Case report

Unusually rapid $\beta$-cell failure in a patient newly diagnosed with type 2 diabetes presenting acutely with unprovoked severe hyperglycaemic hyperosmolar state: a case report

Joey Yeoh$^1$, Judy Chien-Chun Huang$^2$, Harriet Cheng$^2$ and Kenneth Ross Muir$^1$

Abstract

Pancreatic $\beta$-cell failure on a background of insulin resistance results in the inability to compensate for fasting hyperglycaemia and eventually produces type 2 diabetes mellitus. We describe an interesting case of a patient who presented acutely with unprovoked severe hyperglycaemic hyperosmolar state and was subsequently diagnosed with type 2 diabetes mellitus on a background of only impaired first phase insulin secretion 4 months prior. Glucagon stimulation test detected significant $\beta$-cell failure necessitating long term exogenous insulin therapy which is highly unusual by virtue of the rapid apparent deterioration.

Introduction

Type 2 Diabetes Mellitus (T2DM) is undoubtedly the scourge of the developed world and a source of considerable socioeconomic burden [1]. It accounts for 90-95% of diabetes cases worldwide [2]. Although the exact aetiology of T2DM remains unknown, it involves a combination of insulin resistance and some degree of pre-existing $\beta$-cell secretory dysfunction conferring a state of relative rather than absolute insulin deficiency which progressively deteriorates with time irrespective of treatment [2-5]. Individuals at risk of developing overt T2DM, including the pre-diabetic states of impaired glucose tolerance (IGT) and impaired fasting glycaemia (IFG) already exhibit $\beta$-cell dysfunction and the weight of evidence suggests that this occurred long before the onset of pre-diabetes, when normal fasting glucose was still present [5,6].

Pancreatic $\beta$-cell compensation initially keeps glycaemia near normal despite underlying insulin resistance by increasing insulin secretion during the normoglycaemic and pre-diabetes stages [3-5]. The failure of $\beta$-cells to compensate at some point leads to development of overt T2DM [5,6]. This $\beta$-cell failure develops in a progressive fashion and continues after diagnosis, frequently resulting in secondary failure and exogenous insulin requirement [5,6]. It has not been reported <6 months from detection.
of impaired glucose metabolism (IGM) [7]. We report a case where the duration between onset of initial abnormality in glucose homeostasis and apparent β-cell failure necessitating long term exogenous insulin therapy is unusually rapid.

**Case presentation**

A 73-year-old Caucasian male presented to the Emergency Department (ED) in June 2008 with a history of collapse without loss of consciousness while mobilizing to the toilet in the early hours of the morning. This occurred on the background of a 6 day history of general unwellness. He was conscious and orientated on arrival but was slightly drowsy. Other than clinical features of dehydration and mild central abdominal adiposity with a waist: hip ratio of 1.04, there were no other significant clinical signs. His random plasma glucose (RPG) on arrival was 59.1 mmol/L. His height was 1.76 meters (m) and weight was 90 kilograms (kg) making the body mass index (BMI) 29.0 kg/m². The patient was previously fit and well without any known co-morbidities. He was on no regular medication. There was no family history of diabetes or other significant conditions. He was a non-smoker and consumed no alcohol. He works as a farm equipment evaluator. There was no history of specific weight loss over the preceding months. His general practitioner (GP) had performed a fasting plasma glucose test on him in February 2008 which showed a value of 6.4 mmol/L. This was repeated within 2 weeks and the repeated value was 6.1 mmol/L. Glycosylated haemoglobin (HbA1c) was also done and was 6.3%. Subsequent 75 gram (g) oral glucose tolerance test (OGTT) produced a value of 15.3 mmol/L, 14.1 mmol/L and 7.6 mmol/L 30 minutes, 1-hour and 2-hours post glucose load respectively. He was managed with dietary modification and exercise. His sodium (Na) was 150 mmol/L, potassium (K) 4.6 mmol/L, urea 26.1 mmol/L and creatinine (Cr) 237 micromol/L. Calculated osmolality was 394.4 mOsm/kg. Arterial blood gas sampling (ABG) showed a pH of 7.36, pCO₂ 4.9 kPa, pO₂ 11.0 kPa, bicarbonate 20 mmol/L and BE -1 mmol/L. Lactate was mildly elevated at 1.4 mmol/L. pO₂ 11.0 kPa, bicarbonate 20 mmol/L and BE -1 mmol/L.

Patient had been fasted for 15 hours overnight and medications were withheld until after the test. The glucose values were 6.3 mmol/L, 15.1 mmol/L, 16.3 mmol/L and 16.6 mmol/L at baseline, 5, 10 and 15 minutes respectively. Corresponding C-peptide values were 0.21 nmol/L, 0.42 nmol/L, 0.47 nmol/L and 0.51 nmol/L respectively. He was finally discharged after 7 days of hospital care with clinic follow up in 2 months. His average capillary blood glucose (CBG) was 6.7 mmol/L and range 6.5-7.7 mmol/L on discharge.

**Discussion**

The OGTT results prior to admission provided surrogate information on first phase insulin secretion with the relative inability to clear glucose 30 minutes post load being suggestive of first phase insulin secretory defect characteristic of IGM [8]. At that stage, the patient could be classified as having mixed IFG/IGT but did not meet criteria for diagnosis of T2DM [2]. Subsequent admission with overt, apparently uncompromised HHS and full blown T2DM requiring insulin to achieve glycaemic control within 4 months of identification of IGM in a previously treatment naïve individual is highly unusual [6]. In addition, the IV glucagon stimulation test demonstrated the inability of β-cells to respond to hyperglycaemia which not only confirmed established T2DM, but also significant β-cell failure [3-5].

Homeostasis model assessment-insulin resistance (HOMA-IR), reciprocal HOMA-IR and quantitative insulin sensitivity check index (QUICKI) were not done to confirm insulin resistance or state of insulin sensitivity since these calculations require intact β-cell function (fasting insulin and glucose levels) [9]. Other complex tests and calculations for insulin sensitivity/insulin resistance or for assessment of β-cell function were not performed as they were deemed to be more appropriate in controlled research settings and beyond the ethical scope of a case report.

The patient was finally discharged after 7 days of hospital care with clinic follow up in 2 months. His average capillary blood glucose (CBG) was 6.7 mmol/L and range 6.5-7.7 mmol/L on discharge.
possible cause(s) of progressive β-cell dysfunction and failure are a matter of intense interest due to the obvious potential for therapy. The β-cell dysfunction and eventual failure involves β-cell secretory defects (manifesting as loss offirst and then second phases of insulin secretion, reduced diurnal oscillations and impaired rapid pulsatile secretion) and reduction of β-cell mass [3-6]. Research has revealed several potential causes including glucotoxicity, lipotoxicity, proinflammatory cytokines secreted by adipose tissue and immune system and islet amyloid deposition [3-6]. Recent research even provided a possible cellular level common pathway by which β-cell failure can occur, involving the unfolded protein response (UPR) signalling pathways in the endoplasmic reticulum (ER) [10]. Regardless, it is fair to say that we do not yet know the exact cause(s) of β-cell dysfunction and failure. With regards to the possible factor(s) we do know about, there is no conclusive data about the actual duration needed to cause significant β-cell dysfunction and hence onset of T2DM, although the process is thought to occur over years rather than months [3-6]. In this patient, the rapid apparent deterioration of β-cell function cannot be adequately explained by lipotoxicity or glucotoxicity since there was no hyperlipidaemia or significantly raised plasma glucose previously. The possibility that acute glucotoxicity during the HHS affected the IV glucagon stimulation test result is unlikely since the test was done 5 days later, when the patient already had good glycaemic control. Even if the above mentioned causes existed in this patient, the rapid progression profile makes it unlikely that they were of any major importance.

**Conclusion**

Although progression from IGM to T2DM is an expected course in the spectrum of the disease, rapid β-cell failure and development of insulin requiring T2DM within a few months of IGM is unusual. Therefore, this case suggests the pervasive likelihood of other, yet undetected factor(s), which may cause uncharacteristically rapid β-cell failure and rapid progression from normal glycaemia to IGM and overt insulin requiring T2DM.

**Abbreviations**

ABG, arterial blood gas; Anti-GAD 65, anti-glutamic acid decarboxylase 65; BID, Bis in Die; BMI, body mass index; ER, endoplasmic reticulum; HbA1c, glycosylated haemoglobin; HHS, hyperglycaemic hyperosmolar state; HOMA-IR, homeostasis model assessment - insulin resistance; IA-2, insulinoma antigen-2; ICA, islet cell antibody; IFG, impaired fasting glycaemia; IGM, impaired glucose metabolism; IGT, impaired glucose tolerance; OGTT, oral glucose tolerance test; QUICKI, quantitative insulin sensitivity check index; RPG, random plasma glucose; T2DM, Type 2 Diabetes Mellitus; UPR, unfolded protein response; WCC, white cell count.

**Consent**

Written informed consent was obtained from the patient for publication of this case report. A copy of the written consent is available for review by the Editor-in-Chief of this journal.

**Competing interests**

The authors declare that they have no competing interests.

**Authors’ contributions**

JCCH collected the details of the case and was responsible for compiling the case presentation. HC and KRM were responsible for the literature search and compiling background information. JY coordinated the entire process and was responsible for the literature review and preparation of the first draft of the manuscript. JY and KRM managed the patient during in-patient hospital stay. All the authors read and approved the final manuscript.

**References**