A Case of suspected Sick Sinus Syndrome due to combined Beta-blocker and Calcium Channel Blocker Therapy: Anesthesia Management

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ABSTRACT

Sick sinus syndrome (SSS) is a generalized abnormality of cardiac impulse formation. Abnormalities encompassed by this syndrome may include inappropriate sinus bradycardia, sinus arrest, bradyarrhythmias, or tachyarrhythmias. We present a case of a 54-year-old hypertensive male posted for L4–L5 decompression, whom we suspected to develop SSS due to overdose of combined beta-blocker (BB) and calcium channel blocker (CCB) therapy.

Keywords: Beta-blocker–calcium channel blocker toxicity, Inotropes, Preanesthesia checkup, Sick sinus syndrome.

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INTRODUCTION

Sick sinus syndrome is a generalized abnormality of cardiac impulse formation. Abnormalities encompassed by this syndrome include sinus bradycardia, sinus arrest or exit block, and combination of sinoatrial and atrioventricular nodal conduction disturbances.1,2

Beta-blockers and CCBs are the main groups of cardiac drugs in use for several years. Unfortunately, they also remain common causes of cardiovascular collapse following accidental or intentional overdose. Toxicity is associated with significant mortality.3

We present a case of a 54-year-old hypertensive male, posted for L4–L5 spine decompression, in whom we suspect BB and CCB toxicity, which manifested as an SSS-like state.

CASE REPORT

A 54-year-old male patient weighing 70 kg was posted for lumbar spine decompression at L4–L5 level, with instrumentation. Patient was a known hypertensive since 4 years, for which he was on tablet amlodipine and atenolol combination (5–50 mg), once a day, and a known diabetic on injection insulin 4-4-4 subcutaneous. Preoperative investigations were within normal limits. Preoperative pulse was 50/min, sinus rhythm was regular, and blood pressure (BP) 100/70 mm Hg. Patient was premedicated with injection glycopyrrolate 0.004 mg/kg, injection midazolam 0.05 mg/kg, injection fentanyl 1.5 µg/kg and was induced with injection propofol 2 mg/kg. Ventilation was confirmed and patient was intubated with portex cuffed endotracheal tube no 8. After confirmation of placement of endotracheal tube, ventilation was maintained with O2 + N2O (50:50) and sevoflurane at a minimum alveolar concentration of 1.5. Patient was given injection rocuronium 1.2 mg/kg. Patient’s eyes were taped, all pressure points were padded, and patient was given a prone position.

Air entry was checked again in prone position. His pulse was 46/min, regular, and BP was 80/60 mm Hg. After cleaning and draping the patient’s back, the orthopedic surgeon located the spinal level by a 16 G intravenous (IV) cannula, which was repeated two to three times. Suddenly, patient went into bradycardia (pulse 40 beats per minute) and his BP measured 70/40 mm Hg. His carotids were palpable. Injection atropine 0.1 mg and injection mephentermine (12 + 12 + 6 mg) were administered, and patient was immediately made supine from prone position.

No response to atropine was observed, so IV atropine 0.1 mg was repeated. Patient went into cardiac arrest (pulseless electrical activity). Cardiac compressions were started immediately, and injection adrenaline 1 mg was given. Patient could be revived successfully after 1 minute following cardiopulmonary cerebral resuscitation. Patient was kept intubated and right-sided internal jugular vein was cannulated with a triple-lumen 7 French (Fr) central line.

After the effect of IV adrenaline wore off, patient’s BP continued to remain low, in the range of 60 to 70 systolic and 35 to 45 diastolic. His pulse rate remained...
around 45 to 50/min, irregular, with bradyarrhythmias and junctional rhythms. Patient was put on inotropic support. Injection dopamine was started @ 15 µg/kg/min and injection adrenaline was started @ 0.1 µg/kg/min. Blood sugar was within normal limits.

It was later discovered that the patient had received an additional dose of his antihypertensive the previous night.

Troponin T was negative.

We suspected the patient had a BB and CCB overdose. Patient required inotropic support for 24 hours and was gradually weaned off the supports and extubated.

A cardiologist’s opinion was taken, and patient was started on dual antiplatelet therapy as a prophylaxis. Patient was advised a fresh two-dimensional (2D) Echo and a coronary angiography (CAG) to rule out coronary artery disease.

Patient’s 2D Echo was within normal limits. Patient did not give consent for a CAG. He was advised a pacemaker for his sinus arrhythmias.

**DISCUSSION**

The importance of a thorough history taking in a preanesthetic evaluation has been emphasized time and again. The preoperative vital parameters of our patient were stable, and the patient did not complain of dizziness/palpitations, which are often associated with SSS (Fig. 1).

The SSS is characterized by sinus node dysfunction with an atrial rate inappropriate for physiologic requirements because the sinus node is unable to perform its pacemaking function (Figs 2 and 3).

The combination of CCB and BB is increasingly used in the management of hypertension. Anesthesia can cause myocardial depression in patients on BB and silent myocardial ischemia in those on CCB. However, the effect of anesthesia on patients receiving a combination of CCB and BB is unique and was studied by Samad et al, who found that hypotension (systolic BP <90 mm Hg) and bradycardia (HR <50/min) were more common in CCB and BB group (55%) than in BB group (15%) alone. Our patient was receiving a combination of BB and CCB.

The shock resulting from BB and CCB toxicity can be very resistant despite the use of all available catecholamines for resuscitation. Predictably, our patient needed a high dose of IV dopamine and IV adrenaline infusion for treating his hypotension and bradycardia. The IV fentanyl given as a premedication aggravated the bradycardia of the patient. A thorough history is of utmost importance not only in the preanesthetic checkup, but also in the immediate perioperative period.

Traditionally, BB toxicity was treated with glucagon and CCB with calcium gluconate infusion. However, glucagon was not available in our institution and our patient was maintaining vitals on high dose of inotropic support.
CONCLUSION

Profound and refractory hypotension can be encountered in hypertensive patients on chronic BB and CCB therapy in the absence of toxic levels. We have described our management of this case of a suspected SSS, who required a high dose of inotropic support of injection dopamine and injection adrenaline to maintain his pulse rate and BP. Also, a thorough history from the patient’s relative led us to arrive at a conclusion of an overdose of BB and CCB, leading to an SSS-like situation.

REFERENCES