Long Term Medical Treatment of Congenital Hyperinsulinemic Hypoglycemia

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Abstract

Background: Hyperinsulinism is the most common cause of recurrent Hypoglycemia in early infancy. Surgical treatment with partial/near total pancreatectomy has been the mainstay of treatment but does not result in complete remission of Hypoglycemia and is associated with high risk of diabetes mellitus.

Design and methods: We retrospectively reviewed 23 patients (1979-2009) with congenital hyperinsulinism treated medically and surgically. Patients are divided in three groups: 1) treated with diazoxide 2) treated with octreotide alone or along with diazoxide 3) treated with surgical resection. Main outcomes measured are prevention of Hypoglycemia, treatment side effects, onset of diabetes mellitus and effects on growth.

Results: Eight patients were treated with diazoxide, 10 with octreotide alone or along with diazoxide and 5 had surgical resection. Four patients came off diazoxide between 5.5-10.5 years age, 3 are less than 5 years and still on treatment. Two patients came off octreotide at 5.5 and 7 years and six remain on treatment with their ages between 6 months and 12 years. No patient could come off medical treatment after surgical resection. 2 patients had diabetes in surgical group and none in medical group.

Conclusion: Medical therapy with diazoxide, octreotide, glucagon and extensive feeding plan is an effective treatment for control of Hypoglycemia in congenital hyperinsulinism patients.

Keywords: Congenital hyperinsulinism; Neonatal Hypoglycemia; 18F-Dopa PET scan; Diazoxide; Octreotide; Near total pancreatectomy

Introduction

Neonatal Hypoglycemia is a clinical challenge for neonatologist as well as for paediatric endocrinologist. Hypoglycemia in infants and children can result in seizures, developmental delay and permanent brain damage. Neonatal Hypoglycemia aetiology ranges from simple delay in fasting adaptation to complex endocrine or metabolic diseases.

Congenital hyperinsulinism (CH) is the most common cause of transient and permanent Hypoglycemia of infancy [1]. Congenital Hyperinsulinemic Hypoglycemia is a consequence of dysregulated and inappropriate secretion of insulin due to dysfunctional K<sub>ATP</sub> channels. Prevention of hypoglycaemic episodes is the key to avoid long term neurological sequelae associated with congenital hyperinsulinism (CH) [2-4].

Inadequate suppression of insulin causes increase uptake of glucose by insulin sensitive tissues and inhibition of glycogenolysis, gluconeogenesis, lipolysis and ketogenesis [5]. All these processes make developing brain vulnerable to irreversible brain injury. The estimated incidence of CH in general population is 1/50,000 live births, but it is reported higher in parts of Finland and Saudi Arabia (1/2675 with high consanguineous marriages) [6-8].

The aim of management in infants with CH is to avoid recurrent episodes of Hypoglycemia and to maintain blood glucose level above 70 mg/dl (3.9 mmol/l). This is critical to protect brain and prevent seizures associated with Hypoglycemia. Our understanding of genetics, histology and pathogenesis is significantly improved but paediatricians are still struggling to find satisfactory treatment option.

High carbohydrate content feed is necessary to maintain normoglycaemia. Addition of raw corn starch and polyacose in feed can improve the carbohydrate content [9]. Intensive feeding plan may be achieved by nasogastric or gastrostomy tube [10].

A number of pharmacological agents are being used to achieve normoglycaemia in CH patients. There must be a rationale approach in introduction, increase in dose and change of drugs. Diazoxide is the first line drug introduced in 1965 for treatment of CH [11]. Diazoxide is given orally and it is always reasonable to prove that child is not responsive to oral therapy. Diazoxide may cause fluid retention that can precipitate cardiac failure. The most common side effect is the stimulation of generalised
hypertrichosis. The response to diazoxide can be examined by prolong fasting after 5 days; failure to tolerate fasting indicates insensitive KATP channel [10].

Second-line medical therapy for infants unresponsive to diazoxide is octreotide. Octreotide is a long-acting somatostatin analog that inhibits insulin secretion distal to the KATP channel by inducing hyperpolarization of β-cells, direct inhibition of voltage dependent calcium channels, and more distal events in the insulin secretory pathway [12]. Octreotide is administered either subcutaneously every 6-8h or via continuous infusion. Side effects include suppression of growth, steatorrhea, cholelithiasis and abdominal distension [13]. A rare life threatening effect necrotising enterocolitis is recently reported [14].

Glucagon has been used to increase blood glucose levels by stimulation of hepatic glycogenolysis. The rationale of this treatment comes from the observation of an increased glycogen content of the liver when glycogenolysis is inhibited by the inappropriately high insulin levels [15].

Surgical therapy is indicated in infants who cannot be managed medically or who may have a focal CH lesion that can be surgically cured [10]. The best available imaging technique is 18F-DOPA PET scan. This advance imaging modality can identify 75% of focal cases and precisely localise the lesion in all cases [16]. Near total pancreatectomy in patients with diffuse disease often fails to cure hypoglycemia and also lead to early diabetes mellitus [17].

Study Method and Subjects

The aim of this study is to evaluate the response and long-term outcome of pharmacological treatment in children with Hyperinsulinemic Hypoglycemia. Medical treatment may achieve normoglycaemia and complete resolution of hyperinsulinism with no long-term side effects.

A cohort of 23 children (1979-2009) with multi-ethnic background diagnosed with severe persistent Hypoglycemia in British Columbia children hospital is reviewed. Data is collected retrospectively from hospital charts. Patients are divided in 3 groups on the basis of the treatment modality: 1) Treated with diazoxide 2) Treated with octreotide only or octreotide and diazoxide together 3) Surgical resection.

Outcome measures are: effectiveness of therapy in achieving normoglycaemia, age of complete resolution, frequency of severe Hypoglycemia requiring hospital admission, and growth parameters. History of epilepsy and any focal neurological deficit is also reviewed.

Results

Twenty three patients are divided in three groups: Diazoxide sensitive (n=8), Octreotide group (n=10) and Surgical group (n=5). Patients sensitive to diazoxide presented from age of day 1 to 15 months. Fifty percent (n=4) presented with seizures. Gestational age is full term (38-42 weeks) in most of them. Birth weight ranges from 2.7-4.65 kilograms. Diazoxide treatment is commenced soon after diagnosis and supplemented with extensive feeding plan. Three patients were given Polycose in first year and 6 received corn starch after 1st year. Four patients were able to stop diazoxide after 5 years of age (5.5-10.5) with euglycaemia. Three patients still less than 5 years and one with GLUD1 mutation are continuing on diazoxide. Hypertrichosis is consistent side effect in all patients. No case of diabetes mellitus in this group yet. Three patients were admitted for Hypoglycemia; total number of admissions is four in whole group. Four patients with learning difficulties, epilepsy, ataxia and global developmental delay were diagnosed late (10 days, 6 months, 14 months and 15 months). Average height is on 67th percentile (10th-95th).

Ten patients were treated either octreotide alone or octreotide and diazoxide. Age of presentation is from day 1 to 4.5 months; 4 of 10 cases presented in first twenty four hours. Seven patients (70%) presented with seizures. Gestational age is 40-42 weeks except one who was 36 weeks. Birth weight ranges from 2.72 to 4.96 kilograms. All patients were given a trial of diazoxide as standard practice except in one whose sibling was previously diagnosed with diazoxide resistant hyperinsulinism. Diazoxide was discontinued in three patients because of side effects; raised liver enzymes, fluid retention with CCF and paralytic ileus. Octreotide is commenced either due to non-responsiveness to diazoxide or its side effects. Lypodystrophy at injection sites was noted in two patients. Two patients came off treatment at 5.5 and 7 years of age. One patient came off octreotide at 3.5 years of age but continued on diazoxide. Seven are still on treatment with their ages from 6 months to 12 years. Extensive feeding plan with polycose or corn starch via nasogastric or gastrostomy tube and continuous feeding is instituted. No major side effects are noted. Average height is on 51st percentile (10th-90th), only one patient had height of -2SDS. No case of diabetes mellitus is reported in this group yet. Eight patients were admitted due to Hypoglycemia; total number of admissions was 14 in the group. Three patients have developmental issues; they were diagnosed at 15 hours (global developmental delay), 4 months (ADHD and learning issues) and 7.5 months (ADHD and learning issues).

Five patients were treated with subtotal pancreatectomy. Gestational age in this group is from 34-41 weeks; birth weight was 3.74-4.03 kilograms. Four patients presented with seizures; age of presentation is immediately after birth, one of them stabilised with feed initially but then presented at 6 months. All patients were given diazoxide as first line treatment. One patient was given octreotide prior to surgery. Sub-total pancreatectomy decision was taken clinically due to poor response and burden of medical therapy. There was no pre-operative imaging to localise focal lesion. No patient could come off medical treatment after surgical resection. Three continued on diazoxide and two on octreotide. Three patients were admitted due to Hypoglycemia; total number of admissions was 4in the group. Two patients developed diabetes at 9 and 12.5 years. Two cases were 7.5 and 12.5 years of age at last follow up visit and they have not developed diabetes yet. Two patients had normal development and 2 developed ADHD and learning issues. One patient follow up was lost at 5 years.
Discussion

We presented 23 patients in this observational study who were diagnosed with congenital hyperinsulinism and required prolong course of treatment. We treated 18 patients conservatively and 5 had near total pancreatectomy along with medical therapy. The goal of management in CH patients is to achieve early and stable euglycaemia and a good neurodevelopment with minimal intervention and limiting the short and long term side effects of therapy. We used diazoxide as first line medical therapy and octreotide are given to those who are nonresponsive to diazoxide. Medical therapy is strongly supplemented with extensive feeding plan which include nasogastric or gastrostomy tube and continuous feed if necessary. Carbohydrate contents of feed is increased with polycose and corn starch. Families are provided home and school nursing support. Hypoglycaemia home management plan of sugar water, glucagon and octreotide is given to all carers. Patients who had only medical treatment no one developed diabetes till last follow-up. In surgical resection group (n=5) 2 developed diabetes at 9 and 12.5 years. Surgical treatment has shown promising results in focal lesions. Majority of patients require less than 50% pancreatectomy. Near total resection for diffuse lesions has not proven to be effective and not without major side effect of insulin deficiency. Leibowitiz G et al. [17] reported 14 patients with CH, 8 were treated with subtotal resection and 6 had medical treatment. Six patients developed diabetes by puberty; all of them from surgical group. Shilyansky J et al. [18] observed failure of controlling Hypoglycaemia in 25% patients undergoing near total resection and two third of them developing diabetes by the age of puberty. Andrew A et al. [10] reported more than 90% cure for focal CH, they also reported that cases with diffuse lesion and had 90-95% resection 50% of them continued to have Hypoglycemia requiring medical treatment, 25% were controlled without medical treatment and 25% developed insulin deficiency. The outcome of our surgical group is comparable with these studies as 100% continued on medical treatment after resection and 40% developed diabetes.

Neurodevelopmental outcome is difficult to measure and compare because of highly heterogeneous patients in different reported literature as well as in the same series. 26-44% patients have shown developmental delay in different studies. Menni F et al. [3] noted neonatal onset as a major risk factor for severe psychomotor retardation and epilepsy. Medical treated patients are less severely affected then those treated by surgical resection. Meissner et al. [19] described 114 patients and noted poor psychomotor outcome with infancy onset and 51% patients with mental retardation had undergone surgery. Steinkrauss et al. [20] reported 68 patients with CH; one third had developmental delay which was more prevalent in surgical group. Mazor et al. [21] studied 21 Ashkenazi children 12 with diffuse and 9 with focal disease; all of them received medical treatment. Most of the developmental problems resolved by school age and all children were enrolled in regular education. No case of diabetes was noted. In our cohort of 23 patients 13(56.5%) showed normal development among them 4 were treated with diazoxide, 7 with octreotide and 2 had surgical resection. Over all in medical group 61% (11 out of 18 patients) and in surgical group 40% (2 out of 5) showed normal development. Children with developmental delay mostly presented with onset of symptoms in infancy. 3 out of 10 children with developmental delay presented on day one and one of them initially stabilised with feed and represented at 6 months. One child presented at day 10 and others were from 4 months to 15 months old at onset of symptoms. Psychomotor delay or epilepsy in these kids may be secondary to unnoticed Hypoglycemia in neonatal period or in first couple of months.

Octreotide suppresses growth hormone secretion and may have effect on final height. A trend of transient decrease in growth reported previously although ultimate height was not affected [9]. Our patients did not show compromise in their heights as shown at last clinic visits. There is some trend of excessive weight gain seen due to extensive feeding plan.

Conservative management of CH involves a significant burden on family. It has obvious effects on social life and financial means. A good social support and where ever possible home and school nursing care are necessary. Regular blood glucose monitoring with prompt application of Hypoglycemia management plan is challenging as well as stress full for carrier. Most of these children require bolus or continuous nasogastric or gastrostomy tube feeds supplemented with polycose or corn starch. Intermittent illnesses also further complicate the situation. Medical management work load and responsibility would not be suitable for every family. Motivated and sensible parents need to be reassured that gradually this challenge would ease off as severity of disease decreases after first year of life.

In conclusion medical therapy with diazoxide, octreotide, glucagon and extensive feeding plan is an effective treatment for control of Hypoglycemia in CH patients. Neurodevelopmental outcome is good if prolong episodes of Hypoglycemia can be prevented. Surgical treatment is a good option for focal disease especially with improved imaging techniques in term of 18F-DOPA PET scan. Near total resection in diffuse lesions does not prevent continuing Hypoglycemia and freedom from medical therapy is not possible. Near total resection also bears significant risk of developing diabetes mellitus by the age of puberty.

There are limitations in our study. Data is collected retrospectively from hospital charts. We do not have definite diagnosis of focal or diffuse lesion as imaging was not performed in most of the patients. Genetic mutations are requested with research laboratories but results are not available. Neurodevelopmental outcome is measured by parent’s response and assessment at clinic visits rather than by any formal assessment.

References


