CASE REPORT

Development of Exogenous Insulin Antibody Syndrome in a Patient with Newly Diagnosed Type 1 Diabetes Successfully Treated with Oral Immunosuppressive Monotherapy

Terri Jerkins · David S. H. Bell

Received: June 2, 2021 / Accepted: July 23, 2021 / Published online: August 17, 2021 © The Author(s) 2021

ABSTRACT

Exogenous insulin antibody syndrome (EIAS), which rarely occurs in the patient with type 1 diabetes, results in antibody-induced insulin resistance, hyperglycemia, ketosis, ketoacidosis, and hypoglycemia when insulin is released from the saturated insulin antibodies. Recommended treatment regimens include glucocorticoids, immunosuppressants, and plasmapheresis. In the patient with type 1 diabetes, glucocorticoids may by inducing and/or worsening ketoacidosis be contraindicated. With immunosuppressants, various anecdotal treatment regimens have been reported. Currently the most commonly recommended regimen is intravenous immunosuppressive therapy in combination with oral immunosuppressants. Herein we describe a patient in whom oral immunosuppressant monotherapy with mycophenolate resulted in the cure of EIAS, thus avoiding the expense associated with intravenous immunosuppressant therapy and/or hospitalization for plasmapheresis.

Keywords: Type 1 diabetes; Insulin antibodies; Oral immunosuppressants; Exogenous insulin antibody syndrome

Key Summary Points

- Insulin antibody syndrome results in insulin resistance, hyperglycemia and later hypoglycemia.
- Therapies for insulin antibody syndrome are anecdotal and include plasmapheresis, immunosuppression and glucocorticoids.
- The most common current recommendation is combined intravenous and oral immunosuppressants.
- In this case we show the efficacy of the more convenient and economic oral immunosuppressant monotherapy.

INTRODUCTION

Insulin resistance is defined as the need to use 200 or more units of insulin a day or more than 2 units of insulin per kilogram of body weight. Insulin resistance caused by the generation of insulin antibodies has been name exogenous...
insulin antibody syndrome (EIAS) [1]. First isolated in 1956, insulin antibodies were shown to be present in 98% of patients who utilized insulin when only insulins derived from animal sources were available [2]. Since that time, with the use of more purified insulins the incidence has dropped significantly but EIAS still occurs with the use of both human and analogue insulins [3, 4]. Some analogues have shown decreased antibody formation but there is no consistent recommendation for changes in insulin formulation to decrease insulin antibodies [5, 6].

Severe insulin resistance due to exogenous insulin antibodies is extremely rare and is estimated to occur in less than 0.17% of patients with diabetes. Insulin antibody levels have not been shown to correlate with glycemic control. However, the EIAS causes extreme insulin resistance leading to post-prandial hyperglycemic and ketosis and severe hypoglycemia due to release of insulin from the insulin antibody complex. The majority of case reports are in patients with type 2 diabetes, and EIAS is very uncommon in type 1 diabetes where it has been associated with diabetic ketoacidosis (DKA).

Similar presentations have occurred in the presence of endogenous insulin antibodies (Hirata syndrome) and in the insulin receptor antibody syndrome. Treatment of all of these syndromes has focused on elimination of hypoglycemia because of the increased mortality associated with hypoglycemia.

In 2010, the US National Institutes of Health issued guidelines for the treatment of insulin receptor antibodies causing insulin resistance. Initial therapy is glucocorticoids; if glucocorticoids are not successful, treatment with oral or intravenous immunosuppression and/or plasmapheresis is undertaken [7]. All of these treatments have been tried in EIAS with varying results. However, treatment has not been instituted until the syndrome has been present for at least 5 months. We present a patient who was treated within 2 weeks of the diagnosis of type 1 diabetes due to rapidly accelerating insulin requirements and recurrent DKA despite very high doses of insulin and no hypoglycemia.

Written informed consent was obtained from the patient.

CASE PRESENTATION

A 51-year-old woman was in her usual state of good health until 10 days prior to hospital admission when she suddenly developed polyuria, polydipsia, and weight loss accompanied by a serum glucose of 594 mg/dl, a metabolic acidosis, and elevated serum beta-hydroxybutyrate. She had no history of glucose intolerance or gestational diabetes though her family history was positive for type 2 diabetes in her father as well as autoimmune thyroid disease in her sister and a niece. She was easily treated with a low dose intravenous insulin DKA protocol and quickly transitioned to a basal bolus subcutaneous insulin regimen utilizing detemir and aspart insulins. At the onset, HbA1c was 11.8%. GAD-65 antibodies were positive at 3184.1 U/ml, antipancreatic islet cell antibodies were 1:256, and other than a positive anti-parietal antibody a screen for other autoimmune endocrinopathies was negative. She was discharged on detemir 18 units at bedtime and aspart 8 units with each meal (0.58 units/kg).

She was seen as an outpatient 4 days after discharge and was noted to have had a rapid increase in glucose levels. Insulin antibodies were measured and found to be 13 lU/ml (normal < 5 lU/ml). Over the next week her insulin was rapidly increased. However, even rapid increases in insulin doses (detemir 80 units daily and multiple boluses of 60 units aspart up to five times daily) did not lower her blood sugar levels so that she soon developed ketoacidosis in spite of utilizing 225 units of insulin per day. On readmission to hospital, she required 50% more intravenous insulin than on the previous admission to achieve glycemic control. Transitioning to subcutaneous human NPH and regular insulin at more than 400 units of insulin per day failed to control her hyperglycemia and continuous subcutaneous insulin infusion (CSII) with U-500 insulin was started in conjunction with orally administered mycophenolate mofetil at a dose of 500 mg b.i.d. with the intention of adding intravenously administered rituximab as an outpatient. One day after the first dose of mycophenolate her insulin requirement began
to drop, hypoglycemia occurred even with aspart insulin at a basal rate of 0.85 unit/h and a carbohydrate ratio of 1 unit per 7 g of carbohydrate. With lowering of basal rates her glucose levels were controlled and she was discharged from hospital. As an outpatient her mycophenolate was increased to 1000 mg b.i.d. which resulted in a further decrease in her insulin requirements. Two months after the original presentation her insulin antibodies were negative, her GAD-65 antibody level had dropped, and her HbA1c was 7.8% (see Table 1). The planned addition of intravenously administered rituximab, which was delayed for logistical reasons, was found to be unnecessary.

She was continued on mycophenolate 1000 mg b.i.d. for 5 months after the initial presentation. Over the following month she was slowly weaned off mycophenolate without a recurrence of insulin resistance. One year after initial presentation she was still in the “honeymoon period” with her HbA1c being 6.7% and her total daily utilization of insulin less than 40 units a day (0.55 units/kg) which was 10% of her initial need (see Table 1).

**DISCUSSION**

Antibodies to exogenous insulin with the availability of first purified animal insulins and later recombinant DNA human insulin have been greatly but not entirely reduced. However, sporadic occurrences of high insulin antibody titers while being treated with recombinant human insulin or insulin analogues have been reported in both type 1 and type 2 diabetes usually occurring several months or years after the initiation of insulin therapy. Insulin antibodies can lead not only to hypersensitivity but also to glycemic variability, hyperglycemia, ketoacidosis as well as hypoglycemia when the insulin resistance has been overcome and insulin is released from the saturated antibodies.

Therapies that have shown efficacy include withdrawal of insulin and utilization of oral agents in type 2 diabetes. However, with type 1 diabetes replacement of insulin with oral agents is not an option and the proven efficacious treatments are the use of glucocorticoids, plasmapheresis, and immunosuppressants [8–12]. However, there has never been a placebo controlled or randomized study to compare these options and a randomized study is made less likely to occur because of the rarity of this condition.

Glucocorticoid therapy is often counterproductive in that insulin resistance and hyperglycemia are often worsened and the addition of immunosuppressants is needed to either replace or augment glucocorticoid therapy. In addition, in the patient with type 1 diabetes there is a higher risk of ketoacidosis. If these combinations are not successful plasmapheresis is generally recommended. However, plasmapheresis involves hospitalization with its associated expense. Even in the outpatient setting the administration of intravenous immunosuppressants results in considerable expense and inconvenience.

Therefore, an affordable oral immunosuppressant such as mycophenolate (MMF of Cellcept) which has previously been shown to be effective in children should be the preferred initial therapy for EIAS. In this case, after MMF was started, rituximab therapy was prescribed but was delayed for logistical reasons, and oral MMF monotherapy was found to reduce insulin antibody levels so that the resistance to the action of insulin disappeared. The patient has sustained a prolonged remission but continues to have type 1 diabetes which is in excellent control.

In conclusion, for the first time we have shown in adults that exogenous insulin

| Test                  | Pre-therapy | Post-therapy |
|-----------------------|-------------|--------------|
| Insulin antibodies    | 13 U/ml     | < 0.15 U/ml  |
| GAD-65 antibodies     | 134 U/ml    | Negative     |
| HbA1c                 | 11.8%       | 6.7%         |
| Weight/kg             | 72          | 64           |
| Insulin units per day | 400         | 40           |
| Insulin units per kg  | 5.5         | 0.55         |
antibody syndrome can be treated as an outpa-
tient with mycophenolate monotherapy which
potentially avoids the expense of intravenous
immunosuppressive therapy and/or plasmapheresis and the expense of hospitalization. Also, earlier treatment of the syndrome prior to
the development of hypoglycemia and during
propagation of the immune response may allow
a quicker remission.

ACKNOWLEDGEMENTS

Funding. No funding or sponsorship was
received for this study or publication of this
article.

Authorship. All named authors meet the
International Committee of Medical Journal
Editors (ICMJE) criteria for authorship for this
article, take responsibility for the integrity of
the work as a whole, and have given their
approval for this version to be published.

Authors’ Contributions. Authors con-
tributed equally to the manuscript in all areas.

Disclosures. Terri Jerkins and David S.
H. Bell have no disclosures to declare.

Compliance with Ethics Guidelines. Con-
sent to publish was obtained from the patient
by Dr Jerkins.

Data Availability. Data is available upon
request from Dr. Jerkins.

Open Access. This article is licensed under a
Creative Commons Attribution-NonCommer-
cial 4.0 International License, which permits
any non-commercial use, sharing, adaptation,
distribution and reproduction in any medium
or format, as long as you give appropriate credit
to the original author(s) and the source, provide
a link to the Creative Commons licence, and
indicate if changes were made. The images or
other third party material in this article are
included in the article’s Creative Commons licence, unless indicated otherwise in a credit

REFERENCES

1. Xiaolei H, Chen F. Exogenous insulin antibody
syndrome (EIAS); a clinical syndrome associated
with insulin antibodies induced by exogenous
insulin in diabetic patients. Endocr Connect.
2018;7:R47–55.

2. Berson SA, Yallow RS, Bauman A, Rothchild MA,
Newerly K. Insulin-I131 metabolism in human
subjects: demonstration of insulin binding globulin
in the circulation of insulin treated subjects. J Clin
Investig. 1956;35:170–90.

3. Schernthaner G, Borkenstein M, Fink M, Mayr WR,
Menzel J, Schober E. Immunogenicity of human
insulin (Novo) or pork monocomponent insulin in
HLA-DR-typed insulin dependent diabetic individ-
uals. Diabetes Care. 1983;6(Supplement 1):43–8.

4. Maneshi F, Fineberg SE, Kohner EM. Successful
treatment of immune mediated insulin resistance
by human insulin (recombinant DNA). Diabetes
Care. 1981;5(Supplement 2):175–9.

5. Lahtela JT, Knip M, Paul R, Antonen J, Salmi J.
Severe antibody mediated human insulin resis-
tance: successful treatment with the insulin analog
lispro. A case report. Diabetes Care. 1997;20:71–3.

6. Yanai H, Adachi H, Hamasaki H. Diabetic ketosis
caused by the insulin analog aspart-induced anti-
insulin antibody: successful treatment with the
newest insulin analog glulisine. Diabetes Care.
2011;34:e108.

7. Malek R, Chong AY, Lupsa BC, et al. Treatment of
type B insulin resistance: a novel approach to
reduced insulin receptor autoantibodies. J Clin
Endocrinol Metab. 2010;9:3641–7.

8. Church D, Cardoso L, Kay R, et al. Assessment and
management of anti-insulin autoantibodies in
varying presentations of insulin autoimmune syn-
drome. J Clin Endocrinol Metab. 2018;103(10):
3854–5.
9. Jassam N, Amin N, Barth JH, et al. Analytical and clinical challenges in a patient with concurrent type 1 diabetes, subcutaneous insulin resistance and insulin autoimmune syndrome. Endocrinol Diabetes Metabol Case Rep. 2014;2014:130086.

10. Segal T, Webb E, Viner R, Pusey C, Wild G, Allgrove J. Severe insulin resistance due to insulin antibodies: successful treatment with the immunosuppressant MMF. Pediatr Diabetes. 2008;9(1):250–4.

11. Ahmed M, Subbalaxmi M, Anne B, Deme S. Recurrent diabetic ketoacidosis and extreme insulin resistance due to anti-insulin antibodies: response to immunosuppression and plasma exchange. Diabetes Technol Ther. 2021;23:227–9.

12. Matsuyoshi A, Shimaoda S, Tsuruzoe K, et al. A case of slowly progressive type 1 diabetes with unstable glycemic control caused by unusual insulin antibody and successfully treated with steroid therapy. Diabetes Res Clin Pract. 2006;72(3):238–43.