Persistent hyperinsulinemic neonatal hypoglycemia caused by focal nesidioblastosis. Preoperative diagnosis by pancreatic venous sampling

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ABSTRACT
A female infant with persistent hyperinsulinemic hypoglycemia is described. Persistent euglycemia could not be achieved by diazoxide, somatostatin and frequent, low protein feeds. Preoperative localization of focal insulin hypersecretion by percutaneous pancreatic venous sampling allowed excision of only a small pancreatic portion which was followed by normalization of blood glucose levels.

Key words: persistent hyperinsulinemic hypoglycemia, focal nesidioblastosis, pancreatic venous sampling, diazoxide, somatostatin, pancreatectomy.

INTRODUCTION
Hyperinsulinism, although rare, is the most common cause (up to 50%) of persistent infantile hypoglycemia (PHHI)¹. PHHI is usually associated with diffuse or focal nesidioblastosis and only rarely with pancreatic adenoma². There are sporadic and familial forms of the disease with diverse molecular defects³⁴. In the recessively inherited form, mutations have been demonstrated in the gene for the plasma membrane sulfonylurea receptor (SUR1) or its associated inwardly rectifying potassium channel (Kir6.2) of the beta cells⁵. In the dominantly inherited form, a mutation in the glucokinase gene has been identified⁶. The hyperinsulinism-hyperammonemia syndrome is caused by mutations in the glutamate dehydrogenase gene⁷. Finally, in cases of focal PHHI, a specific loss of maternal alleles of the imprinted chromosome region 11p15 has been detected in the hyperplastic area of the pancreas². The cellular hyperproliferation seems to be secondary to the expression of IGF-2 in the absence of H19³.

We report a female infant presenting with hyperinsulinism due to focal adenomatous hyperplasia of the pancreas. Recognition of the focal lesion by percutaneous pancreatic venous sampling (PVS), preoperatively, permitted resection of the lesion and avoidance of blind, 95% pancreatectomy.

CASE REPORT
The patient, a female, was the first child of unrelated Greek parents. She was born at term after an
uncomplicated pregnancy, by vaginal delivery. The birth weight was 4135 g and the length 47 cm. She had plethora and hepatomegaly at birth. No cardiorespiratory resuscitation was required. Soon after delivery she had hypotonia and the blood glucose (BG) concentration was 0.5 mmol/L (10 mg/dL). She was given an intravenous bolus of glucose followed by constant glucose infusion. Despite this regimen, hypoglycemic convulsions were observed in the ensuing 24 hours. The diagnosis of PHHI was made by demonstrating inappropriately raised levels of plasma insulin (Ins) in the absence of ketosis. Representative values were as follows: BG: 0.8mmol/L (14 mg/dL), Ins:205.2 pmol/L (28.6 μU/ml) and C-peptide: 1nmol/L (3ng/ml). The insulin/glucose ratio (Ins/G) ranged from 0.27 to 2.9. Inborn errors of metabolism were excluded by appropriate testing (organic acids profile, carnitine, NH3). The values of growth hormone (GH) during glucagon test (100 μg/kg IM) carried out at the age of 2.5 months and while she was on diazoxide and somatostatin for about 2 months were normal (Table 1). Magnetic resonance imaging (MRI) and ultrasonography (US) of the pancreas were normal.

Insulin and C-peptide were determined by RIA using commercially available reagents (Insulin: Sorin-Biomedica, intra assay variation 6.6%, inter assay variation 6.2%, C-peptide: DSL 7000, intra assay variation 5.1% and intra assay variation 4.2%) and glucose by glucose oxidase.

Her therapeutic regimen included IV glucose infusion, low protein feeds (daily protein intake of 2-2.5g/kg/d), diazoxide up to 15mg/kg/24h and somatostatin (10μg/kg/day). This combination therapy gave satisfactory glycemia most of the time. Nevertheless, a precipitous drop of BG levels to <1.1mmol/L (20mg/dl) occurred once or twice daily, especially if the IV glucose infusion was interrupted owing to technical problems. Hence, it became evident that the patient could not be maintained on medical treatment alone and surgery was considered necessary. In order to avoid blind pancreatectomy, a transdermal catheterization of pancreatic veins was performed at the age of 4.5 months (Hôpital Necker, France). Samples for Ins determination were obtained at 10 points (3 in the head, 3 in the tail of the pancreas and 1 in the portal vein, as indicated in Figure 1). Very high Ins values were obtained from a restricted area in the tail of the pancreas (Figure 1). During surgery, performed at the age of 4.5 months, foci with increased βeta cell mass were found in the tail and part of the body of the pancreas. This area was subsequently excised. Histological examination of the excised pancreas documented focal nesidioblastosis. Following pancreatectomy, the patient has been normoglycemic without any medication. The Ins/G ratio has been in the range of 0.03 to 0.1. She is now 3.5 years old and is developmentally normal. Her linear growth has improved post operatively. Specifically, her height SDS was -0.5 preoperatively and gradually reached the value of 0.8 at age 2 years (Figure 2).

**DISCUSSION**

Conservative management of PHHI consists of frequent, low protein feeds and inhibition of Ins secretion with diazoxide and/or octreotide, a long acting analog of somatostatin. Surgery is necessary for infants with hyperinsulinism who fail to be controlled by a conservative approach. Currently, most pediatric surgeons recommend subtotal (95%) pancreatectomy at the time of initial surgery since lesser resections are unlikely to control BG levels. Subtotal pancreatectomy, however, is an unphysiologic and essentially blind procedure and, although more effective than partial surgery, carries an almost 50% risk for later development of diabetes mellitus as well as exocrine pancreatic insufficiency. Preoperative localization of a possible focal lesion would allow less extensive pancreatectomy. In our patient, preoperative localization of focal insulin hypersecretion allowed excision of only a small pancreatic portion.

Focal lesions are probably not so rare in pediatric patients as has generally been considered. In one series, 36% of pancreatic specimens obtained after blind pancreatectomy showed localized hypersecretion of Ins. Hence, it is important to recognize focal pancreatic dysfunction prior to or at surgery. It is generally assumed that focal nesidioblastosis is associated with

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**Table 1. Results of the glucagon test.**

<table>
<thead>
<tr>
<th>Time (min)</th>
<th>BG (mmol/L)</th>
<th>Ins (pmol/L)</th>
<th>GH (μg/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>3.94</td>
<td>69.6</td>
<td>17</td>
</tr>
<tr>
<td>30</td>
<td>6.32</td>
<td>215.25</td>
<td>10</td>
</tr>
<tr>
<td>60</td>
<td>7.7</td>
<td>63.14</td>
<td>13</td>
</tr>
</tbody>
</table>
less severe forms of congenital hyperinsulinism. However, certain patients with focal lesions, like ours, have severe disease.

The value of imaging to differentiate diffuse from focal pancreatic lesions related to adenomas has been questioned and different noninvasive diagnostic techniques have been proposed. Standard imaging studies like computed tomography (CT), MRI, US, and angiography correctly localize insulinomas in less than 50% of adult patients. The sensitivity of CT, MRI, US and angiography in localizing insulinomas is 24%, 45%, 13% and 43%, respectively. Intraoperative US scan detects 86-90% of insulinomas in adults but is very difficult in small infants and is not expected to be helpful in focal nesidioblastosis. PVS has the advantage that it is not directly dependent on tumor size and has been recommended in adults before surgery when other tests fail to localize the tumor. It also appears as a valuable technique for the detection of focal pancreatic lesions, whether adenoma or focal nesidioblastosis, in infants with hyperinsulinism. Nevertheless, PVS has the disadvantage of being an invasive and difficult procedure which requires experience and carries the risks of irradiation to the child and the radiologist. Nowadays, intra-arterial calcium stimulation with right hepatic vein sampling for insulin gradients is considered a more sensitive preoperative test for localizing insulinomas in adults (sensitivity 88%). In children however seems to be less accurate than PVS in the evaluation of PHPH1.

A final comment should be made concerning the linear growth pattern of our patient prior to and post surgery. Contrary to expectations, body length was at the lower centiles, despite hyperinsulinemia, and moved to higher percentiles following surgery. This must be attributed to the effect of somatostatin which, however, did not seem to be mediated through GH (normal base line GH values and post glucagon). Chil-
Figure 2. Growth data of the patient with focal nesidioblastosis pre- and post pancreatectomy. The arrow indicates the age at operation.
dren in our center who take only diazoxide have normal linear growth;¹² therefore, the impaired growth prior to surgery should not be attributed to diazoxide and is possibly related to somatostatin treatment.

REFERENCES