Testing for Autoimmunity and β-Cell Function in a Young Patient with Diabetes Mellitus

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Abstract

This is a case of a 22-year-old Filipino male, morbidly obese, not known to have diabetes mellitus (DM) who presented with diabetic ketoacidosis on initial diagnosis. He had a phenotype of type 2 DM (T2D) but an initial presentation consistent with type 1 DM (T1D). Insulin therapy was eventually discontinued but he maintained good glycemic control with diet alone. C-peptide showed adequate increase after a mixed-meal diet and GAD65Ab was negative, and he was diagnosed with ketothesis-prone DM (KPD). The increasing prevalence of obesity challenges the classic phenotype of patients with DM, with many patients presenting as an obese type T1D, and being diagnosed with T2D at a younger age. This complicates how to classify the patient's diabetes, and the clinical profile is sometimes insufficient to make the proper diagnosis. In these cases, immunologic markers and assessment of β-cell function are important tools to differentiate between T1D and T2D, to direct management plans and to anticipate complications.

Key words: ketothesis-prone diabetes mellitus, autoimmune, c-peptide

INTRODUCTION

Diabetes mellitus (DM) is classified into 4 major categories: Type 1 Diabetes Mellitus (T1D), Type 2 Diabetes Mellitus (T2D), Gestational Diabetes Mellitus (GDM), other types due to other specific causes that reflect its underlying pathophysiology. Immunologic markers and assessment of β-cell function are useful tools when clinical parameters alone are inadequate to differentiate between T1D and T2D.

There are five antibodies [Islet Cell Antibody (ICA), Insulin Autoantibody (IAA), Glutamic Acid Decarboxylase 65 Antibody (GAD65Ab), antibodies to Tyrosine Phosphatase like IA-2 (IA-2Ab) and Zinc Transporter 8 (ZnT8)] that are used to test for autoimmunity in DM. GAD65Ab is detected in 70-80% of cases during the evolution of a patient with T1D. It is likewise the most sensitive marker (91%) for detecting multiple antibody positivity compared with ICA (82%).

β-cell functional reserve can be assessed with a Mixed-Meal Tolerance Test (MMTT) or Glucagon Stimulation Test (GST) by measuring fasting and post stimulation C-peptide levels. In a recent study, basal C-peptide of less than 0.6 ng/ml indicates absolute insulin deficiency; more than 0.6 ng/ml suggests T2D or maturity onset diabetes in the young (MODY) in a patient with presumed T1D, and more than 3 ng/ml suggests MODY or T2D in young onset DM at diagnosis.

The classification of DM is often difficult based on clinical parameters alone. The face of diabetes has evolved over the years with the increasing prevalence of obesity. Younger patients are frequently diagnosed with T2D and obesity in T1D is becoming more common. In these equivocal circumstances, immunologic markers and assessment of β-cell function are important tools to differentiate between T1D and T2D but is underutilized in routine clinical practice.

This gray area between the types of DM is discussed and how accurate diagnosis leads to proper management.

CASE

The patient is a 22-year-old Filipino male who was admitted due to diabetic ketoacidosis (DKA). He did not have a prior history of DM but experienced polydipsia, polyuria and 8 kg weight loss one month prior to admission. He eventually developed generalized weakness and fatigue.

The patient has an unremarkable past medical history. Family history includes a diabetic mother on metformin, and a hypertensive grandmother. There was no autoimmune disease in the family.

On examination, the patient’s blood pressure was 140/90 mm Hg, RR 20 breaths per minute, heart rate of 79 per minute, and temperature of 37 °C. He is morbidly obese (BM 41.1; WH 1.29) with acanthosis nigricans. Initial work-up showed blood glucose of 462 mg/dl, arterial

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blood gas was uncompensated metabolic acidosis and +3 urinary ketones. There were no infections noted. He was admitted as a case of DKA and was managed accordingly with fluid and an insulin drip. Other work-ups showed absence of microvascular complications.

Patient was discharged with NPH insulin 40u SC in AM and 20u SC in PM, metformin 500 mg/tab BID and amlopidine 5 mg/tab QD. One month after discharge, patient was noncompliant with insulin but noted to have good glycemic control with metformin alone (preprandial CBGs 103-133 mg/dl). He was also noncompliant with his antihypertensive medication but BP monitoring at home was stable at 120/80 mm Hg. We requested for C-peptide and GAD65Ab two weeks after discharge to determine whether the patient has T1D or T2D. GAD65Ab was negative and C-peptide was detectable at fasting state and increased after a mixed-meal diet. Patient has no pancreatic islet antibodies and has adequate β-cell functional reserve, and was diagnosed as unprovoked Aβ Ketosis Prone Type 2 DM. Metformin was eventually discontinued after 6 months. One year after the diagnosis, he was able to achieve good glucose control with HbA1c of 6.2%, FBS 4.3 mmol/L with diet alone.

The biochemical profile of the patient is shown in Table 1.

| Biochemical Profile | Patient |
|---------------------|---------|
| ImmunoLogic Parameter | <2.0 U/l (-) |
| GAD65Ab (NV: <30 U/l Negative; ≥30 U/l Positive) | |
| β-Cell Function | 4.93 ng/ml |
| C-peptide (NV: 1.10-4.40 ng/ml) | 15.87 ng/ml |
| Fasting | |
| Post-Prandial (mixed meal) | |
| Other Metabolic Profiles (on Admission) | |
| FBS | 174.79 mg/dl |
| HbA1c | 12.2 % |
| Total Cholesterol | 168.17 mg/dl |
| Triglycerides | 100.89 mg/dl |
| HDL | 35.18 mg/dl |
| LDL | 119.07 mg/dl |

**DISCUSSION**

The increasing incidence of ketoacidosis without precipitating cause has been reported among pediatric and adult patients with T2D, primarily among Africans, African Americans and other minority ethnic groups. These patients are usually obese and have a strong family history of DM. They are able to regain sufficient β-cell function and insulin sensitivity within months of treatment allowing the discontinuation of insulin therapy. The overlapping features of both types of diabetes have been previously referred to as ‘atypical diabetes,’ ‘diabetes type 1 ½,’ and has been recently called ‘ketosis prone DM (KPD).’

A patient with KPD has severe hyperglycemia with associated ketosis that can be managed without insulin after a few months; and can maintain acceptable glycemic control with diet or oral agents. Ketoacidosis is not rare in T2D but occurs in settings of acute illness and infections. Our patient had a phenotype compatible with T2D but was diagnosed with DM after an episode of DKA which is a presentation more commonly seen among those with T1D. Our patient had no precipitating cause for DKA and was later noted to have near-normoglycemic remission from insulin and oral hypoglycemic agents (OHA).

Assessment of autoimmunity and β-cell secretory reserve of newly diagnosed DM patients after 1–3 weeks of resolution of the index DKA episode was reported to predict the ability of the newly diagnosed KPD patient to discontinue insulin and remain in near-normoglycemic remission. They are divided into four groups based on autoimmunity and β-cell function (Aβ) classification scheme: those patients with autoimmune disease with absent (Aβ) or preserved (Aβ) β-cell function and those without autoimmune diabetes with absent (Aβ) or preserved (Aβ) β-cell function.

Patients with negative β-cell function, with or without autoimmune markers, have clinical and biochemical characteristics of T1D implying lifelong insulin dependency. These groups are comparable to the American Diabetes Association (ADA) classification of autoimmune and idiopathic T1D respectively.

Adequate β-cell function after an episode of DKA, regardless of autoimmunity is compatible with T2D. Types Aβ and Aβ are termed as ‘ketosis prone type 2 DM.’ β-cell reserve determines the definitive management of diabetes and is the best predictor of remission in KPD. Insulinopenia will require lifelong insulin (T1D), while those with adequate β-cell function benefit from lifestyle modification and OHA.

Autoimmunity, on the other hand, is a determinant of future β-cell function. Presence of autoantibodies (GAD65Ab) at diagnosis is highly predictive for future insulin use at a rate of 1-2% per year after 1-5 years of therapy with diet, exercise, OHA. Fifty percent of DKA patients classified as Aβ (represents 7%) have progressive decline of β-cell function eventually becoming insulin requiring following a clinical course that resembles T1D within the first two years of diagnosis. Serial C-peptide determination is recommended every 6 months to track β-cell deterioration.

The majority of KPD is represented by Aβ (74%), whose clinical course mimics that of T2D after resolution of DKA. They are able to achieve near-normoglycemic remission within 10 weeks (~70%) and 40% remain insulin independent a decade after diagnosis. The cause of the sudden, severe but reversible β-cell dysfunction is unknown in 50% of cases (unprovoked), whereas the other half is precipitated by an acute illness or noncompliance with medications (provoked). The unprovoked ketosis among β-cell adequate groups is postulated to be secondary to
impaired ketone oxidation and fatty acid utilization. The provoked AβKPD group has better glycemic control (twice the frequency of attaining HbA1c <7%), 2-fold greater improvement in β-cell function (after 12 months of treatment), and a higher rate of insulin independence.5-12-14

All newly diagnosed KPD patients should be discharged with basal insulin regimen after resolution of DKA. Any attempt to discontinue insulin should be based on the Aβ classification scheme. If blood glucose monitoring of β+ patients after 2 weeks is within ADA target goals (preprandial 90-130 mg/dl; postprandial <180 mg/dl), the insulin dose can be decreased by 50% and is reassessed after another week. If the patient achieves ADA target by the second visit, insulin regimen can be discontinued but close monitoring is warranted. If blood glucose is uncontrolled, insulin regimen is intensified.3

Our patient (unprovoked AβKPD) has sufficient insulin secretion that enables him to sustain good glycemic control through diet alone. Absence of autoimmunity makes him less susceptible to future β-cell failure and recurrence of DKA. However, we need to be vigilant since the patient’s DKA was unprovoked and an underlying β-cell dysfunction may be present. Close follow-up is prudent to avoid recurrence of life-threatening ketoasis, to anticipate complications and to monitor eventual shifts in the therapy of patients with KPD.

CONCLUSION

Diabetes can present in various ways. Phenotypical characteristics are not enough to classify them into either T1D or T2D. Tests for autoantibodies and β-cell function are recommended for patients with overlapping features of both types of DM, and may be considered for patients on insulin who want to be shifted to oral agents, and patients who are uncontrolled with oral agents and are suspected to have inadequate β-cell function. These tests will guide clinicians regarding diagnosis, response to treatment and anticipation of complications.

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This patient exemplifies the increasing difficulty in differentiating between Type 1 and Type 2 DM among young patients, based on clinical grounds alone. This is especially true in the presence of obesity. The category to which the patient in the case was classified would correspond to Type 1 DM or monogenic diabetes (maturity onset diabetes of the young, MODY). Both will have Aβ-+ designation with C-peptide between 0.6 and 2 ng/mL. A young patient eventually diagnosed to have Type 2 DM is a cause for alarm. A comparison of the outcomes among diabetic individuals showed a two-fold increase in case fatality in young-onset T2DM compared with T1DM of a similar age of onset and disease duration.1 This is accounted for by the greater number of cardiovascular deaths,1 which has thus created the notion that young-onset T2DM is a more aggressive disease compared to T1DM. Thus, the prevention of obesity among children cannot be over-emphasized, as this is the single most important factor for the rising trend in the occurrence of Type 2 DM in this age group.

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* A- means negative antibody while B+ means reserved beta cell function.

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