Paternal Mutation of the Sulfonylurea Receptor (SUR1) Gene and Maternal Loss of 11p15 Imprinted Genes Lead to Persistent Hyperinsulinism in Focal Adenomatous Hyperplasia

Virginie Verkarre,* Jean-Christophe Fournet,* Pascale de Lonlay,* Marie-Sylvie Gross-Morand,* Martine Devillers,* Jacques Rahier,** Francis Brunelle,† Jean-Jacques Robert,§ Claire Nihoul-Fékété,¶ Jean-Marie Saudubray,‡ and Claudine Junien*†

*INSERM UR 383, †Department of Pathology, ‡Department of Pediatrics, §Department of Radiology, ¶Department of Surgery, Hôpital Necker-Enfants Malades, University Paris V, 75743 Paris Cedex 15, France; and **Department of Pathology, University Hospital St-Luc of Louvain, Brussels, Belgium

Abstract

Congenital hyperinsulinism, or persistent hyperinsulinemic hypoglycemia of infancy (PHHI), is a glucose metabolism disorder characterized by unregulated secretion of insulin and profound hypoglycemia. From a morphological standpoint, there are two types of histopathological lesions, a focal adenomatous hyperplasia of islet cells of the pancreas in ~30% of operated sporadic cases, and a diffuse form. In sporadic focal forms, specific losses of maternal alleles (LOH) of the imprinted chromosomal region 11p15, restricted to the hyperplastic area of the pancreas, were observed. Similar mechanisms are observed in embryonal tumors and in the Beckwith-Wiedemann syndrome (BWS), also associated with neonatal but transient hyperinsulinism. However, this region also contains the sulfonylurea receptor (SUR1) gene and the inward rectifying potassium channel subunit (KIR6.2) gene, involved in recessive familial forms of PHHI, but not known to be imprinted. Although the parental bias in loss of maternal alleles did not argue in favor of their direct involvement, the LOH may also unmask a recessive mutation leading to persistent hyperinsulinism. We now report somatic reduction to hemizygosity or homozygosity of a paternal SUR1 constitutional heterozygous mutation in four patients with a focal form of PHHI. Thus, this somatic event which leads both to β cell proliferation and to hyperinsulinism can be considered as the somatic equivalent, restricted to a microscopic focal lesion, of constitutional uniparental disomy associated with unmasking of a heterozygous parental mutation leading to a somatic recessive disorder. (J. Clin. Invest. 1998; 102:1286–1291.) Key words: K⁺/ATP channel • persistent hyperinsulinism • Beckwith-Wiedemann syndrome • neonatal hypoglycemia • loss of alleles

Introduction

Persistent hyperinsulinemic hypoglycemia of infancy (PHHI) (OMIM 256450) or congenital hyperinsulinism is a glucose metabolism disorder characterized by unregulated secretion of insulin and profound hypoglycemia. The incidence of PHHI in the general population is 1:50,000 live births (1, 2) in which 95% of the cases are sporadic. The incidence for inbred populations, where most of the familial cases are present, is 1:2,500 (3). Treatment of patients with diazoxide and/or somatostatin analogues is not always effective, necessitating an intervention such as pancreatectomy (4). Rare familial forms may be caused by recessive or dominant defects in four different genes: the high-affinity sulfonylurea receptor (SUR1) gene (5–7); the inward rectifying potassium channel subunit (KIR6.2) gene (8, 9); the glutamate dehydrogenase-1 gene (GUD1) (10); and the glucokinase gene (GK) (11). Two types of histopathological lesions are associated with PHHI: a focal form, FoPHHI, and a diffuse form, DiPHHI (12–15). FoPHHI, which represents ~30% of operated cases (16), is characterized by focal hyperplasia of islet-like cells, including hypertrophied insulin cells with giant nuclei. In DiPHHI, all the islets of Langerhans throughout the pancreas are irregular in size and contain distinctly hypertrophied insulin cells. Two forms can be distinguished by pancreatic venous sampling (17). Peroperative extemporaneous histological examination is performed to determine whether subtotal or partial pancreatectomy is required (15).

We have shown previously that focal hyperplasia of islet-like cell clusters is clonal in origin. In 10 out of 10 cases of FoPHHI examined, we detected a specific loss of maternal alleles (loss of heterozygosity; LOH) in the p15 region of chromosome 11 (18). The minimal region of LOH in FoPHHI patients is between markers D11S922 in 11pter to D11S921 in 11p15.1.

1. Abbreviations used in this paper: BWS, Beckwith-Wiedemann syndrome; DiPHHI, diffuse form of PHHI; FoPHHI, focal form of PHHI; HI, hyperinsulinemia; KIR6.2, inward rectifying potassium channel subunit; LOH, loss of heterozygosity; NBF2, nucleotide binding fold 2; PHHI, persistent hyperinsulinemic hypoglycemia of infancy; PKD, polycystic kidney disease; SUR1, high-affinity sulfonylurea receptor 1.
11p15.1 (18). This region includes two clusters of candidate genes. The first cluster corresponds to the 11p15.5 region involved in the Beckwith-Wiedemann syndrome (BWS) which contains an imprinted domain including several imprinted genes, characterized by monoaicular expression (19–22). The maternal LOH at 11p15 associated with FoPHHI involves the loss of a maternally expressed allele of either one or both of the candidate tumor suppressor genes encoding H19 (23) and p57KIP2 (24) and possibly the duplication of the paternally expressed IGF2 gene coding for the growth factor insulin–like growth factor II (25). Similar mechanisms are observed in embryonal tumors (26) and in the BWS which is also associated with neonatal but transient hyperinsulinism (27). The second cluster of genes in 11p15.1 includes the SUR1 gene (28) and the KIR6.2 gene that code for the two subunits of the β cell K<sup>+</sup><sub>ATP</sub> channel (29). Recessive mutations in these genes were identified in familial forms of PHHI (5, 8, 9, 30, 31). Although the parental bias in loss of maternal alleles in FoPHHI did not argue in favor of the direct involvement of the SUR1 and KIR6.2 genes not known to be imprinted, the somatic loss of the 11p15 region could lead to reduction of hemi- or homozygosity of a paternal mutation in either one of these genes.

To understand whether the LOH may also unmask a recessive phenotype responsible for hypersecretion of insulin we looked for deleterious mutations in the SUR1 gene. We found that four FoPHHI patients were constitutionally heterozygous for three new missense mutations in the paternal allele of the SUR1 gene when examined from leukocyte samples and samples taken from pancreas. In all four cases the mutation was also found in leukocyte DNA from their respective fathers. In the hyperplastic cells, because of the maternal allelic loss of the 11p15 region, reduction to hemizygosity or to homozygosity of the inherited paternal SUR1 mutation was observed. Therefore, these data describe the first example of a one-event two-hit mechanism, i.e., a somatic event resulting in both focal hyperplasia and limited hypersecretion of insulin due to loss of functional K<sup>+</sup><sub>ATP</sub> channel.

**Methods**

**Clinical data.** Among the 12 patients studied, 10 (Fo1 to Fo10) have been described previously (18) and 2 patients (Fo11 and Fo12) are newly reported. No familial history of hyperinsulinemic hypoglycemia was found. All patients had a neonatal PHHI onset (revealed within 72 h of birth), except patient Fo11 who was diagnosed at 2.5 mo. Both were resistant to diazoxide and underwent partial pancreatectomy. A complete relief of hyperinsulinemic hypoglycemia was obtained and the postoperative plasma glucose levels were strictly normal after a mean follow-up of 33.4 mo (21–45 mo) after surgery.

**Tissue samples.** Histological diagnosis of focal adenomatous hyperplasia was performed on extemporaneous frozen sections as described (15) and further confirmed after fixation and paraffin embedding. The focal lesion of PHHI is a limited adenomatous hyperplasia of islet cells which consists of an agglomerate of apparently normal well-defined islets surrounded by exocrine elements (ductal and acinar cells) (Fig. 1 A) and including hypertrophied endocrine cells with giant nuclei (32). Outside the lesion, the pancreatic parenchyma is morphologically normal; the islets have a normal size and have a resting appearance with normal nuclear size, no anisokaryosis, and no cellular hypertrophy (15). Pancreatic samples from normal pancreas, and from carefully identified focal adenomatous hyperplasia were frozen in liquid nitrogen and stored at −80°C. DNA from the lesion was extracted as previously described (33).

**PCR analysis for LOH.** The PCR analyses for LOH were performed as described previously (18) and the markers D11S922, D11S909, D11S1307, D11S921, D11S899, D11S1324, D11S1360, D11S897, D11S1325, and D11S912 (34) were used to genotype paired samples of DNA extracted from frozen samples of normal pancreas and hyperplastic islet cells, and leukocytes of patients and their parents.

**PCR-SSCP and sequence analysis.** PCR primer pairs used to amplify exons 33, 34, 35, 36, 37, and 38 coding the NBF2 (nucleotide binding fold 2) domain of the SUR1 gene have been described previously (5). PCR and SSCP analyses were performed using a modified protocol (35). Samples that contained aberrantly migrating fragments were directly sequenced as described (35).

**Results**

**Loss of 11p maternal alleles in focal adenomatous hyperplasia.** Using several polymorphic markers, losses of maternal alleles of the 11p15 chromosomal region were found limited to the hyperplastic lesion in 12 patients with focal adenomatous hyperplasia. 10 patients (Fo1 to Fo10) have been published previously (18) and 2 patients, Fo11 (Fig. 1 B) and Fo12 (data not shown), are newly described.

**Detection of heterozygous mutations in the SUR1 paternal allele.** Since the most commonly deleted region in focal forms also contains the genes for the two subunits of the β cell K<sup>+</sup><sub>ATP</sub> channel, the SUR1 (28) and the KIR6.2 (29) genes involved in
recessive familial forms (5, 7–9, 36–38), we looked for deleterious mutations in the SUR1 gene. Since most published mutations were located in the NBF2 domain, we studied the six exons coding for this domain (exons 33–38). As shown in Fig. 2, using SSCP analysis we detected a mobility shift by electrophoresis in constitutional DNA from leukocytes (L) samples and from normal pancreas (P) samples from 4 out of 12 patients, Fo1, Fo9, Fo11, and Fo12. In all four cases the mobility shift was also found in leukocyte DNA from their respective fathers (F). In agreement with the loss of maternal alleles, the signal of the mutant band inherited from the father was enhanced in the DNA from the hyperplastic lesion (H) while the signal of the normal wild-type band inherited from the mother was greatly diminished. The same pattern was observed for all four patients, Fo1, Fo9, Fo11, and Fo12 for exons 35, 37, 33, and 37, respectively. Persistence of a faint signal of the wild-type maternal allele in pancreatic hyperplasia (H) can be explained by contamination with stromal or exocrine cells.

new missense mutations (Table I). In all four cases, the father was also constitutionally heterozygous for the same mutation found in the proband (Fig. 3 and data not shown). 212 chromosomes from a population of 106 unrelated individuals were tested for the presence of these newly described mutations. In this population and in other individuals reported in the literature (5–8) no such missense mutations were detected, precluding a polymorphism. Furthermore, in all four cases an arginine, a basic amino acid, was replaced by a nonpolar amino acid such as proline (1353), cysteine (1421), or tryptophan (1494), increasing the likelihood that the mutation would result in a change in the protein conformation. Finally, comparison of homologous sequences of five different members of the ATP-binding cassette protein superfamily, the cystic fibrosis transmembrane conductance regulator gene (CFTR, GenBank accession No. M28668), the multidrug resistance associated protein gene (MRP1, No. L05628), the bile canalicular multispecific organic anion transporter gene (MOAT, No. U63970), the yeast cadmium transporter gene (YCF1, No. L35237), and the P glycoprotein multiple drug resistance 1 (MDR1 No. M14758), revealed that positions 1353, 1421, and 1494 are well-conserved and code for an arginine or another basic amino acid. Taken altogether these data strongly suggest that these missense mutations are deleterious. For the eight remaining patients screening of the remaining exons of the SUR1 gene and of the KIR6.2 gene is underway.

Discussion

We describe for the first time the occurrence of a recessive endocrine disorder which is due to somatic reduction to hemizygosity or homozygosity of a paternally inherited mutation limited to a focal hyperplastic lesion in four patients. In the focal lesion a loss of maternal alleles therefore resulted simultaneously in two different consequences, hyperinsulinemia (HI) and proliferation: the reduction to hemi- or homozygosity of the paternal defective allele of the SUR1 gene with loss of the wild-type maternal allele responsible for hyperinsulinism. HI is the direct consequence of loss of function of the KATP channel as already shown by patch clamp studies (39). In all four cases the father carried the mutant allele therefore excluding the hypothesis of preferential paternal de novo germline mutation. The loss of maternal alleles (18) includes the cluster of imprinted genes in the 11p15.5 region (40) comprising growth factors such as IGF2 (19) and tumor suppressor genes, such as H19 (23) and P57KIP2 (20). This necessarily leads to altered expression of these and other genes in the imprinted domain, suggesting that unbalanced expression of the corresponding gene products may give rise to the increase in proliferation of

<table>
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<th>Patients</th>
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<th>Codon</th>
<th>Amino acid substitution</th>
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<td>Arg→Pro</td>
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<td>1421</td>
<td>Arg→Cys</td>
<td>R1421C</td>
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<td>Fo9 and Fo12</td>
<td>Exon 37</td>
<td>nt 4480 C→T</td>
<td>1494</td>
<td>Arg→Try</td>
<td>R1494W</td>
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Nucleotide and codon positions are according to the full-length human SUR1 cDNA sequence incorporating the alternative splice form of exon 17 (GenBank accession No. L78208 and L78224).

Figure 2. Somatic reduction to hemi- or homozygosity of heterozygous PCR-SSCP patterns of the SUR1 gene in four FoPHHI patients. For each patient, leukocytes (L), normal pancreas sample (P), and pancreatic hyperplasia (H) were analyzed and compared with leukocyte DNA of a control (C), the father (F), and the mother (M). For patients Fo1, Fo11, Fo9, and Fo12, mutations were detected by the presence of a different electrophoretic pattern (→) present in constitutional and hyperplasia samples. The same pattern, i.e., heterozygous in lanes F, L, and P and hemi- or homozygous in lane H, was observed for all four patients, Fo1, Fo9, Fo11, and Fo12 for exons 35, 37, 33, and 37, respectively. Persistence of a faint signal of the wild-type maternal allele in pancreatic hyperplasia (H) can be explained by contamination with stromal or exocrine cells.
β cells, a striking feature of focal adenomatous hyperplasia, not observed in the diffuse form (32). This altered expression is already well-documented for embryonal tumors, and remarkably in Wilms’ tumor (26). However, the transient neonatal HI seen in 50% of BWS patients is in sharp contrast with the severe treatment-resistant HI seen in FoPHHI patients. Thus, HI in BWS, which has never been observed in numerous animal models trying to reproduce the BWS phenotype, remains unexplained (41). Understanding the role of alteration of imprinted genes in HI will require further investigations.

Thus, maternal LOH with somatic reduction to homozygosity of the paternal SUR1 mutation can be considered as the somatic equivalent, restricted to this microscopic focal lesion, of constitutional uniparental disomy associated with unmasking of a heterozygous parental mutation leading to a recessive disorder (42). There are several examples of abnormal phenotypes associated with uniparental disomy which can result both from the presence of imprinted genes on the chromosome involved in this non-Mendelian inheritance mechanism and/or from reduction to homozygosity of a mutation associated with an autosomal recessive (42).

In a recent study we report the observation and follow-up of 52 neonates (>72 h of onset) treated surgically, out of a total of 62 neonatal PHHI patients referred to the Department of Pediatrics (De Lonlay-Debeney, P., F. Poggi-Travert, J. Rahier, C. Sempoux, C.D. Vicci, J.C. Fournet, F. Brunelle, G. Touati, C. Junien, C. Nihoul-Fékété, et al., manuscript submitted for publication). In this series, the minimal proportion of focal cases (22) can be estimated at 35% (22/62). Interestingly, in three other series the percentage of focal forms or so-called “neonatal adenoma” or “focal adenomatosis” was 33%, i.e., 24 cases in a total of 72 patients (4, 12, 16, 43). Thus focal forms represent a significant proportion, one-third, of PHHI cases.

The mechanism involved in FoPHHI is similar to that recently reported for type I autosomal dominant polycystic kidney disease (ADPKD). In PKD, renal cysts are monoclonal and show allelic loss of the unaffected haplotype on chromosome 16p13.3 resulting in hemi- or homozygosity for the mutated polycystin allele (44). In PKD the somatic LOH of 16p is a surprisingly frequent event as shown by occurrence of LOH in each cell leading to cyst formation throughout the patient’s life (44). In contrast to this recurrent loss of 16p in PKD, the occurrence of LOH necessary for HI to be detectable is probably a rare event. This event must occur in an islet cell at a certain stage, late enough during development since the size of the proliferative lesion only represents a few millimeters (15). Moreover the predominantly neonatal occurrence and the rarity of relapse after partial pancreas resection in 20 cases (De Lonlay-Debeney, P., F. Poggi-Travert, J. Rahier, C. Sempoux, C.D. Vicci, J.C. Fournet, F. Brunelle, G. Touati, C. Junien, C. Nihoul-Fékété, et al., manuscript submitted for publication), after a mean follow-up of 6 yr, precludes a high rate for 1p15 LOH (17, 45). Thus, the window within which the LOH occurs is probably quite narrow during the fetal or neonatal period.

Due to methodological difficulties, the differential diagnosis of these two forms is not widely performed. While the proportion of focal cases may represent about one-third of all neonatal cases, the remaining two-thirds are diffuse cases with either homozygous mutations of SUR1 or KIR6.2 or mutations in other genes such as the glutamate dehydrogenase-1 or other (the only glucokinase mutant described is associated with a milder form) (11). Since heterozygous individuals are more numerous than homozygous individuals, this means that the probability of losing maternal alleles of region 11p15 and of developing focal PHHI is rare. But in countries with a higher prevalence of the disorder, such as Saudi Arabia, it can be assumed that these cases are not so rare and are probably underrecognized. In these countries, an apparently dominant mode of transmission due to the occurrence of a focal form in a heterozygous sibling would remain unmasked because of the high rate of consanguineous mating (> 50%). Moreover this has not been looked at in genetic studies performed on other well-described HI series. In two recent reports there was a strikingly high proportion of supposedly HI chromosomes

![Figure 3. Constitutional missense mutations in the SUR1 gene in sporadic FoPHHI patients. In patient Fo11, a heterozygous G→C point mutation substitutes an arginine residue for a proline residue at position 1421 in exon 35 (1421C). In patient Fo9 and Fo12, the sequence revealed the same heterozygous C→T point mutation which substitutes an arginine residue for a tryptophan residue at position 1494 in exon 37 (1494W). In patient Fo1, a heterozygous transition C→T substitutes an arginine residue for a cysteine at position 1421 in exon 35 (1421C). Leukocytes (L), normal pancreas sample (P), and pancreatic hyperplasia (H) were analyzed and compared with leukocyte DNA of a control (C), the father (F), and the mother (M).](image-url)


References


