Lesson of the month: Recurrent falls and hypoglycaemia in an older woman

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We present a case of a non-diabetic 84-year-old woman who presented with recurrent falls and hypoglycaemic episodes and was subsequently diagnosed to have a solitary insulinoma. Her multiple fall episodes may have been precipitated by her low blood glucose levels. We discuss the clinical and radiological findings of insulinoma.

KEYWORDS: falls, hypoglycaemia, insulinoma

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Case presentation

An 84-year-old woman was admitted to hospital in view of a near fall after a recent admission for small bowel obstruction secondary to right hernia, for which a diagnostic laparoscopy, mini laparotomy, adhesiolysis and an open right inguinal hernia mesh repair was done. Prior to the fall, patient did not report any symptoms of giddiness, headache, blurring of vision, focal numbness or weakness, chest pain or shortness of breath. During the fall, there was no loss of consciousness or abnormal jerking movement of limbs. The near fall episode was attributed to various degenerative bone and joint diseases.

She was subsequently transferred to a community hospital for further rehabilitation. During her stay, she was found to have recurrent hypoglycaemia which typically occurred in the morning. During her episodes of hypoglycaemia, she remained asymptomatic. Her renal function, liver panel, infection markers, thyroid function and cortisol level were normal, as was a drug screening test. Her serum C-peptide was 1.20 (reference range 0.78–5.19 ug/L), serum insulin was 0.26 (reference range 1.0–30.0 mU/L) and fasting plasma glucose was 2.3 (reference range 3.9–6.0 mmol/L). Previous CT abdomen pelvis (Fig 1) done for intestinal obstruction was found to have a dense focus in the periampullary area which was either duodenal or pancreatic in origin. The lesion was further characterised by an MRI pancreas (Fig 2), which showed the presence of a pancreatic head lesion likely due to a calcified insulinoma. Subsequently, a PET DOTATE scan (Fig 3) was performed with findings in keeping with a tracer-avid lesion in the pancreatic head, which may represent a neuroendocrine tumour. Endoscopic ultrasound-guided fine needle aspiration cytology was done and confirmed the diagnosis of neuroendocrine tumour. The patient was treated with diazoxide 100 mg BD.

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Fig 1. CT abdomen pelvis. Dense focus in periampullary region which was either duodenal or pancreatic in origin.

Fig 2. MRI pancreas. This shows a pancreatic head lesion which may represent a calcified insulinoma.
Fig 3. PET DOTATE scan. In the pancreatic head, adjacent to the distal common bile duct, there is a focal increase in tracer uptake, likely corresponding to known calcification demonstrated in prior MR/CT study. SUVmax 23.3. The findings are suggestive of an insulinoma.

Discussion

Insulinomas may have varied presentations in the elderly population. In the above case example, one of the possible causes of her near fall event could be attributed to her hypoglycaemic episodes. Insulinoma has an annual incidence of four in 1 million persons and is considered the most common neuroendocrine tumour of the pancreas. Hypoglycaemia is a common presenting feature in insulinoma patients. In the absence of diabetes mellitus, the annual incidence of insulinoma is 0.7 per million persons and is considered the most common neuroendocrine tumour of the pancreas. In terms of the pathogenesis of insulinomas, they can occur sporadically or in conjunction with MEN-1 syndrome. Non-diabetic patients with documented Whipple’s triad or severely depressed serum glucose level (<2.2 mmol/L) require further evaluation, with differential diagnosis including insulinoma, drugs, critical illnesses, hormonal deficiency or paraneoplastic syndrome. The gold standard of biochemical diagnosis of insulinoma is to measure insulin, C-peptide, plasma glucose and proinsulin during a 72-hour fast. Studies have reported that helical computed tomography and endoscopic ultrasound can provide higher sensitivity (94% sensitivity in both). A DOTATATE PET/CT can also identify most insulinomas and can be used as an adjunct imaging study when other imaging studies are non-conclusive.

Hypoglycaemia unawareness happens when recurrent hypoglycaemia shifts the glycaemic threshold for counter-regulation, as in this case. Florid neurogenic symptoms are therefore blunted, while non-specific neuroglycopenia signs and symptoms may predominate. Despite being reviewed by a dietician as an inpatient, the patient continued having hypoglycaemic events. A diagnosis of insulinoma was considered after thorough work up for hypoglycaemia. Localisation via radiological studies and tissue diagnosis was pursued. The median duration of symptoms before diagnosis of insulinoma has been shown in one study to be 18 months. A delayed diagnosis and treatment may be potentially dangerous for patients, whereas early diagnosis can prevent falls, syncope, cognitive impairment and improve quality of life for geriatric patients.

Insulinoma can be treated medically or surgically. The definitive treatment for benign insulinomas is surgery, although other techniques, such as injection of octreotide, endoscopic-ultrasound-guided alcohol ablation, radiofrequency ablation (RFA) and embolisation of an insulinoma of the pancreas, have been described. Our patient opted for medical treatment after discussing the pros and cons of the treatment options. No further hypoglycaemic episodes occurred afterwards.

Conclusion

We present a case of a non-diabetic elderly woman who presented with recurrent falls and hypoglycaemic episodes who was subsequently diagnosed to have a solitary insulinoma. Her multiple fall episodes may have been precipitated by her low blood glucose levels.

Patients and clinicians should engage in shared decision making on whether to pursue surgical or medical management.

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