Type B insulin resistance syndrome associated with connective tissue disease and psoriasis

Citation for published version:

Digital Object Identifier (DOI):
10.1530/EDM-20-0027

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Publisher's PDF, also known as Version of record

Published In:
Endocrinology, Diabetes and Metabolism Case Reports

General rights
Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy
The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.
Type B insulin resistance syndrome associated with connective tissue disease and psoriasis

Agnieszka Łebkowska¹, Anna Krentowska¹, Agnieszka Adamska², Danuta Lipińska², Beata Piasiecka², Otylia Kowal-Bielecka³, Maria Górska², Robert K Semple⁴ and Irina Kowalska¹

¹Department of Internal Medicine and Metabolic Diseases, ²Department of Endocrinology, Diabetology and Internal Medicine, ³Department of Rheumatology and Internal Diseases, Medical University of Białystok, Białystok, Poland, and ⁴Centre for Cardiovascular Science, Queen’s Medical Research Institute, University of Edinburgh, Edinburgh, UK

Summary

Type B insulin resistance syndrome (TBIR) is characterised by the rapid onset of severe insulin resistance due to circulating anti-insulin receptor antibodies (AIRAs). Widespread acanthosis nigricans is normally seen, and co-occurrence with other autoimmune diseases is common. We report a 27-year-old Caucasian man with psoriasis and connective tissue disease who presented with unexplained rapid weight loss, severe acanthosis nigricans, and hyperglycaemia punctuated by fasting hypoglycaemia. Severe insulin resistance was confirmed by hyperinsulinaemic euglycaemic clamping, and immunoprecipitation assay demonstrated AIRAs, confirming TBIR. Treatment with corticosteroids, metformin and hydroxychloroquine allowed withdrawal of insulin therapy, with stabilisation of glycaemia and diminished signs of insulin resistance; however, morning fasting hypoglycaemic episodes persisted. Over three years of follow-up, metabolic control remained satisfactory on a regimen of metformin, hydroxychloroquine and methotrexate; however, psoriatic arthritis developed. This case illustrates TBIR as a rare but severe form of acquired insulin resistance and describes an effective multidisciplinary approach to treatment.

Learning points:

• We describe an unusual case of type B insulin resistance syndrome (TBIR) in association with mixed connective tissue disease and psoriasis.
• Clinical evidence of severe insulin resistance was corroborated by euglycaemic hyperinsulinaemic clamp, and anti-insulin receptor autoantibodies were confirmed by immunoprecipitation assay.
• Treatment with metformin, hydroxychloroquine and methotrexate ameliorated extreme insulin resistance.

Background

Type B insulin resistance syndrome (TBIR) is a rare autoimmune disorder characterised by the development of anti-insulin receptor antibodies (AIRAs). It most commonly leads to extreme insulin resistance with severe hyperglycaemia and weight loss; however, hypoglycaemia may be seen, especially where antibody titre is low (1). TBIR is commonly associated with underlying autoimmune diseases, such as systemic lupus erythematosus (SLE), various connective tissue diseases, interstitial lung disease or organ-specific dermatological, haematological, or hepatic disorders (1, 2, 3). Here, we present an unusual case of TBIR associated with connective tissue disease and psoriasis, with insulin resistance confirmed by hyperinsulinaemic euglycaemic clamp and stabilised with
a regimen of corticosteroids, hydroxychloroquine and metformin.

Case presentation

A 27-year-old Caucasian man with a 2-year history of psoriasis was hospitalised because of an unexplained weight loss of 20-kg in one year. He also reported fatigue and a new onset of Raynaud’s phenomenon. One year earlier, he had undergone haematological evaluation because of peripheral lymphadenopathy and enlarged parotid glands, with results deemed consistent with viral infection. Lymphoproliferative disease was excluded on the basis of lymph node histopathological examination. There was no family history of diabetes or autoimmune disease.

Investigation

Physical examination revealed malnutrition. Height was 180 cm and weight 53.3 kg (BMI 16.4 kg/m²). There was widespread acanthosis nigricans in the axillae, antecubital fossae, groins, and on the thumbs (Fig. 1). Sclerodactyly, psoriatic lesions, enlarged parotid glands, palpable cervical and axillary lymph nodes, and tachycardia were also noted. Dual energy X-ray absorptiometry revealed a paucity of subcutaneous and visceral fat and decreased lean body mass (Fig. 2). Table 1 shows laboratory findings on admission. Full blood count revealed anaemia, leucopaenia with lymphopaenia, and thrombocytopaenia. Blood glucose concentration on admission was 20.1 mmol/L with glycated haemoglobin (HbA1c) 12.4% (112 mmol/mol) and associated glycosuria. From day 17 of observation, the patient developed asymptomatic morning fasting hypoglycaemia, more frequently observed in the further...
course of treatment (glycaemic nadir 1.5 mmol/L). The concentration of endogenous insulin, measured after the reduction of insulin doses, when the patient required only low doses of prandial insulin, exceeded 300 μIU/mL. Fasting C-peptide concentration was within the normal range and increased adequately in glucagon stimulation test (0 min: 2.87 ng/mL, 6 min: 4.56 ng/mL). Plasma adiponectin concentration, assayed as previously described, was 13.8 mg/L (4). Insulin autoantibodies, glutamic acid decarboxylase autoantibodies and insulinoma-associated-2 autoantibodies were negative. Immunoprecipitation assay revealed anti-insulin receptor antibodies in two separate blood samples, confirming TBIR (Fig. 3). Hyperinsulinaemic euglycaemic clamping was performed in accordance with the protocol by DeFronzo et al. (5). The rate of whole-body glucose uptake (M value) was calculated as the mean glucose infusion rate during the last 40 min of the clamp, corrected for the glucose space and normalised for fat-free mass (FFM). The clamp confirmed severe insulin resistance with an M value of 2.1 mg/kgFFM/min (the mean M value obtained in a cohort of 60 healthy normal-weight men was 7.1 ± 2.4 mg/kgFFM/min (6)).

Due to clinical features of connective tissue disease, a wider autoimmune screen was undertaken. High titres of antinuclear antibodies and anti-ribonucleoprotein antibodies were found, with positive Ro52 and anti-dsDNA
antibodies. Direct Coombs test was positive, and serum complement concentrations were decreased. Collectively this led us to the diagnosis of mixed connective tissue disease. CT of the lungs and the abdomen, MRI of the brain, pulmonary function tests, and histological assessment of bone marrow, parotid glands and lymph nodes failed to demonstrate neoplasia.

Treatment

The treatment course is presented in Fig. 4. The patient was initially treated with i.v. insulin infusion, followed by intensive insulin therapy. This regimen did not improve blood glucose concentration despite daily insulin doses from 60 IU/24 h to 110 IU/24 h (>2 IU/kg of body weight). Metformin (850 mg three times a day) allowed gradual reduction of insulin doses (from 110 IU/24 h to 40 IU/24 h in 10 days). Given the coexistence of TBIR and mixed connective tissue disease, a cycle of pulsed methylprednisolone therapy was administered (250 mg/day for 3 days), followed by oral prednisone in decreasing doses for 1 month. Simultaneously, hydroxychloroquine (400 mg/day) was started and euglycaemia was maintained, which allowed withdrawal of insulin and reduction of metformin dose to 3 × 500 mg. Due to fasting morning hypoglycaemic episodes, the morning dose of metformin was stopped.

Outcome and follow-up

Six months after the initiation of treatment, weight had increased to 59.8 kg (BMI 18.5 kg/m²). Acanthosis nigricans and sclerodactyly also improved but did not remit entirely (Fig. 5). HbA1c decreased to 7.3% (56 mmol/mol) with only rare episodes of fasting hypoglycaemia. Fasting insulin concentration remained elevated at 126 µU/mL. The daily dose of metformin was reduced to 1000 mg/day in two doses. Thirty-three months later, inflammation of the elbow and interphalangeal joints of both hands developed with the intensification of psoriatic lesions (Fig. 6). Methotrexate treatment was initiated as an adjunct to hydroxychloroquine. During this therapy,
A Łebkowska and others

Type B insulin resistance syndrome

ID: 20-0027; August 2020
DOI: 10.1530/EDM-20-0027

weight gain increased further (BMI 19.7 kg/m²), with the disappearance of acanthosis nigricans, significant improvement of arthritis and reduction of psoriatic lesions. Fasting glucose concentration was 4.3 mmol/L.

Discussion

Type B insulin resistance is rare. The largest longitudinal cohort of 24 patients has been reported by researchers from the United States National Institutes of Health (NIH) (1). It occurs mainly in the fourth to sixth decade of life, usually against a background of a rheumatological illness, most commonly connective tissue disease, lupus erythematosus, or as a paraneoplastic manifestation (1, 2, 7). We report a younger male Caucasian patient with typical symptoms of TBIR in association with mixed connective tissue disease and psoriasis. We assessed insulin sensitivity by hyperinsulinaemic euglycaemic clamp to confirm the severity of insulin resistance.

There is no pathognomonic clinical feature of TBIR, although rapid onset of extreme insulin resistance with widespread acanthosis nigricans, severe hyperglycaemia with attendant catabolic state and weight loss are characteristic. Type B insulin resistance syndrome is caused by AIRAs, which, based on rodent and cellular studies, act as agonists at the insulin receptor when the titre is low and as antagonists due to receptor downregulation when at high titre (8). This means that metabolic consequences of AIRAs are variable, ranging from extreme insulin resistance and hyperglycaemia in the first phase of the disease to recurrent fasting hypoglycaemia during early remission. Klubo-Gwiezdzinska et al. confirmed that all studied patients with TBIR experienced hypoglycaemic episodes when AIRA titre decreased, which allowed to reduce insulin doses (9). Initial presentation with hypoglycaemia alone is rare and was observed in only 13% of patients in the NIH cohort (1). Our patient started to develop fasting morning hypoglycaemia shortly after diabetes diagnosis. Despite reduction of insulin doses and eventually withdrawal of exogenous insulin, these episodes recurred. Thus, the suggestion of prolonged degradation of insulin–insulin receptor complexes or the coexistence of different populations of antibodies might be the possible pathomechanism (1).

The biochemical triad of markedly elevated fasting insulin concentration, hyperadiponectinaemia and low/normal fasting triglyceride concentrations was discussed as a “working” clinical definition of TBIR (10). Adiponectin level >7 mg/L in subjects with symptoms of severe insulin resistance had a 97% positive predictive value for defects of insulin receptor function (11). Our patient had high fasting insulin level, low triglycerides concentration and increased adiponectin concentration, which confirmed earlier findings.

Treatment of TBIR is challenging. The majority of reported cases of TBIR are associated with other autoimmune diseases, most often systemic lupus erythematosus or other connective tissue disease. To control hyperglycaemia at admission, the dose of insulin required intravenously in our patient was lower than presented in other cases, but was markedly higher than in “common” type 2 diabetes (7). Descriptions of use of metformin, sulphonylureas and thiazolidinediones in therapy of hyperglycaemia in TBIR have demonstrated variable efficacy (12, 13). In our patient, metformin allowed significant decrease in the dose of insulin.

The therapy requires intensive monitoring for side effects of immunosuppressive drugs and insulin titration due to glycaemic variability in the course of the disease (9).
The NIH have recently proposed a treatment regimen consisting of rituximab, monthly high-dose glucocorticoids and cyclophosphamide, which has proved effective in an uncontrolled case series and allowed for discontinuation of insulin therapy in the studied patients (9). Combination immunosuppressive therapy, described in a prospective cohort study, followed by maintenance therapy with azathioprine, succeeded in reversing diabetes, signs of insulin resistance, and hyperandrogenism in women and was relatively safe (9). Unfortunately, the availability of rituximab in Poland is restricted in connective tissue disorders other than rheumatoid arthritis due to its high price. Other reported approaches to treatment have included use of prednisolone with hydroxychloroquine and azathioprine, plasmapheresis or i.v. immunoglobulin (3, 14, 15). In this case, corticosteroids and metformin permitted withdrawal of insulin and ameliorated insulin resistance symptoms. In several reported TBIR cases, remission has occurred spontaneously. In a cohort described by Arioglu et al., spontaneous remission was observed in 33% of patients (1). Time to remission in this group ranged between 11 and 48 months and was similar to patients treated with immunosuppressive agents. Mortality rates in both groups were also comparable. However, in the case of coexisting disorders, such as connective tissue diseases, immunosuppressive treatment may be necessary to ameliorate their symptoms. In the presented case, the occurrence of psoriatic arthritis with connective tissue disease in 3 years of follow-up caused changes to the treatment regimen guided by a rheumatologist. Methotrexate with hydroxychloroquine helped reduce the severity of autoimmunological symptoms with concurrent decrease in the concentration of fasting insulin and reduction of hypoglycaemia episodes.

In conclusion, the presented case illustrates TBIR coexisting with other autoimmune conditions. Beyond the clinical signs of insulin resistance, hyperinsulinaemic euglycaemic clamp provided standardised biochemical confirmation. Targeted individualised therapy with a combination of metformin, hydroxychloroquine and methotrexate proved effective.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

Patient consent
Written informed consent to publish these findings was obtained from the patient.

Author contribution statement
Agnieszka Lebkowska and Anna Krentowska were involved in diagnostic and therapeutic process and writing the article. Agnieszka Adamska, Danuta Lipińska, Beata Piaśecka, and Maria Górska contributed to the diagnostic and therapeutic process. Otylia Kowal-Bielecka was involved in the diagnostic and therapeutic process and rheumatological consultations. Robert Semple was involved in the assessment of anti-insulin receptor antibodies and writing of the article. Irina Kowalska contributed to the diagnostic and therapeutic process, writing of the article, and final approval of the version to be submitted.

Acknowledgement
The authors would like to thank Cornelia Gewert for undertaking immunoprecipitation assays.

References
1 Arioglu E, Andewalt A, Diabo C, Bell M, Taylor SI & Gorden P. Clinical course of the syndrome of autoantibodies to the insulin receptor (type B insulin resistance): a 28-year perspective. Medicine 2002 81 87–100. (https://doi.org/10.1097/00005792-200203000-00001)
6 Stefanowicz M, Nikolajuk A, Matulewicz N & Karczewska-Kupczewska M. Adipose tissue, but not skeletal muscle, sirtuin 1 expression is decreased in obesity and related to insulin sensitivity. Endocrine 2018 60 263–271. (https://doi.org/10.1007/s12020-018-1544-1)
8 Dons RF, Havlik R, Taylor SI, Baird KL, Chernick SS & Gorden P. Clinical disorders associated with autoantibodies to the insulin receptor. Simulation by passive transfer of immunoglobulins to rats. Journal of Clinical Investigation 1983 72 1072–1080. (https://doi.org/10.1172/JCI111032)


Received in final form 15 June 2020
Accepted 13 July 2020