Congenital hyperinsulinism and evolution to sulfonylurea-responsive diabetes later in life due to a novel homozygous p.L171F ABCC8 mutation

Short running title: CHI and later diabetes due to ABCC8 mutation

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What is already known on this topic?
Homozygous ABCC8 mutations cause severe persistent diffuse hyperinsulinemic hypoglycaemia (HH) which is usually diazoxide unresponsive and requires surgical therapy.
In medically managed patients with congenital hyperinsulinism (CHI), disease symptoms become milder overtime.
Hyperinsulinemic hypoglycemia at neonatal period and later diabetes have been reported in heterozygous mutation of HNF4A and HNF1A as well as heterozygous ABCC8 mutations.

What this study adds?
We describe the first homozygous ABCC8 mutation with HH at neonatal period and evolution to complete insulin deficient, sulphonylurea responsive diabetes mellitus.
Findings from present work which show a broad range of clinical spectrum from asymptomatic, mild symptomatic hypoglycemia, severe hypoglycemia as well as insulin deficient diabetes mellitus in family members with identical mutation confirms the phenotypical variations in ABCC8 mutations.
Present case report emphasizes the need for long-term follow up of patients with HH at neonatal period due to ABCC8 mutations, particularly those managed with medical therapy for risk of developing diabetes in later life.

Abstract:
Background: Congenital hyperinsulinism (CHI) is the most common cause of persistent hypoglycemia in infants and children. Recessive inactivating mutations in the ABCC8 and KCNJ11 genes account for approximately 50 % of all CHI cases. Hyperinsulinaemic hypoglycaemia (HH) in infancy and diabetes in later life have been reported in subjects with HNF1A, HNF4A and ABCC8 mutations.
Case report: Herein, we present a child who was diagnosed with CHI at birth, then developed diabetes mellitus at the age of 9 years due to a novel homozygous missense, p.L171F (c.511C>T) mutation in the exon 4 of ABCC8. The parents and one sibling were heterozygous carriers, whilst a younger sibling who had transient neonatal hypoglycemia was homozygous for the mutation. The mother and (maternal) uncle, who was also heterozygous for the mutation, developed diabetes within their third decade of life. The preliminary results of sulphonylurea (SU) treatment was suggestive for SU responsiveness. Patients with homozygous ABCC8 mutations can present with CHI in the newborn period, the hyperinsulinism can show variability in terms of clinical severity and age at presentation and can cause diabetes later in life. Patients with homozygous ABCC8 mutations who are managed medically should be followed as they may be at increased risk of developing diabetes.
Keywords: Congenital hyperinsulinism, MODY, ABCC8 mutation, children

Introduction
ATP-sensitive potassium (KATP) channels play an essential role in the regulation of insulin secretion from the pancreatic beta-cell; the key mechanism maintaining the blood glucose level at a narrow range of 3.5-5.5
mmol/L (1,2,3). K<sub>ATP</sub> channels are open at low glucose levels (1). Increased metabolism, resulting in an increased adenosine triphosphate (ATP)/adenosine diphosphate (ADP) ratio, leads to closure of the K<sub>ATP</sub> channel, depolarisation of the beta cell membrane and subsequent calcium influx through voltage-gated calcium channels. This in turn leads to insulin secretion via the exocytosis of secretory granules (2,3). Dysfunction of the K<sub>ATP</sub> channel can cause either congenital hyperinsulinism (CHI) or diabetes (neonatal or adult onset) (1,4,5,6,7,8,9). CHI occurs when K<sub>ATP</sub> channels are absent on the cell membrane or when they remain closed despite low glucose levels. In contrast diabetes occurs if K<sub>ATP</sub> channels remain open despite high blood glucose concentrations and increased metabolism in the beta cell (1,4). Recessive inactivating mutations of the K<sub>ATP</sub> channel genes (ABCC8 and KCNJ11) are the most common cause of severe diazoxide unresponsive diffuse CHI which usually requires pancreatectomy (1,10). Patients with dominant mutations of K<sub>ATP</sub> channel genes, ABCC8 and KCNJ11, may cause variable phenotype ranges from asymptomatic macrosomia, mild diazoxide responsive CHI to severe persistent HH as well as diabetes mellitus in later life (7,8,9,11).

CHI within the neonatal period and evolution to diabetes later in life has been reported in individuals with heterozygous inactivating mutations in the Hepatocyte nuclear factor 4A and 1A (HNF4A & HNF1A) (12,13,14) and dominant mutations in ABCC8 genes in very limited number of cases (1,7,11,13,15,16,17,18,19,20,21). To the best of our knowledge, CHI due to homozygous ABCC8 mutations and evolution to complete insulin deficient-diabetes later in life has not been reported. Herein, we present a patient with a novel homozygous ABCC8 mutation who was diagnosed with CHI at neonatal period and developed diabetes at the age of 9 years.

Case report

A 9 year-old (VI.2 in Figure 1) Turkish boy presented with abdominal pain and fever. He was diagnosed with perforated appendicitis and was referred to the endocrine clinic for coexisting hyperglycaemia (blood glucose was 27.75 mmol/L). A detailed family history revealed the presence of diabetes in multiple members of the maternal family (see details on the pedigree and footnotes). Specifically the patient’s mother was on insulin therapy to treat diabetes mellitus which had been diagnosed during the first trimester of pregnancy when she was 24 years of age and a maternal uncle was also affected. There was also a history of neonatal hypoglycaemia of various duration and severity affecting two of the patient’s siblings.

The patient was the first living child of the family and was born with a birth weight of 3750 grams (6.6 SD) via C/S at a gestation age of 29 weeks. He presented with a hypoglycaemic episode at postnatal day one (blood glucose was 1.33 mmol/L and simultaneous insulin was 22.7 µU/ml, C peptide 5.42 ng/ml (0.9-7.1). A diagnosis of hyperinsulinaemic hypoglycaemia (HH) was considered and diazoxide was commenced. The patient developed pulmonary edema, which was considered likely to be a complication of treatment with diazoxide. Diazoxide was subsequently stopped and octreotide therapy was introduced. Hypoglycaemia remitted at the age of 3 months and the child remained free of hypoglycaemic episodes until 9 years of age when he admitted to our hospital.

On admission to hospital the child was lethargic and had pale and grayish colour skin. His height was 140 cm (0.7 SDS), weight was 35 kg (0.8 SDS) and BMI 17.8 (0.7 SDS), respiratory rate was 20/minute, heart rate was 72 beats/minute, and blood pressure was 90/60 mmHg. There was abdominal distention, rigidity and rebound tenderness on physical examination. There was commenced and the patient underwent emergency appendectomy. During the post-operative period hyperglycaemia persisted and subcutaneous insulin therapy was introduced. Laboratory investigations revealed a blood glucose of 13.2 mmol/L with a simultaneous insulin of 8.82 µU/mL (2.6-25), C peptide: 1.28 ng/ml (0.9-7.1). Glycosylated haemoglobin (HbA1c) was 9.1 % (76 mmol/l). Islet cell, anti-insulin and anti-glutamic acid decarboxylase antibodies were negative. Over the following two months the insulin requirement gradually decreased until insulin treatment could be completely withdrawn. During the follow-up, HbA1c remained within a range of the high normal limits (6.2% to 6.4%) with dietary intervention and lifestyle changes. At the age of 11.7 years HbA1c was shown to be significantly raised to 9.6% (81 mmol/l). At this time his weight was 46 kg (0.7 SDS), height was 152 cm (0.8 SDS) and BMI was 19.9 (0.9 SDS). The patient and family refused recommencement of insulin therapy. Therefore, HbA1c increased to 11.4% (101 mmol/l) at 12 years when an oral glucose tolerance test suggested insulin deficient-diabetes mellitus (Table 1).

Genetic Testing

Genomic DNA was extracted from peripheral leukocytes using standard procedures and the coding regions and intron/exon boundaries of the ABCC8, KCNJ11, HNF4A and HADH genes were amplified by PCR (primers available on request). Amplicons were sequenced using the Big Dye Terminator Cycler Sequencing Kit V3.1 (Applied Biosystems, Warrington, UK) according to manufacturer's instruction and reactions were analysed on an ABI 3730 Capillary sequencer (Applied Biosystems, Warrington, UK). Sequences were compared with the reference sequences (NM_001287174.1, NM_000525.3, NM_175914.4 and NM_005327.4) using Mutation Surveyor v5.0.1 software (SoftGenetics, State College, Pennsylvania, USA). The variant was classified using the American College of Medical Genetic and Genomics (ACMG)/Association for Molecular Pathology guidelines (22). A written informed consent was obtained from the patients and/or their legal guardians.

Results
The index patient (VI.2) was found to be homozygous for a novel missense c.511C>T (p.L171F) variant in exon 4 of ABCC8 (Figure 2). The p.L171F variant affects a highly conserved amino acid and in silico analysis predicted the variant to be disease-causing (Alamut Visual V2.10 Software, Rouen, France). Mutation testing showed that the variant co-segregated with diabetes and hypoglycemia within the family with an incomplete penetrance of heterozygous carriers (see figure 1).

**Treatment and follow up**

Following detection of the ABCC8 mutation, a trial of sulphonylurea (SU) treatment was commenced in the index case and his mother who had been on insulin therapy for 13 years in an outpatient setting. The mother’s daily insulin dose requirement was reduced to about 50% from the baseline at the first week of the SU therapy with improved blood glucose measurements (BMs). The index case had also well-responded to SU therapy, and even developed one hypoglycaemic episode following SU therapy. Although the SU doses was adjusted accordingly, the family avoided giving the glibenclamide regularly due to the severe hypoglycaemic episodes which had not been observed while he was on insulin therapy or during fasting.

**Discussion**

Herein, we present a patient with a novel homozygous ABCC8 mutation who was diagnosed with HH in the neonatal period and diabetes at the age of 9 years. Hyperglycemia was first recognized during acute appendicitis which suggested stress-induced hyperglycemia. However, the patient had persistent hyperglycemia which required insulin therapy, a history of HH and relatives with autoantibody-negative diabetes. These findings were suggestive for monogenic diabetes which was confirmed by molecular genetics analysis. Homozygous Kₐ₅₉ channel gene mutations are the most common cause of severe, diazoxide unresponsive HH which often require pancreatectomy (3,13). However, clinical heterogeneity is observed in patients with dominant ABCC8 mutations (9). Kapoor et al. reported a marked clinical heterogeneity in siblings with identical mutations in ABCC8 ranging from asymptomatic hyperglycaemia to macrosomia, transient HH or severe HH and development of diabetes mellitus later in life (23). Besides, a heterogenous nature is observed in regards to severity and response to medical treatment and age for onset of the symptoms (23). Variations in the severity of HH and clinical course have also been reported in a mother and her daughter with an heterozygous E1506K mutation in ABCC8. In this report the child had severe symptoms and hypoglycaemic convulsion at 3 months while the mother had subtle symptoms of hypoglycaemia followed by gestational diabetes which persisted after delivery (20).

Similarly, a marked clinical heterogeneity was also observed in our family. While, one of homozygous siblings (VI.2, index case) had prolonged HH and required medical therapy, the other sibling with the identical homozygous mutation (VI.3) suffered from transient hyperglycemia in the first week of life which remitted within 3 months. Diabetes was observed in the heterozygous mother and a maternal uncle but the father (V.1) who was also heterozygous for the ABCC8 mutation, who was normal fasting plasma glucose and HbA1c levels. Unfortunately, family denied to perform an OGTT to the individuals who carry the mutation whilst not yet developed fasting hyperglycemia or elevated HbA1c. We also could not performed genetic analysis in other relatives who also had diabetes with (renal and ocular) complications. We, therefore, were not able to confirm whether diabetes of these additional family members is due to the ABCC8 mutation.

Indeed, the clinical course for patients with ABCC8 mutations is also substantially variable (1,7,13,15,16,17,18,19,20). Clinical features include hypoglycaemia within the neonatal period which remits overtime, this can be followed by coexistence of hyperglycaemia and post-fed hyperglycaemic episodes, impaired fasting glucose or impaired glucose tolerance in response to glucose load and in a few cases the development of diabetes (1,7,11,15,16,17,18,19,20). Regarding the type of mutation reported in ABCC8 which caused neonatal HH and diabetes later in life only few cases had a homozygous mutation (18), whilst the majority had heterozygous or compound heterozygous mutations (7,11,15,17,19,20).

Biallelic (either homozygous or compound heterozygous) ABCC8 mutations usually cause severe, diazoxide unresponsive HH which often requires surgical management (3). Therefore, the number of medically managed cases, particularly those with long-term follow-up is very few. This limits our understanding of the underlying mechanisms and experience in the management of cases who developed hyperglycemia later in life. The data from the reported cases and experimental studies suggest the key mechanisms are the dysregulated insulin secretion; impaired first phase insulin secretion, delayed insulin response and β-cell apoptosis mediated via enhanced β-cell depolarisation, resulting in increased Ca²⁺ entry into the cell (7,8,9,13,15,16,17,18,19,20,24,25,26).

Taking into account the previously reported cases, our patient is the only case with a homozygous ABCC8 mutation who presented with CHI (confirmed by clinical, biochemical evidences and mutation analysis) within the neonatal period which evolved to a complete insulin deficient diabetes later in life. Therefore, this family provides novel insights into the clinical heterogeneity of CHI and later onset diabetes in patients with homozygous ABCC8 mutations.

Neonatal diabetes due to a dominant activating mutation of a Kₐ₅₉ channel gene (KCNJ11 or ABCC8) is usually sulphonylurea responsive (27,28). We also performed a trial of SU therapy in index case and his heterozygous
mother who had insulin dependent diabetes. Preliminary results suggested a favorable SU responsiveness. Since SU drugs work by binding to the SUR1 subunit of K<sub>A<sub>ATP</sub> channel, a positive response to SU therapy suggested that the presence of a homozygous mutation may not completely abolished the channel function.

In conclusion, we present the novel missense c.511C>T (p.L171F) <i>ABCC8</i> mutation causing neonatal HH and sulphonylurea-responsive diabetes mellitus later in life. There are however some limitations in interpreting the phenotype-genotype relationships observed in this family. Firstly, we were not able to analyse the mutation status of other family members with diabetes mellitus. Secondly, although, clinical evidences and bioinformatic tools confirmed pathogenicity of the novel mutation, functional analysis have not been undertaken to assess the role of the variant <i>in vitro</i>. These results highlight the need for the long-term follow up of a larger series of CHI patients with homozygous <i>ABCC8</i> mutations who have been managed medically as well as further evaluation of these variants including functional analysis to better understand the underlying molecular mechanism and phenotype-genotype relationships.

| Table 1. Oral glucose tolerance test results of index case at age of 12 year-old |
|-----------------|-----------------|-----------------|
| Glucose(mmoll)  | Insulin(uIU/ml) | C peptide(ng/ml) |
| 0'              | 16.5            | 9.05            | 1.68           |
| 30'             | 21.8            | 10.09           | 1.94           |
| 60'             | 25.2            | 9.66            | 1.93           |
| 90'             | 28.2            | 7.76            | 1.92           |
| 120'            | 25              | 7.62            | 1.88           |
| Normal lab. values | (3.5-5.5)     | (2.6-25)        | (0.9-7.1)      |

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<th>Table 2: The results of sulphonylurea treatment trial for index case and his heterozygous carrier mother with diabetes mellitus</th>
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Figure legends
Figure 1: Pedigree of the family. The members developed either hypoglycaemia, diabetes or both are indicated as affected and shown with black-filled boxes. M: Mutated, WT: Wild type, IV.4: Insulin dependent diabetes since 35 years-old, developed diabetic nephropathy (chronic renal failure) (Reportedly), IV.5: Had insulin dependent diabetes and diabetic nephropathy (Reportedly), IV.6: Diabetes and bilateral visual loss was reported, V.1: Father, 41 years old, apparently healthy with normal glucose and HbA1c (5.6%) levels, V.2: Mother 37 years old, developed insulin dependent diabetes during pregnancy and is on insulin treatment since 24 years old, shifting the treatment to SU therapy is in progress (see the section of sulphonylurea treatment), V.3: 40 years old, had insulin dependent diabetes mellitus since 32 years-old, VI.1: Born at term, macrosomic birth weight (4750 gram; 2.8 SDS), hypoglycaemia was detected during the neonatal period, died at 1-month during hospitalization with unknown etiology, VI.2: Index patient, VI.3: 9.5 years-old, born at term, birth weight was 3750 gram (0.4 SDS), had transient hypoglycemia during the neonatal period, latest blood glucose and HbA1c levels were normal, VI.4: 6.5 years old, born at 7 months of gestational age from twin pregnancy, birth weight was 1250 gram, HbA1c is normal.
Figure 2. Electropherograms of the reference, index case and parents for c.511C>T (p.L171F) mutation.
References


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