Refractory Shock from Amlodipine Overdose Overcomed with Hyperinsulinemia

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Abstract
Intoxication from calcium channel blockers exhibits almost 50% mortality rates. Amlodipine is a long-acting dihydropyridine and inappropriate dosage poses a great threat for profound vasodilation, hypotension, and refractory vasopressor-resistant shock. A 72-year-old woman with unremarkable medical history presented to the emergency department due to amlodipine overdose after a suicide attempt attributed to COVID-19 pandemic severe anxiety disorder. Vital signs at presentation: heart rate 82 beats/min, arterial pressure 72/55 mmHg, and oxygen saturation 98%. Resuscitation was initiated with intravenous infusion of normal saline 0,9%, noradrenaline, and calcium chloride, while activated charcoal was orally administrated; however, blood pressure remained at 70/45 mmHg. Abruptly, she experienced acute pulmonary edema and was finally intubated. We commenced high-dose insulin infusion with Dextrose 10% infusion to maintain euglycemic hyperinsulinemia. Hemodynamic improvement occurred after 30 min, systolic blood pressure raised to 95 mmHg, and decongestion was achieved with intravenous furosemide. Insulin effect was dose-dependent and patient’s hemodynamic status improved after insulin uptitration. Eight days later, the patient was weaned from the mechanical ventilation and she was successfully discharged after 14 days. High-dose intravenous infusion of insulin up to 10 units/kg per hour appears as an inotropic agent possibly through alterations in myocardial metabolism of fatty acids and augmentation of insulin secretion and uptake. This regimen possibly exhibits additional vasotropic properties. We conclude that euglycemic hyperinsulinemia is a potentially advantageous treatment in CCB toxicity.

Keywords Amlodipine · Overdose · Hyperinsulinemia · Insulin · Calcium channel blocker

Introduction
Calcium channel blockers (CCB) intoxication is a leading cause of cardiovascular medication overdose exhibiting high—almost 50%—mortality rates [1]. Amlodipine is a long-acting dihydropyridine derivative with a half-life of 30–50 h [2]. Inappropriate dosage poses a great threat due to profound vasodilation, hypotension, and refractory vasopressor-resistant shock. High-dose insulin infusion has been described as a potential shock reversal agent induced by CCB poisoning [3].

We describe a clinical case of a 72-year-old woman effectively treated with euglycemic hyperinsulinemia after developing refractory shock due to amlodipine overdose.
Case Presentation

A 72-year-old woman with unremarkable medical history presented to the emergency department (ED) due to amlodipine overdose (200 mg of amlodipine by oral route) after a suicide attempt three hours ago. The incident was attributed to severe anxiety disorder arising from the pandemic of coronavirus disease 2019 (COVID-19).

Vital signs at presentation: heart rate 82 beats/min, arterial pressure 72/55 mmHg, oxygen saturation 98%, and temperature at 36.4 °C. Physical findings were normal and she was oriented and fluent. Electrocardiogram demonstrated that sinus rhythm and cardiac ultrasound was within normal limits.

Resuscitation was initiated with intravenous Normal Saline 0.9% (NS) and 60gr (1gr/kg) of activated charcoal was orally administrated to inhibit the absorption of amlodipine. She was transferred to the Acute Cardiac Care Unit (ACCU) due to shortage of beds in the Intensive Care Unit during the first wave of COVID-19 outbreak in Europe. Within the first hours of ACCU monitoring blood pressure decreased at 70/50 mmHg. Intravenous noradrenaline infusion was initiated with gradual uptitration to 1 mcg/kg/min. We introduced simultaneously calcium chloride 10% infusion at 15 mg/hr in an attempt to overcome the competitive blockage of calcium channels. After administrating 2 L of NS (11 ml/kg/hr) and increasing noradrenaline dose at 2.5 mcg/kg/min, blood pressure decreased further at 65/45 mmHg. Abruptly, she experienced dyspnea with mental status deterioration and signs of acute pulmonary edema provoked by a combination of iatrogenic congestion with amlodipine vasoplegia and led to intubation and mechanical ventilation.

We commenced high-dose insulin infusion (45 IU/hr) alongside with intravenous 10% dextrose solution to maintain euglycemia. After 30 min, systolic blood pressure elevated to 95 mmHg and the patient was treated with intravenous furosemide to gradually regularize volemia and oppose organ congestion. Further insulin uptitration to 60 IU/hr (1 IU/kg/hr) was followed by systolic blood pressure augmentation at 105 mmHg.

Calcium chloride infusion was terminated 3 days later, insulin infusion after 5 days, as well as noradrenaline. Gradual downtitration of insulin and noradrenaline was driven by the patient’s hemodynamic status. It is remarkable that attempts to reduce the dose of insulin during the first 3 days were followed by immediate decrease of blood pressure; insulin effect was dose-dependent (Fig. 1). Eight days later, the patient was weaned from mechanical ventilation. She was discharged after 14 days. Cardiac function tests were unremarkable and she received antipsychotic treatment.

One-month clinical and laboratory follow-up was performed and the patient exhibited improved mental status with no cardiovascular comorbidity.

Discussion

CCB act at the L-type calcium channels in heart and vessels resulting in vasodilation and reduction of both inotropism and chronotropism. Effects of CCB in pancreas include inhibition of insulin secretion and favored insulin resistance with consequent reduced glucose uptake by cardiac and vascular myocytes (the leading factor of CCB cardiovascular toxicity) [4]. Regarding the gastrointestinal tract, motility alterations and possible increase of bleeding have been described.

![Fig. 1 Systolic blood pressure (mmHg) fluctuation during insulin administration. 0 h (hrs) represents the initiation of insulin infusion](image-url)
during CCB administration [5]. CCB exhibit metabolic effects as well by modifying metabolism and kinetics of beta-adrenergic receptors and cardiac glycosides [6]. Dihydropyridines have a predominant effect on vascular motility and do not interfere with cardiac conduction properties, pacemaker cells, and contractility. This selectivity, however, fades in significant overdose resulting in bradycardia, conduction abnormalities, and reduced inotropism (Fig. 2) [1].

The patient presented in the ED as hypotensive and asymptomatic, nevertheless, the situation rapidly evolved in refractory shock. Noradrenaline, fluids, and calcium exerted no clinical effect and hypotension was followed by respiratory insufficiency and mental status alteration in the context of acute pulmonary edema. The latter is a frequent complication of CCB overdose and the most dominant pathophysiological pathway is massive pulmonary capillary dilatation and increased permeability, possibly sharing a similar mechanism as peripheral edema caused by pre-capillary vasodilatation [7].

The CCB-induced shock is vasogenic and first-line management includes fluid resuscitation and vasopressors. High intravenous calcium doses as calcium chloride or calcium gluconate may be thought to antagonize the CCB toxicity by increasing serum ionic calcium level enough to overcome calcium channels blockade and drive calcium into the cells. Atropine and cardiac pacing are required in the cases of bradycardia and conduction abnormalities. Lipid emulsion, methylene blue, and glucagon are included in our quiver as additional alternative therapeutic approaches, and ECMO (Extracorporeal membrane oxygenation) has been described as a potential bridging therapy until the involved drug has been metabolized [8]. CCB are strongly bounded to plasma proteins so that conventional hemodialysis is not effective. Gastrointestinal decontamination with gastric lavage, whole-bowel irrigation, and activated charcoal remains the only option for preventing the drugs’ absorption, but should be ideally administered shortly after ingestion [9].

CCB plasma levels correlate directly with increased serum glucose concentrations primarily because they block pancreatic L-type calcium channels leading to inhibition of insulin release [10]. This “diabetogenic” effect triggered investigation on CCB toxicity monitoring through measuring serum glucose levels [11]. Case reports and animal studies suggest that hyperinsulinemia improves hemodynamics in CCB overdose. The dose varies among 1–10 units/kg/hour, while glucose and potassium levels should be meticulously monitored and corrected [12]. Some reports support a loading bolus dose of insulin followed by maintenance infusion and other infusion drip only for adequate control of blood pressure and organ perfusion [13]. Hyperinsulinemia acts through improving cardiac contractility by overcoming carbohydrates’ starvation and therefore providing heart’s metabolic demands [14]. Experimental models demonstrated that insulin accelerates myocardial lactates oxidation and facilitates calcium handling from myocardial cells. The

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**Fig. 2 Calcium Channel Blockers toxicity effects**

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**Calcium channel blocker toxicity**  
**L-type Calcium channels inhibition**  
Vessels  
- Vasodilation (decreased vascular tone)  
- Insulin resistance  
- Decreased afterload  
Gastrointestinal tract  
- Motility alterations  
- Increased risk of bleeding  
Pancreas  
- Insulin secretion inhibition  
Heart  
- Insulin resistance  
- Reduced inotropism/chronotropism (non-dihydropyridines)
The aforementioned effects render insulin as an effectual inotrope, even better than epinephrine which increases oxygen consumption [15].

In our case report, we exhibited that euglycemic hyperinsulinemia was the key to stabilize the patient in accordance to existing literature. As the dose regimen has not been well established, we commenced the infusion at less than 1 unit/kg/hour with gradual uptitration and observed that blood pressure elevation was proportional to the insulin dose. Interestingly, any dose reduction led to blood pressure drop. Even after intubation and concomitant propofol administration, the patient remained stable receiving insulin. Interestingly, our patient’s systolic function was normal and remained unaffected during the hospitalization, therefore, there was no need for inotropic support. Based on the hemodynamic stabilization after insulin initiation, we could assume that the insulin exerted pleiotropic vasotropic actions. The effectiveness of the euglycemic hyperinsulinemia possibly renders this regimen as a first-line treatment for CCB toxicity-induced shock.

Conclusion

We conclude that hyperinsulinemia could be an advantageous treatment for CCB intoxication and possibly included in the first-line regimens. Hemodynamic response is in proportion with insulin administered dose, and for maximal beneficality, careful uptitration is suggested in conjunction with the conventional treatment.

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Data Availability No additional data are available. This case report is supported by the facts and data included in the manuscript.

Declarations

Conflict of interest None.

Ethical Approval No ethics approval is required.

Informed Consent Signed consent has been obtained.

Consent for Publication The patient and the authors consent for publication.

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