Diagnosis and management of congenital hyperinsulinism: a case report

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Introduction

Congenital hyperinsulinism (CHI) is a common cause of persistent hypoglycaemia in neonates, with an estimated incidence of one in 50 000 live births (1). The molecular basis for CHI can be established in approximately 50% of the cases and is categorised as channelopathies – defects in the ATP-sensitive potassium (K<sub>ATP</sub>) channels that regulate insulin secretion, and metabolopathies – defects in metabolic pathways altering β-cell function (2). Channelopathies form the largest group and can be further subdivided into focal and diffuse forms. In contrast to metabolopathies, most channelopathies do not respond to pharmacological treatment and require surgical removal of either focal lesions or, in diffuse forms, near total (>97%) pancreatectomy. This case report describes the diagnostic and therapeutic challenges in a patient with CHI. More extensive background information on CHI can be found elsewhere (2).

Case report

A girl born large for gestational age (birth weight > +2.5 SDS) after an uneventful pregnancy, presented 1 h after birth with apnoea and convulsions, caused by severe hypoglycaemia (serum glucose < 0.6 mmol/l). Hypoglycaemia was treated with 5 ml/kg 10% glucose i.v. followed by a continuous i.v. glucose infusion. Laboratory investigations at the time of hypoglycaemia (Table 1) showed hyperinsulinaemia, hypoketotic, hypofattyacidaemic hypoglycaemia, and a glucose infusion rate of 19 mg/kg per min was required to maintain serum glucose above 3 mmol/l. A diagnosis of ‘hyperinsulinism’ was made. Maternal gestational diabetes was excluded by an oral glucose tolerance test, and at the end of the first week the patient was transferred to our hospital for further assessment and treatment.

She had no dysmorphic features suggestive of a syndromic cause for CHI. Treatment was started with diazoxide (15 mg/kg per day in three doses) and hydrochlorothiazide (7 mg/kg per day in two doses), which resulted in a 25% decrease in glucose requirement. Higher doses of diazoxide were not tolerated, so a continuous i.v. infusion of octreotide 15 µg/kg per day was added, which produced a modest effect on glucose requirement. Increasing the octreotide dose was ineffective and glucagon 5 µg/kg per h i.v. was needed to normalise glucose intake.

The poor response to diazoxide and the results of additional investigations (Table 1) indicated a channelopathy as the most likely cause of CHI. Analysis of the ABCC8 and KCNJ11 genes (encoding for the SUR1 and Kir6.2 proteins of the K<sub>ATP</sub> channel respectively) was undertaken. As the mother was from central Finland and the father had Finnish ancestors, initial screening focussed on the Finnish founder valine 187 aspartic acid substitution in the SUR1 protein (3).

The patient was transferred to another centre for [18F] fluoro-L-DOPA positron emission tomography (PET), in order to differentiate between focal and diffuse CHI. It was requested that medication need not be discontinued and carbidopa was given before the PET. No pancreatic uptake of [18F] fluoro-L-DOPA was observed; the patient was therefore transferred to a third centre for diagnostic laparoscopy with pancreatic biopsies. No focal lesions could be identified and a large biopsy (approximately 50% of the pancreas) was taken. Immunohistochemistry of the pancreatic biopsy showed diffuse intense staining for insulin and pro-insulin compatible with a diffuse form of CHI. Mutation analysis revealed a maternally derived V187D substitution. The result excludes the focal form of CHI, which results from a paternally derived mutation in SUR1 combined with a loss of maternally derived unaffected allele in the hyperplastic islets (4).

Diazoxide and hydrochlorothiazide were restarted after surgery and the patient remained normoglycaemic with a glucose intake of 7.5 mg/kg per min. Most patients with a channelopathy do not respond to diazoxide, but several attempts at stopping diazoxide resulted in an increase in glucose requirement. No hypoglycaemic events occurred over the next 4 weeks, including monitoring glucose levels continuously during 96 h using a continuous glucose monitoring system (Medtronic Minimed, Inc., Northridge, CA, USA).

The patient was, however, readmitted 3 weeks after discharge for recurrent hypoglycaemia. Each therapeutic...
intervention to reduce insulin secretion had only a temporary effect. Six weeks later, glucose intake was still 11 mg/kg per min with starch-enriched formula, administered by a 3-hourly feeding schedule and continuous nocturnal feeding via a gastrosome tube. Maximum tolerable doses of diazoxide 16 mg/kg per day, hydrochlorothiazide 8 mg/kg per day and octreotide 25 μg/kg per day continuously s.c were needed. Consequently, laparoscopic near-total pancreatectomy was performed, leaving only a small slip of pancreatic tissue remaining beneath the distal end of the common bile duct. During surgery, ectopic pancreatic tissue was removed from the duodenal wall.

Within a week, diabetes mellitus had developed and required insulin therapy. However, the patient became hyperinsulinaemic again and medical treatment for hyperinsulinism was reinstated. As maximal therapy did not prevent hypoglycaemia completely, she was transferred to a specialised centre abroad for a third pancreatectomy. Hypoglycaemia did not recur during this period and after 1 month of being euglycaemic, she was discharged.

The paternally derived SUR1 mutation was subsequently identified (splice site mutation intron 10:c.1630 + 1 G>T). Over the next 6 months, hypoglycaemia (glucose < 2.6 mmol/l) occurred less than once a month and at the age of 1 year, psychomotor development was completely normal. She was admitted for an episode of gastroenteritis at 13 months of age. Diazoxide treatment was also stopped. Continuous s.c. octreotide was successfully replaced by lanreotide 60 mg s.c. every 4 weeks by 18 months of age. Her height develops along the +1.5 and +2.0 SDS line (target height +0.9 SDS) and weight for height between +1.5 and +2.0 SDS.

Table 1 Results of laboratory investigations at the time of hypoglycaemia.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Result</th>
<th>Response to hypoglycaemia</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glucose</td>
<td>1.8 mmol/l</td>
<td>Adequate</td>
</tr>
<tr>
<td>Growth hormone</td>
<td>98.3 mU/l</td>
<td>Adequate</td>
</tr>
<tr>
<td>Cortisol</td>
<td>1950 nmol/l</td>
<td>Adequate</td>
</tr>
<tr>
<td>Insulin</td>
<td>12 mU/l</td>
<td>Inadequate suppression</td>
</tr>
<tr>
<td>C-peptide</td>
<td>0.62 nmol/l</td>
<td>Inadequate suppression</td>
</tr>
<tr>
<td>β-OH-butyrate</td>
<td>0.35 mmol/l</td>
<td>Inadequate ketogenesis</td>
</tr>
<tr>
<td>Total free fatty acids</td>
<td>1078 μmol/l</td>
<td>Limited lipolysis</td>
</tr>
<tr>
<td>Urine ketone bodies</td>
<td>Absent</td>
<td>Inadequate ketogenesis</td>
</tr>
<tr>
<td>Ammonia</td>
<td>44 μmol/l</td>
<td></td>
</tr>
<tr>
<td>Acyl-carnitine profile</td>
<td>Normal</td>
<td>—</td>
</tr>
<tr>
<td>Transferrin iso-electric focussing</td>
<td>Normal</td>
<td>—</td>
</tr>
</tbody>
</table>

Conclusions

- Differentiation between focal and diffuse forms of CHI can be achieved by [18F] fluoro-l-DOPA PET or by laparoscopy with pancreatic biopsies. In addition, a paternally derived pathogenic mutation in SUR1 or Kir6.2 excludes focal CHI.
- There is insufficient evidence to support the laparoscopic approach for therapeutic pancreatectomy in diffuse forms of CHI.
- Somatostatin analogues, such as octreotide and lanreotide, can be used as long-term treatment in patients with persistent hyperinsulinism despite therapeutic pancreatectomy.
- Management of CHI requires medical intensive care, modern imaging and surgical expertise combined in designated specialist centres.
References


Received 26 June 2006
Accepted 27 July 2006