A challenging case of recurrent disabling severe hypoglycemic episodes

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Case history

- 33 year old landscape gardening labourer
- Insulin treated diabetes diagnosed at 3 weeks of age
- Referred to us at the age of 31 with 6-7 months of worsening hypoglycaemia with reduced / loss of warning symptoms
- No family history of Type 1 or Type 2 diabetes

Collateral history from mother

- Born at term by Forceps delivery
- Intubated for 3 minutes after birth
- Found soon after collapsed, 'looked blue' required re-intubation
- Persisting hypoglycaemia requiring IV dextrose, hydrocortisone and glucagon
- Continued to become hypoglycaemic and 'started fitting'
- Further management:
  IV Diazoxide, Paraldehyde, Phenobarbitone, Diazepam

Surgical review during 1st week of life

- 5 days old:
  * Partial pancreatectomy
- 3 weeks old:
  * Subtotal pancreatectomy
  * Insulin started (sliding scale initially)

Histology – Nesidioblastosis

- Irregular shaped hypertrophied islets
- Islet cell pleomorphism. Marked difference in size of cells within same islet

Clinic visit - 2010

- 6-7 months of worsening hypoglycaemia with reduced / loss of warning symptoms
- 4-5 episodes of severe hypoglycaemia needing to go to local A&E
- Medications:
  Novorapid 12 units with breakfast, 20 units with lunch and dinner
  Detemir 36 units mane, 38 units nocte
  Creon ?dose 8-9/day with meals

Clinical findings

- Weight 79.5kg, Height 1.78m (BMI 25.1)
- CVS/RS/CNS – no abnormality detected
- Abdomen – small midline scar, no organomegaly

Investigations

- HbA1c 6.6%, 49 mmol/mol
- C-peptide detectable at 0.12 ug/L (NR 1.1-4.4)
- Insulin <0.5 mIU/L (NR 2.6-24.9)
- Sulphonylurea screen negative
- Normal renal, liver and thyroid profile
- MRI and CT pancreas - possible small area of residual pancreatic tissue in the region of head of pancreas
- Octreotide scan - no evidence of increased uptake in this area

Discussion

- Nesidioblastosis was first identified in 1938, when Laidlaw coined this term to describe the neodifferentiation of islets of Langerhans from pancreatic ductal epithelium (a term since replaced by PHHI and CHI)
- Persistent hyperinsulinemic hypoglycaemia of infancy (PHHI) represents the most common cause of hyperinsulinism in neonates. Currently, many authors prefer the term congenital hyperinsulinism (CHI)
- Severe recurrent hypoglycaemia associated with an inappropriate elevation of serum insulin, C-peptide, and proinsulin levels defines Congenital hyperinsulinism
- If left untreated, this condition can lead to brain damage or death secondary to severe hypoglycaemia
- CHI is often poorly responsive or unresponsive to medical management, necessitating 95% or near-total pancreatectomy

Histological features of Nesidioblastosis

Questions for audience

This has been a challenging case of hypoglycemia.

Nesidioblastosis is a well recognized cause of persistent hyperinsulinaemic hypoglycaemia of infancy. Although it was initially thought to affect only infants and children, numerous cases have been reported in adults of all ages but at a much lower incidence. Regardless of the pancreatectomy procedure used, hypoglycaemia may recur during long term follow up

- Should total pancreatectomy be considered for at this stage?
- How do we control his hypoglycaemia if this gets worse in the future?