltage electrical injury at the age of 12, and otherwise showed no prior history of cardiac illness. After a careful interview of the patient, we learned that the symptoms, such as dizziness and transient loss of consciousness of less than 1-min duration, developed after the accident associated with electrical energy and were precipitated by exercise.

On physical examination, the arterial blood pressure and heart rate were 120/80 mmHg and 44 beats per minute, respectively. Cardiovascular assessment was normal. On chest examination, there was a large cutaneous burn scar extending from the right subclavicular to the left subscapular area on his chest wall. Other systemic examinations were normal. Surface electrocardiogram demonstrated a junctional rhythm (Fig. 1). Echocardiographic assessment revealed a normal left ventricular size and function, and no valvular pathology. Holter monitoring showed sinus bradycardia with a sinus pause of 4.6 sec and concomitant dizziness. Routine laboratory tests, including serologic tests for collagen tissue diseases, were in normal values.

After 8-hr-fasting, a written informed consent was obtained, electrophysiological study (EP) was performed for the evaluation of the functions of sinoatrial (SA) and atrioventricular (AV) nodes. The findings were as follows: baseline cycle length (BC L), 1,200 msec; atrial-His interval (AH), 90 msec; His-ventricle interval (HV), 48 msec; Wenckebach point, 480 msec; sinoatrial conduction time, 300 msec; sinus-node recovery time (SNRT), 2,800 msec; corrected sinus node recovery time (cSNRT), 1,600 msec (Fig. 2). An injection of 2 mg of atropine failed to increase the sinus rate over 90 beats per minute, suggesting chronotropic incompetence. SNRT and cSNRT did not shorten after atropine injection.

We decided that the sinus node was not functioning normally. This abnormal function of sinus node, associated with clinical symptoms, led us to a diagnosis of SSS. Coronary angiography was performed to exclude possible coronary artery disease, even if there was no family history related to coronary heart disease or other risk. The angiography showed a normal coronary anatomy. Since the patient had chronotropic incompetence, we implanted a DDD-R pacemaker (Kappa, Medtronic, Inc., Minneapolis, MN, U.S.A.) to the left subclavian region. However, during implantation procedure, the pacing thresholds of both ventricle and atrium were higher than expected. The values measured during the procedure were as follows: p-wave amplitude, 5.6 mV; atrial pacing threshold, 1.5 V; atrial lead impedance, 476 Ohm; R-wave amplitude, 15.2 mV; ventricle pacing threshold, 1.2 V; and ventricle lead impedance, 538 Ohm.
SSS is a disease frequently encountered in the elderly population, especially in the 6th and 7th decades. It may frequently result from decreased pace cells in SA node with aging, and fibrotic and calcific process of SA and AV node regions. Other etiologies may include hypothyroidism, ischemic heart disease, a number of pharmacologic agents such as digoxin, calcium-channel blockers, beta-blockers, class-I anti-arrhythmic agents, and amiodarone, prior cardiac operation, and heart transplantation.

Heart is one of the most vulnerable organs against electricity. Various myocardial manifestations develop at the time of injury. These include asystole, ventricular fibrillation, which may cause immediate death, QT-prolongation, right bundle branch block, complete AV block, valvular or myocardial rupture, CK-MB elevations caused by myocardial injury, structural changes in the small coronary vessels, and pericardial effusion (1). In young patients, SSS is usually related to congenital heart disease (CHD) or seen after operation for correcting congenital heart disease (2, 3). In our case, there was no history of CHD or operation for CHD.

James et al. (4) reported autopsy findings of 4 patients died from electrical injury. They noticed an increased myocardial thickness in two of them, fibromuscular changes in the coronary arteries in one of them, and fatty infiltration in the vicinity of both SA and AV regions in the last one. We speculated that a probable infiltration in the SA node could be responsible for the clinical manifestations of our case. The probable myocardial injury due to electricity may explain why the levels of thresholds of both atrial and ventricular pacing were higher than expected during pacemaker implantation. Unfortunately, the patient refused endomyocardial biopsy.

Bognolo et al. (5) reported a 50-yr old patient with SSS, suggesting non-penetrating chest trauma as a causative factor. As described previously, electricity can cause mechanical trauma to the body at the same time. This kind of trauma could also cause SSS. Carleton (6) recommended that all patients injured by electrical energy should be followed because of the risk of developing cardiac manifestations for at least one year, and avoid any cardiac operation during the first six months. In our case, the symptoms developed several months after the accident and unfortunately, the diagnosis was confirmed 8 yr after the accident. This case merits attention in that there has been no case report on SSS caused by electrical injury.

In conclusion, electrical injury can infrequently cause SSS. In such cases, high pacing threshold levels can be encountered during pacemaker implantation.

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