Iatrogenic Hypoglycemia Induced by Valproic Acid in an Adult Patient

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Abstract

Literature on antiepileptic-induced iatrogenic hypoglycemia is scanty. Due to its broad spectrum of activity and mechanisms of action, valproic acid (VPA), a fatty acid, is one of the most widely prescribed epilepsy treatments worldwide. Hence, we describe an adult epileptic patient, in whom persistent, otherwise unexplained, hypoglycemia was most likely induced by VPA, as suggested by the VPA and glucose blood level time course. Indeed, no further hypoglycemic episodes occurred after VPA discontinuation and the diagnostic work-up ruled out other possible causes of hypoglycemia. This case supports the hypothesis that VPA may induce hypoglycemia, even in the absence of a VPA hepatotoxicity syndrome, due to still not well-defined metabolic mechanisms of action. Moreover, it emphasizes the fact that an iatrogenic pathogenesis should be considered if an apparently unexplained hypoglycemia occurs in a patient on chronic therapy with VPA, even at a therapeutic dosage.

Keywords: Hypoglycemia; Valproic Acid (VPA); Epilepsy; Side Effects; Diabetes Mellitus; Iatrogenic Effects; Seizure

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INTRODUCTION

Valproic acid (VPA), a branched short-chain fatty acid, is a widely used anti-epileptic drug with a broad spectrum of activity and mechanisms of action [1]. It is used for the treatment of several types of seizures and epileptic syndromes, including generalized and focal seizures [2]. Moreover, it is commonly used for migraine prophylaxis and in the treatment of bipolar disorder, as a mood tone stabilizer. VPA, as a therapeutic agent, is commercially available as Depakote®, Depakote ER®, Depakene®, Depacon®, Stavzor®, Mylproin®, Ergenyl®, Dipropylacetic acid, Myproic Acid, and Convulex®.

To date, its mechanism of action has not yet been fully clarified. Although VPA does not directly interact with postsynaptic GABA receptors, it does increase regional neuronal GABA concentrations by both inhibiting its metabolism and increasing its synthesis [1]. Preclinical research on animal models has shown the involvement of ion channels, monoamines, the corticotropin-releasing
factor, and intracellular signaling proteins in the mechanism of action of VPA [2,3]. It has also recently emerged that VPA inhibits histone deacetylation and regulates the transcription of various genes. This effect implies it has a potential role as anti-cancer drug and as possible antidiabetic agent [4].

However, it has been demonstrated that VPA administration may cause numerous side effects and/or pharmacological interference and, thus, requires strict clinical and laboratory control, with serum level monitoring [5]. Indeed, VPA can lead to hypoglycemia due to an inappropriate antidiuretic hormone secretion. It can also cause hematologic toxicity (thrombocytopenia, leucopenia, macrocytic anemia), severe liver damage, kidney failure, hypothroidism, hyperammonemia, rhabdomyolysis, hyperandrogenism, pancreatitis, as well as teratogenicity. The mechanisms underlying most of these side effects have not yet been completely understood [3,6,7].

It has been reported that drug-induced hypoglycemia as a cause of acute medical admissions ranges from 0.1% to 1.7% [8]. Literature reports several types of non-diabetic drugs that may induce hypoglycemia, including: NSAIDs, analgesics, antibacterials, antimalarials, pentamidine, ß-adrenergic receptor antagonists (ß-blockers), antiarrhythmics, ACE-inhibitors, fibrates, and antidepressants [9]. Ben Salem et al. also reported that some antiepileptic drugs (gabapentin, phenotoin, topiramate) caused iatrogenic hypoglycemia [9]. It was hypothesized that the underlying mechanisms were insulin release or the reduction of its clearance, and/or an interference in the glucose metabolism [9].

Herein, we report a case of persistent hypoglycemia episodes, which were most likely induced by VPA, administered at an appropriate dosage in an epileptic patient.

**CASE PRESENTATION**

A 49-year-old male, affected by an epileptic syndrome from the birth, was admitted to our Neurology Unit for recurrent focal seizures to bilateral tonic-clonic seizures. His clinical history included diabetes mellitus (DM) type 2 treated by metformin, oligophrenia, associated with the epileptic syndrome, which was probably structural, related to perinatal hypoxic damage, and secondary psychosis, treated with risperidone.

He had been on VPA therapy, at 1000 mg bid, for about a year before being hospitalized in our department. Two EEGs performed in the previous 2 months had shown no evidence of epilepsy.

The patient had been admitted to our Emergency Department the previous month for respiratory failure and bilateral pneumonia, which had evolved into septic shock requiring Intensive Care. Although recurrent hypoglycemia was observed, it was attributed both to the septic state and his reduced food intake.

His hepatic function was normal, i.e., alanine transaminase 30 mg/dl, aspartate transaminase 35 mg/dl, and ammonium levels were within normal levels (35 mg/dl), as was the renal function.

When he arrived in our Division, he was still on maintenance therapy, i.e., methylprednisolone (40 mg daily), prescribed when he had been admitted to the ICU, and risperidone 1 mg daily: no variation was made to the treatment. He had continuous glucose infusions.

The patient was transferred to an Internal Medicine Department after a week, where the hypoglycemia episodes persisted and a generalized epileptic seizure occurred, even though his VPA plasma levels were in the therapeutic range.

A more in-depth medical history brought to light two previous diabetology consultations for hypoglycemia in the previous year. As the first hypothesis was that his symptoms were attributable to metformin, it was discontinued. Clinical investigations were made, firstly a diabetology assessment, dosing of C-peptide (it was within normal limits), and an abdomen CT scan.

The simultaneous determination of insulinemia and blood glucose, the insulin-blood glucose ratio and the concomitant determination of insulinemia and blood glucose during 24-48-72 hours of fasting were unremarkable.

No neuroendocrine tumor was found by the abdomen CT scan, and neuron-specific enolase and chromogranin A were normal.

An iatrogenic effect was then hypothesized, with the suspicion that VPA played some kind of role. Therefore, the patient was transferred back to the Neurology Unit for rapid VPA withdrawal (500 mg bid for two days, then withdrawn), switching to levetiracetam (LVT) at 1000 mg bid.

The hypoglycemic episodes persisted for another week, especially during the morning.
It took 7 days for the blood glucose values to normalize after VPA withdrawal (Figure 1). A steady normoglycemic state was then maintained and epilepsy was kept under good control by levetiracetam.

**DISCUSSION**

VPA is known to have several side effects, including teratogenicity and hyperammonemia, with possible serious encephalopathy [3,6] and it is difficult to predict the individual response to treatment and the appearance of side effects.

The recent developments of pharmacogenomics has brought to light the fact that there are numerous genes which may influence VPA metabolism, efficacy and safety. This may, in part, explain the individual variability of plasma levels and the onset of some serious side effects, including liver toxicity and teratogenicity [7].

Our patient had type 2 DM and a good metabolic compensation. He was hospitalized for sepsis associated with respiratory failure and admitted to the ICU, where the emerging episodic hypoglycemia was attributed to his septic state. But the hypoglycemia progressed, despite the fact that the inflammatory state remitted. A diagnostic work up investigated into the possible pathogenesis of the hypoglycemia and reasonably ruled out: an iatrogenic nature (metformin), organic endogenous causes (insulin, congenital metabolic diseases, and extra-pancreatic cancers), as well as functional endogenous causes (postprandial reactive hypoglycemia, pituitary and cortico-surrenalic insufficiency, liver disease, malnutrition), and a VPA hepatotoxicity syndrome.

As the C-peptide connects the insulin A-chain to its B-chain in the proinsulin molecule, it was dosed. Proinsulin is the precursor of the insulin synthesized in the B-pancreatic cells. As C-peptide and insulin are released into the circulation in equimolar quantities by pancreas B-cells, assessing circulating C-peptide levels provides an indirect evaluation of insular B activity. C-peptide assessment is often preferred to insulin assessment as it is metabolized in the proximal renal tubes and not by the liver and has a slower metabolism and lacks cross-reactivity with antibodies directed towards insulin [10]. In our patient, the C-peptide tuned out to within the normal limits.

Moreover, the VPA level decrease observed after withdrawal was followed by a blood glucose increase (Figure 1), supporting the hypothesis that VPA played a direct role in hypoglycemia as a side effect, even though the decline in VPA serum concentrations took somewhat longer than expected.

This time course of VPA level decrease and glucose rise, after a week interval from the VPA nadir, pointed to a possible role of other factors, mainly pharmacological (the other drugs co-administered), but also clinical (the comorbidities), which might have af-

**Figure 1. Glucose and Valproic Acid level time course (on the abscissa the days from admission to the Neurology Unit)**
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Iatrogenic Hypoglycemia Induced by Valproic Acid in an Adult Patient has a metabolic influence on insulin release at a pancreatic level [9]. Moreover, it has been hypothesized that VPA acts as a substrate for the fatty acid β-oxidation pathway, leading to carnitine consumption. Therefore, when fatty acid β-oxidation is inhibited by a low carnitine level, the fasting hepatic gluconeogenesis may be decreased, leading to hypoglycemia [17].

CONCLUSIONS

Antiepileptic-induced iatrogenic hypoglycemia, in particular VPA-induced, has not been commonly reported, nor is it included in the side effects described in the VPA datasheet. The case herein reported in an adult patient supports the hypothesis that VPA may induce hypoglycemia, by mechanisms and metabolic effects that remain to be defined, even at a therapeutical dosage.

Therefore, we strongly suggest that whenever there is a case of apparently unexplained hypoglycemia in patients on chronic VPA therapy, the possibility of there being an iatrogenic pathogenesis should be carefully evaluated.

As currently only anecdotal reports are available, the effect of VPA on glucose metabolism should be better clarified by future investigations. Indeed, a better understanding of these mechanisms may well lead to better and safer management of VPA therapy and may even prompt research on the possible, novel indications for metabolic control in patients with glucose intolerance.

Key points

- VPA may induce hypoglycemia, even at a therapeutical dosage.
- Cases of apparently unexplained hypoglycemia in patients on chronic VPA therapy should be carefully evaluated to rule out an iatrogenic pathogenesis.
- Hopefully, future studies will shed light on the mechanisms underlying the hypoglycemia induced by VPA and the effect VPA has on the glucose metabolism.

Consent to publication

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REFERENCES