CASE REPORT

An anticipatory complete atrio-ventricular block and state of shock due to hyperkalemia

Laachach Houssam, Anass Hbali, Bachrif Mohammed, Ilham Benahmed, Ofkir Mohammed, Nabila Ismaili, Elquafi Noha


ABSTRACT

Even hyperkalemia is not a common etiology of a complete interruption of nodal conduction; it was the case of signs of shock and complete atrio-ventricular block in association with myocardial ischemia or a low level block (4,9,10).

BACKGROUND

The important role of potassium in the electrophysiologic regulation of myocardial function is known. Any change in extracellular potassium concentration may have a significant effect on myocyte electrophysiologic gain and bring about many electric dysfunctions (1). Hyperkalemia occurs in 1–10% of hospitalized patients in medical department, with a mortality rate of 1 in 1,000 patients (2). In general, High serum potassium levels results in a gradual decrease in the excitability and conduction velocity of specialized pacemaker cells and conducting tissues throughout the heart. Typical Electrocardiogram (ECG) findings in Hyperkalemia progress from tall, “peaked” T waves and a shortened QT interval to a lengthening of PR interval and loss of P waves, and then to widening of the QRS complex, culminating in a “sine wave” morphology and death if not treated (3-6).

Hyperkalemia is thought to impair the conduction in Purkinje fibers and ventricles more than that in the AV node; although complete Atrio-ventricular block (AVB) can occur, it is one of the rarest initial presentations (7).

CASE REPORT

A 50-year-old woman who had a history of renal failure undergoing hemodialysis at 3 sessions per week, admitted into emergency department due to generalized weakness, chills, somnolence and Coldness of the extremities. The symptoms have started early the morning.

At the admission she had a respiratory rate of 24 per min, a pulse rate of 28/min, collapsed blood pressure and here blood sugar level was 120 mg/dL. Here ECG revealed a complete atrioventricular block (Figure 1) Ventricular escape rate at 20/min.

The results of Laboratory examination were as the following: Creatinine=71 mg/L; urea=4.6 g/l; Na=137 mmol/L; K=8.5 mmol/L; CRP=40 mg/L; Hb=10.8 g/dl, and INR=1.94. Besides, a respiratory acidosis was been objectified.

The patient was stabilized by vasoactive drugs: Dobutamine and Noradrenaline, infusion of glucose with insulin and gluconates of calcium.

Hyperkalemia. However, cardiomyopathies, myocarditis, structural heart disease, hyperthermia, and severe hypothyroidism are much less common causes. Acquired third-degree AV block often has a wide complex escape rhythm, with symptoms of hypo perfusion at rest or with minimal exertion. CHB may be sub-acute or chronic and relatively well tolerated providing that the block occurs at a relatively high level, (AV nodal to His bundle region). Alternatively, the patient may be very unwell, usually in acute onset in association with myocardial ischemia or a low level block (4,9,10).

RESULTS AND DISCUSSION

Diagnosis of hyperkalemia is usually based on laboratory studies, although the electrocardiogram (ECG) may contain changes suggestive of hyperkalemia. Typical ECG findings in hyperkalemia progress from tall, “peaked” T waves and a shortened QT interval to lengthening PR interval and loss of P waves, and then to widening of the QRS complex culminating in a “sine wave” morphology and death if not treated (1,2). Even with profound serum potassium elevations, the ECG cannot reliably be used to exclude Hyperkalemia or to monitor therapy designed to lower the Serum potassium concentration (3,4). Many studies demonstrated that important hyperkalemia can occur in auriículo-ventricular block (5-7). AVB has been demonstrated experimentally in animals. Potassium, according to plasma level and infusion rate, demonstrated a quantitatively different, and sometimes opposite, effect on AV conduction, intraventricular conduction, automaticity and excitability (7). The second or third degree BAV caused by hyperkalemia has been rarely documented in humans (3,8,9). Complete heart block (CHB) is the consequence of the loss of electrical transmission between the supraventricular tissues to ventricles. The lower level of block, the slower and the less reliable will be the ventricular escape rhythm. Third-degree AV block can be congenital or acquired. Three causes are known to induce an acquired CHB: Ischemic heart disease, drugs, and Hyperkalemia. However, cardiomyopathies, myocarditis, structural heart disease, hyperthermia, and severe hypothyroidism are much less common causes. Acquired third-degree AV block often has a wide complex escape rhythm, with symptoms of hypo perfusion at rest or with minimal exertion.

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Figure 1) ECG revealed a complete atrio-ventricular block.
The treatment is based on two main sources: Treatment of the atrioventricular block and palliation against tissue hypoperfusion on the one hand; On the other hand decreasing the plasma potassium concentration, combating its effects on myocytes and treating the cause of hyperkalemia. When the heart block is complete, Patients with clinical hypoperfusion or not responding to medical measures may have a continuous monitoring and treated by percutaneous pacing as a temporary or conclusive solution (9,10). Hyperkalemia should be treated as quick as possible, by antagonizing the extracellular potassium effect thus decreasing the serum potassium level by transferring on intracellular. The next line is to remove potassium from the body by 'Kayexalat' or ultimately by hemodialysis (11-13). As the case of our patient, the evolution is good after early treatment.

CONCLUSION

Complete heart block is not a common complication of hyperkalemia, it can occur when serum potassium level is very high. Syncope and heart failure is some of clinical signs. Treatment is an imperative decreasing potassium serum level and repairing the etiology of hyperkalemia. As this case has demonstrated, the heart block is mostly regressive after kalemia normalization.

CONFLICT OF INTEREST

None

AUTHORS CONTRIBUTION

All the authors have contributed simultaneously to elaborate this article

REFERENCES