Hyperinsulinaemic Hypoglycaemia

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Abstract

Insulin secretion from pancreatic β-cells is tightly regulated to maintain fasting blood glucose level between 3.5–5.5 mmol/l. In hyperinsulinaemic hypoglycaemia (HH) insulin secretion becomes unregulated so that insulin secretion persists despite low blood glucose levels. HH can be due to a large number of causes and recent advances in genetics have begun to provide novel insights into the molecular mechanisms of HH. Defects in key genes involved in regulating insulin secretion have been linked to HH. The most severe forms of HH are clinically observed in the newborn period whereas in adults an insulinoma is the commonest cause of HH. This review provides an overview on the molecular mechanisms leading to HH in children and adults, it describes the clinical presentation and diagnosis, and finally the treatment options for the different forms of HH are discussed.

Introduction

Hyperinsulinaemic hypoglycaemia (HH) is due to the unregulated secretion of insulin in the presence of hypoglycaemia. HH occurs in all age groups (neonates, children and adults), but due to different mechanisms. In the newborn and infancy periods it is a major cause of persistent and recurrent hypoglycaemia associated with hypoglycaemic brain injury. The biochemical basis of the HH involves dysregulated insulin secretion with defects in glucose counter regulatory hormones [1–3]. The unregulated insulin secretion drives glucose into the insulin sensitive tissues especially skeletal muscle, adipose tissue and liver causing profound hypoglycaemia. This is compounded by the fact that insulin simultaneously inhibits glycogenolysis (glycogen breakdown), gluconeogenesis (glucose production from non-carbohydrate sources), lipolysis and ketogenesis. The normal physiological glucagon and cortisol counter-regulatory hormonal response to hypoglycaemia are blunted in the newborn period further exacerbating the hypoglycaemia [2,3]. This biochemical milieu is a recipe for depriving the brain of its most important fuel namely glucose. This brain glucopaenia is accompanied by the lack of alternative substrates such as ketone bodies and lactate. It is under these conditions that the risk of brain damage is highest.

In the newborn and infancy periods HH can be either congenital or secondary to certain risk factors (such as intrauterine growth retardation) [4]. Congenital HH involves either defects in the genes ABCC8 and KCNJ11 (encoding for the 2 proteins SUR1 and Kir6.2 of the pancreatic β-cell KATP channel, respectively), or abnormalities in the enzymes glucokinase, glutamate dehydrogenase and Short Chain Acyl-CoA Dehydrogenase (SCHAD) [5–10]. Mutations in ABCC8/KCNJ11 have been reported to account for 1/3rd to 2/3rd of the causes of congenital HH in large series of patients [11,12]. Mutations in GLUD1 are the second commonest cause, responsible for approximately 5% of patients. Other known genes altogether account for less than 4% of patient with congenital HH [11,12]. Loss of function mutations in the genes ABCC8 and KCNJ11 cause the most severe forms of HH, which is usually medically unresponsive. Histologically, HH can be classified into 2 broad categories: diffuse (affecting the whole pancreas) and focal (localised to a single region of pancreas) disease [13]. Recent development, fluorine-18F 3,4-dihydroxynaloxylaline position emission tomography (¹⁸F-DOPA-PET) scan, helps to differentiate focal from diffuse disease and accurately localises the focal lesion preoperatively [14]. With the advent of¹⁸F-DOPA-PET scan and laparoscopic surgery, the clinical approach has changed dramatically.
In adults, apart from an insulinoma, HH has been reported with several conditions including insulin autoimmune syndrome, non-insulinoma pancreatic islet cell tumor, following post-gastric bypass surgery and in patients with insulin receptor mutations [15–18]. This state of the art review will describe first the molecular mechanisms regulating insulin secretion from pancreatic β-cells, then summarise the clinical presentation of HH in children and adults and finally focus on the molecular mechanisms of HH in different age groups and their management.

### Physiological Mechanisms Regulating Insulin Secretion from Pancreatic β-Cells

The main regulator of insulin secretion is the plasma glucose concentration. Insulin secretion is modified by other nutrients, circulating hormones and the autonomic nervous system, as well as local paracrine and autocrine signals. In pancreatic β-cells, mitochondrial metabolism translates glucose sensing into signals regulating insulin secretion. Under normal physiological conditions the metabolism of glucose is intricately linked to insulin secretion in pancreatic β-cells (Fig. 1) [19]. Glucose enters the β-cell and is converted to glucose-6-phosphate by the enzyme glucokinase. Glucokinase plays a unique role in acting as a glucose sensor providing a link between the extracellular glucose concentration and the metabolism of glucose in the β-cell [20]. When the blood glucose concentration is increased, the activity of glucokinase is also increased hence increasing insulin production from the β-cell. Similarly as the blood glucose concentration decreases serum insulin becomes undetectable at plasma glucose concentrations below 3 mmol/l [21, 22].

The pancreatic β-cell possesses a unique signal transduction system, which links the metabolism of the fuel stimulus to initiate insulin secretion, the so-called “stimulus-response coupling” [23]. Glucose is the most important fuel involved in the stimulus-response coupling mechanism. This stimulus response-coupling event is controlled by potassium channels (K<sub>ATP</sub>) located in the pancreatic β-cell membrane [24]. Each K<sub>ATP</sub> channel consists of a heteromultimeric complex of at least 2 proteins designated SUR1 (ABCC8 gene) and Kir 6.2 (KCNJ11 gene) [25]. The functional integrity of both of these proteins is necessary for potassium channel movement and the genes responsible for them have been localised very closely to each other on the short arm of chromosome 11 (11p14-15.1).

Under normal physiological conditions the K<sub>ATP</sub> channels maintain the electrical potential of the β-cell membrane. The metabolism of glucose in the β-cell increases the ratio of ATP/ADP, which has the effect of closing the K<sub>ATP</sub> channels. This in turn causes the opening up of voltage gated calcium channels, which regulate the entry of calcium into the β-cell. The entry of calcium is thought to be the final stimulus for insulin exocytosis [24]. Thus, the K<sub>ATP</sub> channel functions as an “on/off” switch for triggering insulin secretion.

Although K<sub>ATP</sub> channels have an essential role in linking the metabolism of glucose to the secretion of insulin, there is now evidence that there may well be other mechanisms of insulin secretion, the so-called K<sub>ATP</sub> channel independent pathways of insulin secretion [26–28]. This pathway leads to the augmented insulin release in the presence of raised cytosolic calcium (Ca<sup>2+</sup>) concentrations. Increases in the intracellular Ca<sup>2+</sup> concentration in the pancreatic β-cell cause modest increases in insulin secretion, which can be dramatically increased by modulators of protein kinases and phosphatases [29]. This suggests that steps distal to the elevation of cytosolic Ca<sup>2+</sup> are of greater quantitative importance in controlling insulin secretion [29]. It has also been shown that glucose can cause pronounced insulin secretion in Ca<sup>2+</sup> depleted islets in the presence of activators of protein kinases A and C [30].

Glucose stimulates insulin gene transcription in pancreatic β-cells by activation of the homeodomain transcription factor PDX1 (Pancreatic Duodenal Homeodomain transcription factor, PDX1) via a stress-activated pathway involving stress-activated protein kinase 2 [31, 32]. PDX1 plays an essential role in linking the cytosolic events to nuclear signalling [33]. Glucose metabolism causes phosphorylation of an inactive 31-kDa PDX1 protein localised exclusively in the cytoplasm resulting in the conver-
sion of PDX1 to an active 46-kDa form that is predominantly localised in the nucleus [32, 33]. In addition to binding to the promoter sequences of the insulin gene PDX1 also binds to the coding sequences of other genes, which are specifically expressed in the pancreatic β-cell such as GLUT2 and glucokinase [34].

Clinical Presentation of HH

Most common and severe presentation of HH is during the neonatal period. The presenting symptoms of hypoglycaemia may be very nonspecific (irritability, poor feeding and lethargy) and severe (seizures and coma). Typically these infants are macroscopic and require very high dextrose infusion rate to maintain normoglycaemia. Transient forms of HH are observed in infants born to mothers with diabetes mellitus (insulin dependent or gestational), those who have sustained perinatal asphyxia or those with intrauterine growth restriction (IUGR). In these groups of infants the HH tends to resolve after few weeks or months.

HH may be the presenting feature in some developmental syndromes such as Beckwith-Wiedemann syndrome (BWS), Soto’s syndrome and rare metabolic conditions such as congenital disorder of glycosylation (CDG). The most common syndrome associated with HH is BWS. The vast majority of HH in BWS is transient and resolves spontaneously. The other clinical features of BWS include prenatal and/or postnatal overgrowth, macroglossia, anterior abdominal wall defects, organomegaly, hemihypertrophy, ear lobe creases, helical pits, and renal tract abnormalities [35].

During infancy and childhood, the presentation of HH may be very subtle and difficult to diagnose. The presenting symptoms before 1 year of age are seizures, episodes of drowsiness or excitability. After 1 year, the symptoms are typical of hypoglycaemia; pallor, faint, tachycardia, sweating and seizures.

Patients with HH due to GLUD1 mutations (HI/HA syndrome) demonstrate 2 characteristic features. First, they show marked sensitivity to dietary protein (and leucine), and symptoms of hypoglycaemia become manifest or aggravated following a protein-rich meal, rather than a fast [36]. Second, they typically show mild to moderate asymptomatic hyperammonaemia. Hypoglycaemia is usually not as severe as that seen in HH due to $K_{\text{ATP}}$ channels defects. Children with HI/HA syndrome show good response to therapy with diazoxide and in some cases protein restricted diet. In addition, these children usually are not macroscopic at birth and their hypoglycaemia is often not recognised before several months of age.

HH may be observed postprandially, for example, in the dumping syndrome. The “dumping syndrome” is classically observed in infants following gastro-oesophageal surgery [37]. In exercise induced HH symptoms of hypoglycaemia occur within the 30 min following a short period of anaerobic exercise.

Adult onset HH is usually caused by insulinoma, presenting commonly with recurrent episodes of fasting hypoglycaemia. Rarely, functioning insulinoma can present with hypoglycaemia in the postprandial period [38]. Gastric bypass surgery is increasingly emerging as an important cause of postprandial HH in adults as a consequence of islet hyperfunction and hypertrophy [16,17]. These patients can present with neuroglycoenaenia months to years after surgery [16].

Causes of HH (Table 1)

A. HH due to defects in pancreatic β-cell $K_{\text{ATP}}$ channels

Given the key role of pancreatic β-cell $K_{\text{ATP}}$ channels in regulating insulin secretion it is no surprise that genetic defects in the genes regulating the function of these channels lead to severe forms of HH. Recessive inactivating mutations in $K_{\text{ATP}}$ channel subunits are the most common cause of HH. As mentioned earlier, $K_{\text{ATP}}$ channel is a heteromultimeric complex of at least 2 proteins designated SUR1 (ABCC8 gene) and Kir 6.2 (KCNN1 gene).

So far, over 150 mutations have been identified in the ABCC8 and 25 in KCNN1. These include missense, frame shift, nonsense, insertions/deletions, splice site and regulatory mutations, either present in homozygous or compound heterozygous state. In the Ashkenazi Jewish population, 2 common (F1388del and c.3992–9G4A) mutations account for 90% of all cases of congenital HH [6,39].

The molecular basis of recessive inactivating ABCC8 and KCNN1 mutations involves multiple defects in $K_{\text{ATP}}$ channel biogenesis and turnover, in channel trafficking from the ER and Golgi apparatus to the plasma membrane and alterations of channels in response to both nucleotide regulation and open state frequency.
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There are 2 different classes of these loss-of-function mutations. In class I, all K\textsubscript{ATP} Channels are absent from the β-cell plasma membrane, resulting in no K\textsubscript{ATP} current. The most common class I mutations are those leading to defects in trafficking. In class II mutations, K\textsubscript{ATP} Channels are present in the membrane (although less than normal) but show reduced sensitivity to Mg
 nucleotide activation or reduced intrinsic channel open probability [19].

The mutation R1437Q(23)X in exon 35 of ABCC8 is an example of class I mutation. It causes truncation of the C-terminus of SUR1, which contains the signal sequence necessary for exiting the ER. Thus, the channel protein is retained in the ER and cannot be expressed in the membrane [43]. Point mutations such as G1479R in NBD2 of SUR1 or V187D in the TMD of SUR1, which lead to reduced responsiveness to ADP activation in the expressed channels, are examples of class II mutations [19]. Overall, these mutations result in a loss-of-function of K\textsubscript{ATP} channels in the pancreatic β-cell, leading to constitutive exocytosis of insulin-containing secretory vesicles.

Recessive inactivating mutations in ABCC8 and KCNJ11 usually cause severe HH which is in the vast majority of patients is unresponsive to medical treatment with diazoxide. However, some compound heterozygote mutations may be milder and may respond to treatment with diazoxide [44].

Dominant inactivating mutations in ABCC8 and KCNJ11 causing HH have been reported with much milder phenotype than that of patients with recessive inactivating mutations [45]. They usually cause mild medically responsive HH, present later, and do not require a pancreatectomy. However, medically unresponsive dominant HH has also been reported. The molecular basis of dominant ABCC8 and KCNJ11 mutations involves impaired responsiveness to MgADP and low or nonconducting K\textsubscript{ATP} channels [46,47].

B. HH due to gain of function mutation in the GLUD1 gene

Mutations in GLUD1 gene [encodes the intramitochondrial enzyme glutamate dehydrogenase (GDH)] cause the second most common congenital form of HH (HI/HA syndrome) [10]. They are ‘activating’ mutations, leading to a gain in enzyme function by reducing the sensitivity of GDH to allosteric inhibition by the GTP and ATP [9]. GDH sensitivity to its allosteric activator, leucine is increased, which leads to the increased oxidative deamination of glutamate to α-ketoglutarate in the Krebs cycle.

GDH is widely distributed at high levels in the pancreas, liver, brain, kidney, heart and lungs. In pancreatic β-cells, the resulting increased ATP/ADP ratio consequently activates K\textsubscript{ATP} channels, with subsequent cell depolarisation and insulin release. Increased GDH activity in liver may lead to hyperammonaemia because of excessive ammonia production and impaired urea cycle activity. Recent animal model studies suggest the role of renal ammoniagenesis due to activation of GDH as a source of hyperammonaemia in these patients [48].

The GLUD1 gene mapped to the 10q23.3 region contains 13 exons encoding a 505 amino acid mature enzyme. Mutations causing HI/HA syndrome have been identified in the GTP allosteric region of the enzyme encoded by exons 11 and 12 of the GLUD1 gene, as well as in the GTP binding site encoded by exons 6 and 7 [49,50].

The major clinical feature of children with HI/HA syndrome is recurrent episodes of symptomatic HH. These may occur with fasting or can be provoked by protein feeding, as leucine is an allosteric activator of GDH. Some carriers can be asymptomatic, resulting from incomplete expression of enzyme abnormality [51].

Hyperammonaemia, a characteristic biochemical marker of HI/HA syndrome, is typically mild to moderate (up to 3–5 times the upper limit of normal). It is resistant to detoxification compounds (sodium benzoate, sodium phenylbutyrate, N-carbamylglutamate) or protein-restricted diet [51].

Children with HI/HA syndrome appear to have epilepsy more frequently as compared to other congenital forms of HH [52–54]. The increased frequency of epilepsy is thought to be either the result of a) hypoglycaemic brain injury due to recurrent hypoglycaemia or b) chronic hyperammonaemia or c) decreased concentrations of glutamine and the neurotransmitter γ-aminobutyric acid (GABA) in the brain due to raised GDH activity. Overactivity of GDH in the brain may lead to a decrease in glutamate availability for glutamate decarboxylase and GABA synthesis, which results, in turn, in altered GABA concentration [54]. However, measurements of GABA and other neurotransmitters in CSF of these patients have been normal [54]. Finally, mutations in the GTP binding site tend to be more frequently associated with epilepsy than those in the allosteric domain [53,54].

C. HH due to gain of function mutation in the GCK gene

Glucokinase (GCK) is a key regulatory enzyme in the pancreatic β-cells. It plays a crucial role in the regulation of insulin secretion and is referred to as the pancreatic β-cell sensor [20]. Its unique kinetics of low affinity for glucose (high Km) and no inhibition by its end product glucose-6-phosphate helps in modulation of its activity in relation to the concentration of glucose over a range of physiological glucose concentrations (4–15 mmol/l). Hence pancreatic β-cells are able to increase their rate of glucose metabolism in response to a rise in the extracellular glucose concentration.

Activating (or gain of function) mutations in GCK increase the affinity of GCK for glucose and alter the threshold for glucose stimulated insulin secretion [55]. Thus insulin continues to be produced at lower blood glucose concentrations. All reported activated mutations cluster in the allosteric activator site of the enzyme. There is no evidence of increased gene expression as a likely cause of HH.

GCK mutations can lead to a variable phenotype, ranging from asymptomatic hypoglycaemia to medically unresponsive HH, with majority causing mild diazoxide responsive HH [56–59]. In a large study, activating GCK mutations accounted for ~7% of medically responsive HH [60].

D. Transcription factors and HH

The hepatocyte nuclear factor 4α (HNF-4α), encoded by HNF4A, is a member of the nuclear receptor (NR) family of transcription factors. The HNF4A gene is highly expressed in the liver, kidney, gut, and pancreatic islets and is thought to play an important role in the development and function of these organs [61]. Other transcription factors important for the development of the pancreas include HNF-1α and HNF-1β [62]. These along with HNF-4α are thought to play a crucial role in expression of several genes involved in glucose stimulated insulin secretion [63].

Heterozygous mutations in HNF4A lead to a dominantly inherited condition with a dual phenotype of MODY1 and HH [64,65]. Clinically HNF4A mutation is characterised by an early neonatal presentation of HH and macrosomia. The severity may range from mild transient hypoglycaemia not needing medication to
persistent HH requiring treatment with diazoxide for up to 8 years [63]. In all reported cases, the HH improved with age and responded to diazoxide. A variable penetrance is often observed within families, demonstrated by absence of neonatal HH in some HNF4A mutation carriers. A family history of diabetes is a useful indicator of an HNF4A gene mutation, however its absence should not preclude HNF4A sequencing in patients with diazoxide responsive HH [63]. The reason for the differences in clinical presentations is not currently understood; though, it is likely that other genetic and environmental factors may influence the severity of the disease. The HNF4A mutation phenotype may extend beyond β-cells of pancreas and include liver glycoegenosis and renal Fanconi tubulopathy [66]. The exact mechanism behind HH in HNF4A mutations is unclear but may involve reduced expression of the potassium channel subunit Kir6.2, as was found in conditional knockout of HNF4A [67]. The other possible mechanism is HNF-4α deficiency leads to lower levels of PPAR α (peroxisomal proliferator-activated receptor alpha), which is important for insulin regulation [68].

A recent report described 2 cases with diazoxide responsive HH associated with mutations in HNF1A [66]. The phenotype described is very similar to HNF4A mutations – macrosomia, early neonatal presentation and diazoxide responsiveness.

E. HH and defects in the mitochondrial oxidation enzyme short chain 3 hydroxyacyl-coenzyme A dehydrogenase (SCHAD)

Short chain L-3-hydroxyacyl-CoA dehydrogenase (SCHAD) catalyses the penultimate step in fatty acid β-oxidation in the mitochondria. It is encoded by HADH, which is highly expressed in the pancreatic β-cells. Short chain 3-hydroxyacyl-CoA dehydrogenase (SCHAD) deficiency is a recently described disorder of mitochondrial fatty acid β-oxidation [9,69,70]. Unlike other inherited defects of fatty acid β-oxidation, the main clinical feature of this metabolic disease is HH. A number of studies have demonstrated that HADH has a pivotal role in regulating insulin secretion [71,72]. Recently, the mechanism behind unregulated insulin secretion in SCHAD deficiency is becoming clear. These patients were noticed to be severely protein sensitive, suggesting an amino acid triggered unknown pathway of insulin release [73]. Subsequently HADH –/- knockout mice studies demonstrated protein-protein interactions between HADH and glutamate dehydrogenase (GDH) [74]. Similar interactions have been reported in human control lymphoblast, which are lost in patients with HADH mutations [75]. Studies on isolated islets showed an increase in the affinity of GDH for its substrate α-ketoglutarate. It is therefore likely that HADH mutations cause HH by activation of GDH via loss of inhibitory regulation of GDH by HADH.

Metabolic profile in affected individuals may reveal a raised plasma hydroxybutyrylcarntine and urinary 3-hydroxybutyrate levels. However, the reason why not all patients show abnormal organic acid profiles or defects in acylcarntine metabolism is unclear. Most cases reported to date are from consanguineous families [76]. Hence sequencing of HADH is recommended in patients with diazoxide responsive congenital HH who come from consanguineous families and do not have an identifiable mutation in the ABCC8/KCNJ11 genes.

F. Exercise-induced hyperinsulinism (EIHI)

Exercise-induced hyperinsulinism (EIHI) is an autosomal dominant disorder in which strenuous physical exercise causes inappropriate insulin secretion in affected individuals, leading to hypoglycaemia. Heterozygous gain-of-function mutations in the solute carrier family 16, member 1 (SLC16A1) that encodes monocarboxylate transporter 1 (MCT1; required for transmembrane transport of pyruvate and lactate) causes exercise induced hyperinsulinism [77–79].

In normal individuals, expression of the pyruvate transporter (MCT1) is specifically silenced in pancreatic β-cells, despite nearly universal expression across other tissues [79]. Affectet patients have symptoms due to activating mutations in the SLC16A1 promoter in β-cells. Increased expression of MCT1 thus renders the plasma membrane permeable to lactate and pyruvate, allowing the latter to inappropriately stimulate insulin secretion [79]. Affected patients do not normally experience fasting hypoglycaemia. During exercise pyruvate is generated along with lactate by muscle, thereby stimulating inappropriate insulin secretion from the pancreatic β-cell despite low blood glucose levels. The mechanism highlights the importance of MCT1 absence from these cells for the normal control of insulin secretion. Although the HH is usually quite severe in these patients, specific treatment is not usually needed as hypoglycaemic episodes may be prevented by avoiding strenuous exercise.

G. Insulinoma

An insulinoma is the commonest cause of endogenous HH in adults. They are insulin secreting tumours of pancreatic origin, with an incidence of 1–4 per million [80]. Majority (90%) of them are benign, solitary, intrapancreatic and <2 cm in diameter. Classically, symptoms become evident in the fasting state or following exercise. However, it is now known that insulinoma can also present with postprandial symptoms [81]. Diagnosis was previously based on findings of abnormal serum levels of insulin, C-peptide, and more recently, proinsulin at the time of fasting hypoglycaemia [21].

Postprandial Hyperinsulinaemic Hypoglycaemia (PPHH)

PPHH refers to hypoglycaemia within a few hours of meal ingestion secondary to inappropriate insulin secretion in response to a meal.

A. Dumping syndrome

Dumping syndrome seen in infants after Nissen’s fundoplication is a classic example of PPHH [82]. Precipitous emptying of hyperosmolar carbohydrate-containing solutions into the small bowel results in rapid glucose absorption, hyperglycaemia and reactive hypoglycaemia. These children also tend to have abnormally exaggerated secretion of Glucagon Like Peptide-1 (GLP-1), which may contribute to the exaggerated insulin surge and resultant hypoglycaemia [83].

The cause of hypoglycaemia in these circumstances is usually investigated by oral glucose tolerance test (OGTT) or by a mixed-meal provocation test. Physiological dip in blood glucose level seen in OGTT might lead to misdiagnosis. However, corresponding biochemical evidence of endogenous hyperinsulinaemia and symptoms of neuroglycoapnia during a hypoglycaemic episode would help distinguish between pathological PPHH and reactive hypoglycaemia. A decrease of >6mmol/l between peak and nadir blood glucose during OGTT has been used as a diagnostic criterion for dumping syndrome [84].
B. Insulin autoimmune syndrome
Insulin autoimmune syndrome or Hirata disease is a rare condition characterised by HH associated with high titre of antibodies to endogenous insulin, in the absence of pathologic abnormalities of pancreatic islets and prior exposure to exogenous insulin [85]. The disease is extremely uncommon in Western countries. Insulin autoimmune syndrome affects men and women equally and is seen more frequently in patients older than 40 years of age. The binding kinetics of endogenous insulin by the antibodies are thought to lead to physiologically inappropriate levels of bioavailable insulin, causing either hyper- or hypoglycaemia. In this syndrome, the insulin levels are markedly elevated, usually above 100 mU/l [86]. After a meal or glucose load, these patients often demonstrate initial hyperglycaemia, followed by hypoglycaemia a few hours later. The hyperglycaemia is caused by the anti-insulin antibodies that bind the insulin secreted in response to rising blood glucose levels after a meal. This binding reduces the bioavailability of the secreted insulin to the receptors in the liver and peripheral tissues, resulting in hyperglycaemia and further insulin secretion. As the blood glucose concentrations begin to decrease and insulin secretion declines, the insulin bound to the antibodies is released, resulting in inappropriately high free insulin concentrations for the blood glucose, causing hypoglycaemia [86].

C. PPHH in patients with insulin-receptor mutations
Postprandial HH has been described in patients who carried a heterozygote mutation (Arg1174Gln) in the insulin-receptor gene [18]. Hyperinsulinism seems to be associated with decreased degradation rather than increased secretion of insulin, as evidenced by increased fasting levels of serum insulin despite normal levels of serum C-peptide and reduced clearance of exogenous insulin during clamp studies [18].

D. PPHH after gastric bypass surgery
A consequence of the obesity epidemic is the increasing use of gastric bypass surgery for patients with severe, medically complicated obesity, which has led to a number of reports of postprandial HH [16,17]. In a review of Swedish Bariatric Surgery registry, the incidence of hospitalisation for hypoglycaemia in post gastric bypass patients was reported as less than 1% [87]. A number of different explanations have been suggested to explain hypoglycaemia post gastric bypass surgery. This can either be a manifestation of dumping syndrome or improved insulin sensitivity following weight loss unmasking an underlying hyperinsulinemia syndrome. The hypoglycaemia could also be due to an effect on “enteroinsular axis” induced by the diversion of nutrients into the small intestine.

The principal reason seems to be enhanced postprandial insulin secretion, thought due primarily to increased secretion of GIP (glucose-dependent insulinotroppe polypeptide (GIP) and, especially GLP-1. GLP-1 levels are now well documented to be increased hyperinsulinaemia syndrome [85]. These findings are suggestive of the role of growth factors in islet hyperfunction seen in post gastric bypass patients.

E. Noninsulinoma pancreatogenous hypoglycaemia syndrome
Noninsulinoma pancreatogenous hypoglycaemia syndrome (NIPHS) is characterised by postprandial neuroglycopaenia [15]. Investigations reveal negative prolonged fasting tests and negative perioperative localisation studies for insulinoma. However in some patients the selective arterial calcium stimulation tests is positive with the histology of the resected pancreas showing neosidioblastosis [12]. The underlying genetic basis of NIPHS is not known. These patients are negative for ABCC8/KCNJ11 mutations and show islet hypertrophy histologically (as observed in diffuse congenital HH). The positive responses to selective arterial calcium stimulation in some patients with NIPHS, despite negative radiological localizing studies, establish that this technique should be performed in all adults with HH of unknown aetiology. Immunohistological studies of the resected pancreatic tissues have failed to show increased rate of proliferation of β-cells, or abnormal synthesis and/or processing of either proinsulin or amylin. Neither there has been any evidence of overexpression of pancreatic differentiation factors, PDX-1 and Nkx-6.1, as well as the calcium sensing receptor (CaSR) [95]. These patients usually require partial pancreatectomy to relieve further neuroglycoenaic attacks. However in some patients, diazoxide does seem to attenuate the insulin response to meals [95].

Molecular Basis of Diffuse and Focal HH

Despite identical clinical presentation, at least 2 (possibly more) well described histological types are associated with HH: a focal form and a diffuse form [13]. In diffuse form, all of the islets of Langerhans throughout the pancreas are enlarged and contain distinctly hypertrophied insulin producing cells. Focal form is characterised by nodular hyperplasia of islet-like cell clusters, including ductuloinsular complexes and giant β-cell nuclei surrounded by a histologically and functionally normal pancreatic tissue.

These 2 subtypes have different underlying genetic mechanisms. The most common causes of diffuse HH are recessive and dominant mutations in ABCC8 and KCNJ11. The focal form has a unique genetic aetiology and involves 2 independent events – the inheritance of a paternal mutation in ABCC8 or KCNJ11, and somatic loss of the maternal 11p allele (11p15.1 to 11p15.5) involving the ABCC8 and KCNJ11 region within the focal lesion [96]. The maternal 11p loss leads to paternal uniparental disomy unmasking the paternally inherited K<sub>ATP</sub> channel mutation, and leading to altered expression of a number of imprinted genes in this region, including the maternally expressed tumour suppressor.
sor genes H19 and CDKN1C, and the paternally expressed growth factor IG2 [97]. These events form the basis of unregulated insulin secretion and focal increased proliferation of β-cells evolving into a focal adenomatous hyperplasia. The focal disease is mostly sporadic in origin; however, a familial case has been reported in literature [98].

Approach to Diagnosis and Investigations (Table 2)

The management approach involves establishing a clear diagnosis of HH, undertaking appropriate genetic analysis as guided by clinical and biochemical findings, evaluating the potential for controlling hypoglycaemia by medical therapy and determining whether surgery will be required. The primary goal is to prevent neurologic symptoms and sequelae by early identification and maintenance of normoglycaemia (blood glucose levels 3.5–6 mmol/l) [1]. The diagnosis of HH is based on clinical presentation and detection of characteristic biochemical profile of hypoketonemia, hypofattyacidaemic hypoglycaemia arising from the anabolic effects of excessive insulin action at the time of hypoglycaemia. Certain clinical clues for diagnosis of HH includes macrosomia or severe IUGR, and high glucose requirement (>8 mg/kg min, normal range 4–6 mg/kg min) to maintain normoglycaemia. Characteristic metabolic profile can either be identified during spontaneous hypoglycaemia or hypoglycaemia induced by provocation tests (controlled fast/exercise/protein ingestion). However, provocation test should only be done within a controlled environment with appropriate monitoring as it can be potentially life-threatening. Laboratory findings at time of hypoglycaemia would unveil inappropriately elevated insulin and inappropriately low β-hydroxybutyrate and free fatty acids. Plasma insulin levels may not be dramatically elevated. Under normal physiological conditions, insulin production is switched off during hypoglycaemic state. As insulin release is pulsatile and has a short half-life, measurement of C-peptide (which has a longer half-life and reflects the endogenous insulin production) can prove to be more helpful when the diagnosis is in doubt. Additional supportive evidence can be provided by a positive glycaemic response to intramuscular/intravenous glucagon at the time of hypoglycaemia (a clear increment in blood glucose of >1.5 mmol/l despite severe hypoglycaemia), a positive glycaemic response to octreotide and a decreased serum levels of insulin-like growth factor-binding protein 1 (IGFBP-1) (as insulin suppresses the transcription of IGFBP-1 gene) [2,99,100]. A low serum cortisol and/or growth hormone levels at the time of hypoglycaemia is not diagnostic of cortisol or growth hormone deficiency [3]. Appropriate stimulation tests are required to confirm cortisol or growth hormone deficiency.

In the persistent rare forms of congenital HH, certain specific diagnostic findings can suggest likely underlying genetic diagnosis. Elevated serum ammonia and HH imply HI/HA syndrome (GHD-HH) [101]. However, normal ammonia concentrations do not necessarily exclude GDH-HH. In these patients with “activating” GLUD1 mutations, hypoglycaemia may occur after ingestion of leucine- or protein-rich meal and can be provoked by performing a protein/leucine provocation test [102,103]. Abnormal plasma acylcarnitine profile (elevated 3-hydroxybutyrylcarnitine) and urine organic acids (3-hydroxyglutarate in urine) characterise HADH-HH [9]. If HH is related to exercise, consider performing an exercise provocation test or a pyruvate load test [77]. An exercise test with submaximal to maximal exercise over 10 min is diagnostic.

In patients with mutations in HNF4A and UCP2, there are no specific laboratory findings. HNF4A mutations are associated with a considerable increase in birth weight, macrosomia and family history of MODY [64,65]. However, prenatal hyperinsulinism due to other genetic causes may also increase the birth weight. Family history of MODY and postprandial HH may denote GCK-HH.

In adults with HH, a supervised 72 h fast has been the classic diagnostic test as insulinoma, the commonest cause of HH in adults, would be detected in 99% cases [104]. Critical diagnostic findings are plasma insulin concentrations of at least 3 μU/ml (18 pmol/l), plasma C-peptide concentrations of at least 0.6 μg/ml (0.2 nmol/l), and plasma proinsulin concentrations of at least 5 μmol/l when the fasting plasma glucose concentrations are below 55 mg/dl (3.0 mmol/l) [21]. Measurement of sulfonylureas in plasma and urine is recommended in all patients to rule out factitious hypoglycaemia due to administration of sulfonylureas. Measurement of insulin antibodies is essential to rule out insulin autoimmune syndrome.

The patient with a history suggestive of postprandial hypoglycaemia should undergo a mixed-meal tolerance test (MMTT) [21]. A MMTT seems to be superior to an oral glucose tolerance for the evaluation of suspected postprandial hypoglycaemia. Postprandial HH, negative 72 h fasting studies, and positive selective arterial calcium stimulation test characterise noninsulinoma pancreaticogena hypoglycaemia syndrome [NIPHS] [15].

Therapies for Different Forms of HH Congenital HH (Fig. 2)

Medical management

The goal of treatment in patients with HH is to maintain normoglycaemia (blood glucose levels between 3.5–6 mmol/l). Intravenous dextrose infusion of up to 15–25 mg/kg/min through an established central venous access may be required to maintain normoglycaemia in neonates with severe forms of HH. This should be supported with enteral feeding to maintain orality. It

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### Table 2  Biochemical profile diagnostic of HH.

<table>
<thead>
<tr>
<th>Hormone/intermediary metabolite</th>
<th>Result in HH</th>
</tr>
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<tbody>
<tr>
<td>Blood samples</td>
<td></td>
</tr>
<tr>
<td>Insulin</td>
<td>Detectable or elevated</td>
</tr>
<tr>
<td>C-Peptide</td>
<td>Detectable or elevated</td>
</tr>
<tr>
<td>Free fatty acids</td>
<td>Inappropriately low</td>
</tr>
<tr>
<td>β-Hydroxybutyrate</td>
<td>Inappropriately low</td>
</tr>
<tr>
<td>Acetocetate</td>
<td>Inappropriately low</td>
</tr>
<tr>
<td>Lactic acid</td>
<td>Normal</td>
</tr>
<tr>
<td>Hydroxybutyryl carnitine</td>
<td>Elevated in HADH deficiency</td>
</tr>
<tr>
<td>Brain chain amino acids</td>
<td>Low</td>
</tr>
<tr>
<td>Ammonia</td>
<td>Elevated in HI/HA syndrome</td>
</tr>
<tr>
<td>Cortisol</td>
<td>Elevated due to hypoglycaemia</td>
</tr>
<tr>
<td>IGFBP-1</td>
<td>Low</td>
</tr>
<tr>
<td>Urine samples</td>
<td></td>
</tr>
<tr>
<td>Ketones</td>
<td>Negative</td>
</tr>
<tr>
<td>3-hydroxyglutarate</td>
<td>Elevated in HADH deficiency</td>
</tr>
</tbody>
</table>

Blood glucose < 3 mmol/l on glucose infusion rate of > 8 mg/kg/min

HI/HA syndrome: Hyperinsulinism hyperammonaemia; SCHAD: Short chain L-3-hydroxyacyl-coenzyme A dehydrogenase; AADC: aromatic L-amino acid decarboxylase; HH: Hyperinsulinaemic Hypoglycaemia; GHD-HH: growth hormone deficiency hypoglycaemia; MODY: maturity-onset diabetes of the young; NIPHS: noninsulinoma pancreaticogena hypoglycaemia syndrome; GI: gastrointestinal.
is important to note that orality is often affected possibly due to disturbed feeding pattern and gastro-oesophageal reflux, often inducing feeding refusal behaviour. Frequent monitoring of blood glucose levels is required. Once the diagnosis of HH has been confirmed, medical therapy with diazoxide should be initiated, and within the first week of diagnosis, the response to diazoxide would give an indication for the need of further evaluation and surgical intervention. In all new-borns and infants diagnosed with HH, diazoxide (KATP channel opener) is the first-line medical therapy. Diazoxide is administered orally at a starting dose of 5 mg/kg/day in 3 divided doses. The dose can be increased by 5 mg/kg/day every 48 h to an effective and tolerated dose (the maximal dose rate is 15 mg/kg/d) [105, 106]. The responsiveness to diazoxide is determined by a) appropriate fasting tolerance for age; b) feed volume and frequency normal for age; c) normal blood glucose levels at the end of the fast. The use of diazoxide is often limited by its side effects. The most common side effects of diazoxide are hypertrichosis and fluid retention (especially in the newborn), followed by hyperuricaemia, tachycardia, leukopenia, and feeding problems. In newborns, the drug is given in conjunction with the thiazide diuretic chlorothiazide (5–10 mg/kg per day in 2 divided doses), which reduces water retention. If the dose of diazoxide falls below 5 mg/kg/day, a trial off diazoxide should be considered under medical observation in the hospital setting. In those with less severe HH (HH associated with perinatal stress or IUGR), it may be preferable to start at a lower dose (2–3 mg/kg/day) of diazoxide. Diazoxide is an agonist of the KATP channel and a dose range of (5–15 mg/kg/day) is usually effective in all forms of congenital HH, except those caused by autosomal recessive mutations in the ABCC8 and KCNJ11 genes [1]. A functional KATP channel is required for diazoxide to exert an effect. Hence patients with focal or diffuse KATP HH do not respond to therapy with diazoxide. Patients with GDH-HH, SCHAD-HH, HNF4A and transient HH typically respond well to diazoxide. Patients with GCK-HH have a variable response to diazoxide and some may require surgery [56]. Patients unresponsive to maximum doses of diazoxide need urgent genetic analysis to identify those who should undergo 18F-DOPA-PET/CT in search of a focal lesion. Those with a paternally inherited ABCC8/KCNJ11 mutation are likely to have a focal lesion. While these investigations are carried out, normoglycaemia should be achieved with second-line medications. Octreotide is the second line of medical therapy for infants with diazoxide unresponsive congenital HH. Octreotide inhibits insu-
lin secretion by activation of somatostatin receptor-2 and -5 and inhibition of calcium mobilisation in β-cells [107]. Octreotide is administered subcutaneously every 6–8h, beginning at a low dose (5μg/kg/day) and titrating up to a maximum of 30μg/kg/day. Necrotizing enterocolitis is a rare but potentially life-threatening adverse effect of octreotide and therefore, it must be used with caution in neonates [108]. In most patients, there is transient hyperglycaemic response to the initial doses of octreotide. However desensitisation can occur after 2–3 doses, requiring increasing doses (tachyphylaxis) which in some patients makes this drug unsuitable for long-term use. There are reports suggesting that continuous subcutaneous octreotide infusion can overcome tachyphylaxis and lead to reduction in the dosage required as compared to when given by multiple daily injections [109,110].

Treatment with long acting preparations of octreotide has been reported to be successful in older children and needs more research to prove efficacy in younger group of patients [111,112]. In combination with frequent feeding, it may be a long-term treatment option either alone or in conjunction with diazoxide [112]. In diazoxide unresponsive patients, glucagon can be given along with octreotide, as a continuous intravenous infusion as a rescue therapy to help maintain normoglycaemia [113].

GLP-1 receptor may be a new therapeutic target in future for children with KATP HH. In a mouse model of KATP HH (SUR-1 –/–), treatment with exendin-(9–39) (GLP-1 receptor antagonist) had been shown to result in improved fasting blood glucose levels [97]. The authors’ findings suggested cAMP may have a role in KATP HH as cAMP content in SUR-1 –/– was reduced by exendin-(9–39) both basally and when stimulated by the amino acids [114]. More recently, in a randomised, open-labelled, 2-period crossover pilot clinical study involving 9 human subjects with KATP HH, it was shown that significantly higher nadir blood glucose levels were observed with exendin-(9–39) as compared to placebo. These findings propose that GLP-1 and its receptor may play a role in the regulation of fasting glycaemia in KATP HH [115].

Surgical management
The indications for surgery in HH patients include medically unresponsive diffuse disease and confirmed focal disease on 18F-DOPA-PET/CT scan. Despite the huge advances in diagnosing and accurately localising focal lesions preoperatively with novel imaging techniques such as 18F-DOPA-PET/CT there is still a potential for ambiguity. Therefore, it is very important to have an experienced surgeon, endocrinologist, as well as pathologists trained in evaluating intraoperative frozen sections to confirm the focal lesions, which aid in guiding the extent of the surgery. Infants with diffuse disease require a near-total pancreatectomy (95–98% removal) to control the HH. They might require additional therapy post-operatively with diazoxide, octreotide, and/or frequent feedings to maintain normoglycaemia. Laparoscopic pancreatectomy is a new approach to the diagnosis and management of patients with congenital HH associated with less operative trauma and faster recovery than traditional laparotomy [116,117].

Histology
Focal lesions are characterised by nodular hyperplasia of islet-like cell clusters with ductuloisular complexes and scattered giant β-cell nuclei with surrounding normal tissue [118–120]. Focal lesions are usually less than 10mm in diameter. Rarely, large focal lesions that occupy virtually the entire pancreas have been reported and 18F-DOPA-PET/CT scans may be difficult to interpret in these patients with giant lesions [121].

With the aid of the 18F-DOPA-PET/CT scan, molecular genetics, experienced pathologists, and an experienced surgeon, complete resolution of HH can be achieved with a limited pancreatectomy in focal disease [122–124]. Detailed morphology of the islets of Langerhans in diffuse HH reveals large β-cells with abnormally large nuclei [125] throughout the pancreas. Subtotal and even near-total pancreatectomy may be insufficient to relieve the hypoglycaemia in these patients with diffuse disease. Recently, a new atypical histological form of HH was characterised by morphological mosaicism [126]. In this histological form 2 types of islets coexist: large islets with cytoplasm rich β-cells and occasional enlarged nuclei and shrunken islets with β-cells exhibiting little cytoplasm and small nuclei [122]. Large islets were mostly confined to few lobules. This form has the potential of cure with partial pancreatectomy and hence pathologists are recommended to recognise this mosaicism on intraoperative frozen sections [122]. In vitro studies on islets isolated from patients with this atypical form showed elevated insulin secretion at 1 mmol/l glucose and immunohistochemistry revealed undue presence of low-Km hexokinase-I in β-cells of hyperfunctional islets [127]. This represents a novel cause of focal congenital HH.

Follow-up
Focal forms of congenital HH are completely cured after the resection of focal lesion and do not require as intensive follow-up as diffuse HH. After initial successful management of diffuse HH, these children should be closely followed up with regular 24-h glucose profile and controlled fast for ensuring optimal glycaemic control. Parents/carers should also perform rigorous blood glucose monitoring at home. Most children will become milder as they grow older. Few children will need adjustment in the medication doses with increase in weight. Over the long-term period, specific subgroups of patients such as those managed with near-total pancreatectomy are at higher risk of developing diabetes mellitus [128].

Role of genetics in management of congenital severe HH
In diazoxide unresponsive infants, mutation analysis of the KATP channel genes (ABCC8 and KCNJ11) can provide helpful information for differentiating focal from diffuse HH. In patients with homozygous or compound heterozygous mutations in KATP channel genes, the disease will be diffuse, whereas in patients carrying paternally inherited heterozygous mutations, focal disease is likely and further investigations with 18F-DOPA-PET/CT are warranted [96,129]. However, in a significant proportion, heterozygous KATP mutations may be dominant-acting and lead to diffuse disease. Additionally, approximately half of HH patients do not have mutations in known genes with a role in insulin secretion [11,12]. The interpretation of mutation results is complicated with novel variants, with whom it may be difficult to determine whether the defect is expressed recessively or dominantly. Finding a paternally derived ABCC8/KCNJ11 mutation is consistent with but does not guarantee focal HH [11]. A patient with a paternally-derived mutation may have a post-zygotic disease-causing mutation on the maternal allele that is not expressed in peripheral blood cells, resulting in diffuse HH. Finding a recessive, dis-
ease-causing $K_{ATP}$ channel mutation transmitted from the mother excludes the possibility of a focal lesion. Furthermore, as only coding regions and conserved splice sites are sequenced in Sanger sequencing, patients with no mutations identified in known genes regulating insulin secretion may have deep intrinsic mutations, which get overlooked with Sanger sequencing [130].

Role of the $^{18}$F-DOPA-PET/CT in management of congenital HH

In medically unresponsive HH with paternally inherited, de novo or no identified mutation in $K_{ATP}$ channel genes, the 2 histological subtypes, focal and diffuse, needs to be differentiated before planning surgery, as the surgical approaches for them are completely different. The diagnosis of diffuse disease is definitive in patients homozygous or compound heterozygous for $K_{ATP}$ channel genes. $^{18}$F-DOPA-PET/CT combined with contrast-enhanced CT is currently the gold standard technique for differentiating between diffuse and focal forms of congenital HH in infants [131]. However, such imaging should be performed only in centres with the necessary expertise, and the images should be interpreted only by experts in the field of combined PET and contrast-enhanced CT imaging.

The principle of this imaging technique is based on the fact that pancreatic islets take up L-DOPA and convert it into dopamine using the enzyme DOPA decarboxylase, which is expressed in islet cells. $^{18}$F-DOPA is an analogue of DOPA and thus the positron-emitting compound is useful for tracking the uptake of this dopamine precursor. Both diffuse and focal diseases have a high DOPA decarboxylase activity. Apart from differentiating diffuse and focal disease, $^{18}$F-DOPA PET/CT simultaneously permits precise preoperative localisation of the lesion [132–135]. A meta-analysis reported the pooled sensitivity and specificity of $^{18}$F-DOPA PET/CT in differentiating between focal and diffuse HH as 89% (95% CI: 81–95%) and 98% (95% CI: 89–100%), respectively [136]. The pooled accuracy in localizing focal HH was 80% (95% CI: 71–88%). If facilities are available for rapid genetic testing, then $^{18}$F-DOPA PET/CT should only be performed in those patients with paternally inherited, de novo or no identifiable $K_{ATP}$ channel mutations.

Adult onset HH

Insulinoma is the most common cause for HH in adults. Surgery is the treatment of choice for insulinoma and has a relative high success rate. Based on the extent of surgery, lifelong treatment for diabetes mellitus may be required (near total pancreatectomy). Preoperative accurate localisation is necessary for planning the surgical approach. Insulinoma can be localised using noninvasive [transabdominal Ultrasonography, Computed Tomography (CT) and/or Magnetic Resonance Imaging (MRI)] and invasive methods [Endoscopic Ultrasonography (EUS) and/or angiography and arterial stimulation venous sampling (ASVS)].

Invasive modalities are highly accurate in the preoperative localisation and have frequently been shown to be superior to noninvasive localisation techniques [137,138]. An innovative noninvasive technique, glucagon-like peptide-1 receptor (GLP-1R) scintigraphy, has been reported to successfully localise small insulinomas pre- and intraoperatively and may affect the strategy of insulinoma localisation in the future [139,140]. Medical treatment is also available but only for patients who are unable or unwilling to undergo surgical treatment, with variable response to diazoxide and octreotide. Very recently, there have been reports of improved glycaemia control with mTOR (mamalian target of rapamycin) inhibitors in patients with malignant insulinoma and refractory hypoglycaemia [141,142]. The clinical benefit is thought to be either due to the antitumor effect of mTOR inhibitor or a direct effect on glycaemic control. As functional insulin receptors are present on B-cells and mediate insulin-stimulated insulin production, mTOR inhibition downstream of insulin receptors may decrease insulin production and release [143,144].

Patients with NIPHS usually require partial pancreatectomy to relieve further neuroglycopaenic attacks. However, in some patients, diazoxide does seem to attenuate the insulin response to meals [95]. The treatment options for postprandial HH after gastric bypass surgery include modified low-carbohydrate diet, diazoxide, octreotide, α-glucosidase inhibitors (such as acarbose), calcium-channel blockers (such as nifedipine), or post-operative feeding to the bypassed proximal gut by gastrostomy [145–147]. When medical treatment options fail, surgery (pancreatic resection or reversal of gastric bypass) has been advocated due to life threatening risk of neuroglycopaenia [17]. HH due to insulin autoimmune syndrome is managed with steroids [86].

**Conclusion**

HH is a complex challenging disorder which requires early diagnosis to prevent brain injury especially in the childhood and adolescence. Despite the recent advances in understanding some of the causes of HH there are still a significant number of patients where the HH is still not established at a genetic level. Further research is required to develop novel therapies for children with the diffuse form of the disease.

**Conflict of Interest**

The authors declare that they have no conflicts of interest in the authorship or publication of this contribution.

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