UNIQUE PRESENTATION OF AN 8p DELETION IN A DISCORDANT TWIN WITH ATRIOVENTRICULAR CANAL DEFECT AND PROLONGED HYPOGLYCEMIA

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ABSTRACT

We report on a del(8)(p22) in a severe intrauterine growth retarded newborn with balanced atrioventricular canal defect and prolonged hyperinsulinemic hypoglycemia of infancy. Atrioventricular septal defects are associated with terminal deletions of chromosome 8p. Hyperinsulinism during infancy represents a group of clinically, genetically and morphologically heterogeneous disorders and is also associated with mutations in several genes. However, such 8p deletions are not associated with hyperinsulinemic hypoglycemia of infancy.

Key words: Del(8)(p22); Atrioventricular canal defect; Congenital hyperinsulinism; Hypoglycemia

INTRODUCTION

The congenital heart disorders associated with chromosome abnormalities are approximately 12%. Atrioventricular canal defects have extensive genetic heterogeneity [1], occurring frequently in trisomy 21 and are strongly associated with 8p syndrome [2]. A zinc-finger transcription factor GATA4 encoded at 8p22-23 is implicated as a cause of cardiac septal defects [3]. Our patient had prolonged hyperinsulinemic hypoglycemia but this region of 8p is not associated with hypoglycemia.

CASE REPORT

Discordant, dichorionic diamniotic twins were born to a 21-year-old African-American gravida 3 para 1 mother via emergency C-section at 36 weeks gestation. The pregnancy was complicated by tobacco use. The mother had a healthy 3-year-old female and one spontaneous abortion in the first trimester. The birth-weight of female twin A was 860 g (<3%), length 30.5 cm (<3%) and head circumference 25 cm (<3%). The birth-weight of female twin B was 2060 gm (3-10%), length 40.5 cm (<10%) and head circumference 30.5cm (3-10%).

The physical examination of twin A at birth revealed a triangular face, low set and posterior rotated ears, micrognathia and high arched palate. The APGAR scores at birth were 1 and 7 at 1 and 5 min., respectively. On cardiovascular examination, grade 2/6 systolic ejection murmur at the left sternal border was heard. Echocardiography showed a complete balanced atrioventricular canal defect, pulmonary stenosis (peak gradient of 63.2 mmHg), a small muscular ventricular septal defect and a secundum atrial septal defect (Figure 1). Repeated echocar-
diography showed progressive valvular pulmonary stenosis (peak gradient of 104.9 mm Hg). Twin A remained hospitalized for 4 months and during this time developed poor oral feeding, pelviectasis on the left side, hypertension which required hydralazine and methyldopa, bilateral inguinal hernia, distal renal tubular acidosis, Type 4B.

Twin A developed nonketotic hypoglycemia at 2 months of age. The blood glucose ranged between 26.0 and 40.0 mg/dL and was associated with inappropriate insulin levels (Table 1). During a glucagon stimulation test (0.5 mg intramuscular) the patient’s glucose increased from 48.0 to 82.0 mg/dL. The hypoglycemia resolved with diazoxide (5 mg/kg/ day) which was discontinued at 11 months of age without recurrence of hypoglycemia. Cyto genetic testing at the Chicago Laboratory, University of Illinois Medical Center, Chicago, IL, USA, revealed a small deletion of the terminal bands of the short arm of chromosome 8.46,XX, del (8)

Table 1. Results of critical samples during episodes of hypoglycemia

<table>
<thead>
<tr>
<th>Day of Life</th>
<th>Serum Glucose (mg/dL)</th>
<th>Serum Insulin (µIU/mL)</th>
<th>Serum Acetone (&lt;10 mg aceto-acetic acid per dL)</th>
<th>Serum Free Fatty Acids (mmol/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>67</td>
<td>39.0</td>
<td>6.0</td>
<td>Negative</td>
<td>0.40</td>
</tr>
<tr>
<td>70</td>
<td>49.0</td>
<td>4.0</td>
<td>Negative</td>
<td>0.43</td>
</tr>
<tr>
<td>76</td>
<td>48.0</td>
<td>2.0</td>
<td>Negative</td>
<td>n.a.</td>
</tr>
<tr>
<td>88</td>
<td>46.0</td>
<td>2.0</td>
<td>Negative</td>
<td>0.24</td>
</tr>
</tbody>
</table>

n.a.: not available

Figure 1. Two-dimensional echocardiography 4 chamber view showing balanced atrioventricular canal defect.
(p22).ish del(8)(p23.3 p23.3)(D8S504-). Microarray analysis at the Signature Genomics Laboratory, Washington, DC, USA, confirmed single copy loss of 673 oligonucleotide probes at the subtelomeric region of the short arm of chromosome 8. The patient had global developmental delay. Twin B was normal on physical examination and echocardiography and had no hypoglycemia. The limited family history is negative for diabetes mellitus.

The patient’s cardiovascular anomalies were attended to at 6 months of age with repair of the atrioventricular canal defect and pulmonic stenosis. The postoperative period was complicated by supraventricular tachycardia, junctional ectopic tachycardia, atrial fibrillation, functional AV blocks and multifocal atrial ectopic rhythm. The arrhythmias were well controlled with amiodarone and digoxin.

**DISCUSSION**

The first patient with an 8p deletion and congenital heart disease was reported in 1973 [4]. Such children frequently show congenital heart defects and a variable degree of mental retardation. In contrast, facial anomalies are subtle [5]. Congenital heart diseases which are phenotypically linked to chromosome 8p22-23 have a locus between D8S264 and D8S1827 [3], which is associated with GATA4, a zinc-finger transcription factor. Our patient’s deleted locus was (D8S504-). In mouse embryos, this transcription factor is necessary for normal folding that later on gives rise to heart tube and pericardial cavity [6]. Therefore, GATA4 is required for development of all four chambers. GATA4 also interacts with TBX5 for normal development of cardiac septum [3]. Thus, an 8p deletion causes haploinsufficiency of GATA4 which is responsible for congenital heart diseases.

Hyperinsulinism during infancy causes transient or permanent hypoglycemia. Prematurity, intrauterine growth restriction and congestive heart failure may result in transient neonatal hypoglycemia without hyperinsulinemia. Perinatal asphyxia can result in hyperinsulinemic hypoglycemia, but it presents during the first days of life. There are several genetic causes of hyperinsulinemic hypoglycemia including mutations in the genes encoding the SUR1/Kir6.2 complex [7], glutamate dehydrogenase, glucokinase and 3-hydroxyacyl-CoA dehydrogenase, but in many cases the genetic etiology is unknown [8]. Laboratory diagnosis of hyperinsulinemic hypoglycemia of infancy is based on persistent hypoglycemia, inappropriately elevated plasma insulin concentration during hypoglycemia, low ketones and increased glycemic response to glucagon. Hoe *et al.* [9] reported 26 cases of prolonged neonatal hyperinsulinism where genetic etiology underlying glucose regulation was unknown and 95% neonates demonstrated good response to diazoxide. Our patient had late onset prolonged hypoglycemia with good response to diazoxide. The laboratory results confirmed non ketotic hypoglycemia with glycemic response to glucagon, inappropriate insulin levels during hypoglycemic episodes and low serum free fatty acid concentration. The response to glucagon and hyperinsulinism ruled out glycogen storage defects. In our patient, chromosome 8 locus D8S504- onwards is deleted. There are no known genes that regulate insulin secretion or action within this interval. One study has reported linkage of susceptibility to type 2 diabetes on indigenous Australians to the nearby marker D8S549 [10]. It is possible that a mutation vs. deletion in such a gene could cause opposite phenotypes, hyperinsulinemia vs. increased risk for Type 2 diabetes.

**REFERENCES**


