Pregnancy Complicated by Hyperinsulinemic, Non-Insulinoma Hypoglycemia Syndrome: Case Report

Enio Luis Damaso¹, Conrado Sávio Ragazini², Mariane Nunes de Nadai¹, Patrícia Moreira Gomes³, Raphael Del Roio Liberatore Junior⁴, Elaine Christine Dantas Moisés³

Abstract

Hyperinsulinemic hypoglycemia is an unusual metabolic disorder characterized by frequent episodes of hypoglycemia caused by uncontrolled insulin release. In pregnancy it can result in adverse perinatal outcomes. We report a rare case of a pregnant woman diagnosed with this disorder. During the first pregnancy, drug treatment was necessary, and the second pregnancy occurred after partial pancreatectomy, which allowed good control with nutritional support. Hyperinsulinemic hypoglycemia syndrome complicating pregnancy, although very rare, should be included in the differential diagnosis of hypoglycemia during pregnancy.

Keywords: Hypoglycemia; Hyperinsulinism; Pregnancy; High-Risk; Case Report

Introduction

Hyperinsulinemic hypoglycemia is an unusual metabolic disorder caused by uncontrolled insulin release, secondary to either neoplastic or functionally defective pancreatic beta cells. The latter is known as Noninsulinoma Pancreatogenous Hypoglycemia Syndrome (NIPHS) or Congenital Hyperinsulinism, which is usually seen in newborns and rare in adults – only 0.5-5% of all organic hyperinsulinemic diseases in adults are caused by this condition [1]. Because this condition is rare in adults, histopathologic and clinical features are not well known [2].

There are 3 cases published in scientific literature about this syndrome in pregnancy. The major concern of NIPHS and pregnancy is the well-known association between adverse perinatal outcomes and hypoglycemia [3,4], which occurs frequently in these patients.

This article describes the obstetric and clinical care involved and the evolution of two pregnancies of a woman who had the diagnosis of hyperinsulinemic hypoglycemia due noninsulinoma pancreatogenous hypoglycemia syndrome. This paper intends to increase clinical knowledge about this syndrome and report a successful case of treatment.

Case report

A 21-years old primigravida presented for the first time at 27 weeks and 5 days to our high-risk pregnancy antenatal clinic in 2013. She had a personal history of frequent episodes of hypoglycemia since she was three months of age. Her medical background included a family history of hypertension and dyslipidemia. At the presentation, her body mass index was 19.5 kg/m² (body weight 58.4kg, height 160cm). The initial approach consisted of
In her second pregnancy, 3 years after pancreatic surgery, she was 28 years old and started antenatal follow-up at 8 weeks of gestation. At the presentation, her body mass index was 21.8 kg/m² (body weight 55.8 kg, height 160 cm). Nutritional support was based on a fractional diet, not fasting for long periods even during the night. Home glucose monitoring showed only a few episodes of glycemic levels between 60 and 70 mg/dL (3.3-3.8 mmol/l) during pregnancy with no need of additional care. At 39 weeks of gestation induction of labor was indicated and oxytocin after a cervical ripening balloon was used to induce labor. The patient had a vaginal delivery of a healthy female newborn, weighing 3155g (INTERGROWTH-21 40th percentile [5]), Apgar scores at 1 and 5 minutes of 8 and 9. They were discharged after 3 days.

We found a mutation in KCNJ11 gene, g.1350C>G (NM_000525) related to congenital hyperinsulinism. The patient refused pancreatic surgery after delivery as definitive treatment and dropped clinical follow-up for 3 years.

After 3 years she restarted follow-up at our center as the episodes of hypoglycemia had intensified. Verapamil was prescribed as treatment with no control of glycemic levels. Therefore, pancreatic surgery as definitive treatment was once again proposed. Subtotal pancreatectomy was performed, removing approximately 85% of the pancreas. Anatomopathological examination showed pancreatic islets with hypertrophic cells and giant nucleus in a diffuse distribution. Figure 1 shows this examination. She was discharged with normal glycemic levels and no exocrine dysfunction.

Table 1: Serum values of tests collected in 2013 of the reported case, as well as normal reference values and suggestive of NIPHS.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Patient</th>
<th>Normal range</th>
<th>NIPHS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glucose (mg/dL)</td>
<td>19</td>
<td>&gt;70</td>
<td>&lt;55</td>
</tr>
<tr>
<td>Insulin (microU/mL)</td>
<td>3</td>
<td>&lt;3</td>
<td>≥3</td>
</tr>
<tr>
<td>C-peptide (nmol/L)</td>
<td>1.53</td>
<td>&lt;0.2</td>
<td>≥0.2</td>
</tr>
<tr>
<td>Proinsulin (pmol/L)</td>
<td>7.5</td>
<td>&lt;5</td>
<td>≥5</td>
</tr>
<tr>
<td>Beta-hydroxybutirate (mmol/L)</td>
<td>-</td>
<td>&gt;2.7</td>
<td>≤2.7</td>
</tr>
</tbody>
</table>

Reference: Cryer et al. [13] and American Diabetes Association [33]
Hyperinsulinemic hypoglycemia is characterized by unregulated insulin release, leading to persistently low blood glucose concentrations with lack of alternative fuels [6]. Congenital Hyperinsulinism is characterized by endogenous hyperinsulinemic hypoglycemia that is not caused by an insulinoma. Pancreatic specimens from patients show beta cell hypertrophy, islets of Langerhans with enlarged and hyperchromatic nuclei, and increased islets budding from peri ductular epithelium [7-11]. Nesidioblastosis is a term that was introduced for the first time by Laidlaw in 1938 [10] to define the diffuse proliferation of pancreatic islet cells budding from ductal epithelium. After the knowledge of genes involved in the syntheses and secretion of insulin, the name nesidioblastosis was changed to congenital hyperinsulinism [12].

We report a rare case of Congenital Hyperinsulinism on a patient who has been pregnant twice. During her first pregnancy, laboratory tests (blood glucose concentrations, insulin, C peptide and pro-insulin) (Table 1) and radiological exams (US and MRI) were compatible with NIPHS. Drug treatment was prescribed due to insufficient glycemic control with diet alone. The second pregnancy occurred after a partial pancreatectomy, which resulted in satisfactory glucose levels with nutritional support alone.

In adults, hypoglycemia due to endogenous hyperinsulinism can be caused by the following [13]: beta cell secretagogues, such as a sulfonylurea; beta cell tumors (insulinoma); functional beta cell disorders that can occur as a feature of the noninsulinoma pancreatogenous hypoglycemia syndrome; after gastric bypass and insulin autoimmune hypoglycemia.

If nutritional support fails to prevent hypoglycemia, some drugs have been proposed to improve hypoglycemic symptoms in patients with NIPHS [15-18]. Octreotide is a somatostatin analog that inhibits insulin secretion [19]. Verapamil is a calcium-channel blocker that also acts in beta cells inhibiting insulin secretion [16]. Diazoxide is a non diuretic benzothiadiazine and an agonist of the sulfonylurea receptor, used for inhibiting insulin secretion in such cases of hypoglycemia [20]. Finally, acarbose, an alpha-glucosidase that inhibits postprandial glucose and its insulin response, can also be used [21].

For patients with NIPHS and severe postprandial hypoglycemia (e.g., neuroglycopenia with loss of consciousness) or with symptoms despite medical management, surgery is the pillar of therapy. In case series and reports, partial or subtotal pancreatectomy successfully relieved hypoglycemic symptoms in the majority of patients [8,9,14].

Early pregnancy is associated with a reduction in fasting glucose levels by 5 to 10 mg/dL [22,23], presumably secondary to an increase in glucose uptake by the fetoplacental unit, as well as a decrease in glycogenolysis and hepatic glucose production. As pregnancy progresses, a state of "accelerated starvation" develops with increased glucose consumption and reduction in hepatic glucose production, resulting in a lowered blood glucose and a compensatory increase in fatty acid utilization [24]. Cousins et al. [25] demonstrated the wide excursions in glucose and insulin necessary to maintain relative euglycemia throughout pregnancy. Therefore, early pregnancy, in particular, is associated with a reduction in peripheral glucose levels and can potentially increase the risk of hypoglycemia [26].

The occurrence of hypoglycemia during pregnancy raises concerns for the well-being of the pregnant woman and her fetus. It has been hypothesized that hypoglycemia during pregnancy can induce potential adverse effects that lead to fetal malformations, small for gestational age and poor neuropsychiatric development [4,27]. The association between the level of hypoglycemia and diabetic embryopathy remains unclear [28].

No fetal malformation or growth retardation has yet been reported in pregnant patients with insulinomas, even in a patient in whom the diagnosis of insulinoma was not made until delivery [29-31]. This case confirms this, as both newborns had similar birth weight percentiles (33 and 40, respectively). On the other hand, hypoglycemia in pregnant women with insulinomas can have serious maternal consequences, especially repeated seizures, and misdiagnosis can be even fatal [32]. Insulinomas can be surgically treated during the second trimester of pregnancy.

There are 3 more cases published of pregnancy in congenital hyperinsulinism patients. In all cases, the treatment used was octreotide.
In the first case, published in 2014, a 36 years woman with nesidioblastosis was treated until 33 weeks of gestation with octreotide with no effects on the newborn [34].

In the second case, published in 2017, a 24 years old woman with congenital hyperinsulinism caused by a GCK mutation, was treated with octreotide from 23 to 35 weeks of gestation. Intrauterine growth retardation was observed although she was also a smoker [35].

In the third one, published in 2022, a 29 year old woman was treated with octreotide until 38 weeks of gestational age with no complications [36].

In our case, during the first pregnancy, we used diazoxide, an anti hypertensive drug used for treatment of congenital hyperinsulinism and a category C drug for use during pregnancy. During 2 weeks of using diazoxide and glucocorticoid, the nutritional approach and medication were able to maintain normal glucose levels. During the second pregnancy and after pancreatectomy, she had no hypoglycemic episodes. Both newborns were normal.

Octreotide and diazoxide can cross the placental and potentially affect the fetus [37,38]. There was no robust experience on treatment of congenital hyperinsulinism during pregnancy and this is probably the first case of diazoxide use.

It is also uncommon that a woman who had a pancreatectomy becomes pregnant and gives birth to a normal newborn.

**Conclusion**

This report reminds us that episodes of hypoglycemia can be unrelated to the treatment of diabetes mellitus. Physicians need to be aware of differential diagnoses and their main therapeutic approaches. The already lower blood glucose levels during pregnancy associated with NIPHS pose even higher risks for these women and their offspring.

Noninsulinoma pancreatogenous hypoglycemia syndrome complicating pregnancy, although very rare, should be included in the differential diagnosis of hypoglycemia during pregnancy.

We report the use of diazoxide for two weeks during pregnancy with partial control of hypoglycemic episodes and a successful pregnancy after pancreatectomy.

**Ethical Approval**

The Ethics Committee of the University Hospital, Ribeirão Preto Medical School approved this study under protocol number 5.043.448.

**Contributions**

ELD e CSR foram os responsáveis por reunir as informações clínicas e obstétricas do caso relatado. PMG e RDRL foram responsáveis por reunir as informações endocrinológicas e moleculares do caso relatado. MNN e ECDM foram respondidos pela revisão final do manuscrito. Todos os autores contribuíram para a confecção e revisão do manuscrito.

**Conflict of Interest**

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

**References**


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