Global Journals $end{transformula} \mathbb{A}T_{\mathbf{E}} X$ JournalKaleidoscope

Artificial Intelligence formulated this projection for compatibility purposes from the original article published at Global Journals. However, this technology is currently in beta. *Therefore, kindly ignore odd layouts, missed formulae, text, tables, or figures.*

¹ Insulin Autoimmune Syndrome Treated with Plasmapharesis

Kavya Jonnalagadda¹ and Praveen V. Pavithran²

¹ Sree Charith Hospitals, Amrita Institute of Medical Sciences and Research

Received: 8 December 2019 Accepted: 5 January 2020 Published: 15 January 2020

6 Abstract

2

3

4

 $_{7}$ A 66-year male with a history of Central Serous Retinopathy presented with recurrent

⁸ episodes of hypoglycemia. On evaluation, he was found to have insulin-mediated

⁹ hypoglycemia with serum insulin of 300uIU/ml, C peptide 27.51ng/ml, when the blood

¹⁰ glucose was 46mg/dl. High insulin levels above 100uIU/ml, led to suspicion of Autoimmune

¹¹ hypoglycemia and were confirmed by a high anti-insulin antibody titer of 300U/ml. Imaging

¹² was negative for Insulinoma. The patient was started on low dose oral prednisolone under

¹³ ophthalmological monitoring, but as there was no symptomatic improvement, the dose was

¹⁴ increased following which there was a flare-up of CSR. The patient was initiated on

¹⁵ plasmapheresis following which his hypoglycemia improved with drop in anti-insulin antibody

¹⁶ titers to 29U/ml. The patient was maintained on low dose steroids, which were tapered and

¹⁷ stopped over the next six months with complete resolution of hypoglycemia and normalization

- 18 of anti-insulin antibody titers.
- 19

20 Index terms— hypoglycemia, autoimmune, plasmapheresis, anti-insulin antibody.

²¹ 1 Introduction

utoimmune hypoglycemia is a rare but key cause of hyperinsulinemic hypoglycemia, which either resolves by itself
or will need immunomodulators. This case highlights the usefulness of plasmapheresis when routine treatments
like steroids are contraindicated and or ineffective. Lack of awareness of the condition can lead to unnecessary
interventions for an insulinoma.

26 **2 II.**

27 **3** Case Presentation

A 66-year male with a history of Central Serous Retinopathy was referred for the evaluation of recurrent episodes of 28 Hypoglycemia occurring in both fasting and postprandial state. Four months before presentation, he had tinnitus 29 for which he had taken alternative medicines and multivitamins following which he experienced recurrent episodes 30 of fatigue, palpitations, sweating, and day time sleep, which went unexplained. Hypoglycemia was diagnosed 31 when he was found unresponsive in the bed early in the morning, which led to emergency room admission 32 33 and documentation of venous plasma glucose as 38 mg/dl. Physical examination was unrevealing. The patient 34 underwent a supervised fasting test and developed hypoglycemia within four hours of fasting. At this time, venous 35 plasma glucose levels were 46.1 mg/dl, and 32.9 mg/dl, Insulin was 300uIU/ml, C-peptide was 27.51ng/ml. High insulin levels more than 100uIU/ml led to suspicion of autoimmune hypoglycemia and confirmed with high levels 36 of anti-insulin antibody titers >300U/ml. The imaging of the pancreas was normal. HLA typing was not done. 37 The patient was started on a low dose of oral prednisolone 20 mg OD, but as the patient continued to have 38 episodes of hypoglycemias the strength of steroids has been increased to 40 mg/day with which the hypoglycemic 39 symptoms improved, but the patient had flare-up of Central Serous Retinopathy, and the ophthalmologist advised 40 to decrease the dosage of steroids. In view of this, a trial of plasmapheresis was initiated, and he underwent six 41

42 sessions on alternate days following which anti-insulin antibody titers have dropped to 29U/ml, and the frequency 43 of hypoglycemic episodes still reduced. The patient was maintained on low dose steroids, which were tapered 44 and stopped over the next six months. Presently the antibody titer has dropped to 1.3U/ml. Currently, he is

45 free of hypoglycemic symptoms.

46 **4 III.**

47 5 Discussion

Hirata disease or Insulin autoimmune syndrome (IAS), is a rare cause of hyperinsulinemic hypoglycemia with 48 autoantibodies to native insulin in individuals without prior exposure to exogenous insulin. [1] More than 300 49 cases of Insulin autoimmune syndrome were published. [2] Highest number was recorded in Japan, followed by 50 Europe and the US. [3] In these patients, the cause of hypoglycemia was due to the binding of the antibodies 51 to the endogenous insulin, which leads to increased levels of bioavailable insulin, which were physiologically 52 inappropriate, causing either hyper or hypoglycemia. [4] More than half of IAS patients had recent exposure 53 to drugs containing a sulfhydryl group. [5]Methimazole, is the most common trigger for IAS. Other drugs that 54 can cause IAS include imipenem, penicillins, isoniazid, glutathione, carbimazole, tolbutamide, D penicillamine, 55 Interferon-alpha, procainemide. There are many other drugs that can cause IAS, but the above-mentioned 56 drugs are common drugs that can provoke IAS. [6] Alpha-lipoic acid, a health supplement with antioxidant 57 and anti aging properties used in the treatment of neurological symptoms of diabetes, was associated with 58 IAS in recent years. [7] A IAS is a transient condition with spontaneous resolution within 3-6 months of 59 diagnosis in the majority of the patients. [8]Small frequent meals low in carbohydrates remain the first line 60 of treatment in those with intractable hypoglycemia. Drugs responsible for hypoglycemia, if present, should 61 be stopped. Glucocorticoids are the mainstay of treatment. Other therapeutic options include acarbose, which 62 delays carbohydrate digestion and absorption, plasmapheresis to reduce insulin autoantibody titers. All these 63 therapeutic options have demonstrated varying success in the management of IAS. [4] [9] If hypoglycemia persists, 64 combination therapy with Glucocorticoids and immunosuppressants like azathioprine or 6-mercaptopurine along 65 66 with plasmapheresis should be considered. [9]In our patient as there is a flare in the CSR steroids could not be 67 continued at a higher dose and we opted to go ahead with a combination of steroids and plasmapheresis with which we were able to come down on the dose of steroids following which his ocular symptoms and hypoglycemic 68 symptoms have improved. An anti-CD20 monoclonal antibody Rituximab, which acts by blocking antibody 69 responses and suppress insulin autoantibodies, is the last hope for those with refractory hypoglycemia. [10] IV. 70

71 6 Conclusion

1. Autoimmune hypoglycemia is one of the usual causes of hyperinsulinemic hypoglycemia if druginduced etiology

could be ruled out 2. High insulin levels >100uIU/ml during documented hypoglycaemic episodes should initiate

⁷⁴ workup for autoimmune etiology as insulin is rarely above 100uIU/ml in insulinomas. 3. Spontaneous remission

is seen in up to 80% of patients. 4. Most patients respond to steroids. 5. Plasmapheresis is a second line therapy
which is beneficial in many patients, although the effect may be delayed.

- [Yamada et al.], T Yamada, J Imai, Y Ishigaki, Y Hinokio, Y Oka, H Katagiri. of HLA-DRB1*0403.
 (Possible Relevance)
- ⁷⁹ [Lupsa et al. ()] 'Autoimmune Forms of Hypoglycemia'. B Lupsa , A Chong , E Cochran , M Soos , R Semple ,
 ⁸⁰ P Gorden . *Medicine* 2009. 88 (3) p. .
- [Uchigata et al. ()] 'Drug-induced insulin autoimmune syndrome'. Y Uchigata , Y Hirata , Y Iwamoto . Diabetes
 Research and Clinical Practice 2009. 83 (1) p. .
- [Redmon et al. ()] Endocrinology and Metabolism Clinics of North America, J Redmon, F Nuttall, Autoimmune
 , Hypoglycemia . 1999. 28 p. .
- 85 [Haplotype in Insulin Autoimmune Syndrome Induced by -Lipoic Acid, Used as a Dietary Supplement Diabetes Care ()]
- 'Haplotype in Insulin Autoimmune Syndrome Induced by -Lipoic Acid, Used as a Dietary Supplement'.
 Diabetes Care 2007. 30 (12) p. .
- [Paiva et al. ()] 'Insulin Autoimmune Syndrome (Hirata Disease) as Differential Diagnosis in Patients with
 Hyperinsulinemic Hypoglycemia'. E Paiva , A Pereira , M Lombardi , S Nishida , T Tachibana , C Ferrer .
- Pancreas 2006. 32 (4) p. .
- 91 [Y U (1999)] 'Insulin Autoimmune syndrome (IAS, Hirata disease)'. Y U , Y . Ann Med Interne 1999 Apr. 150
 92 (3) p. .
- ⁹³ [Lidar et al. ()] Insulin autoimmune syndrome after therapy with imipenem. Diabetes Care, M Lidar, R Rachmani
 ⁹⁴ , E Half , M Ravid . 1999. 22 p. .
- ⁹⁵ [Ishida et al. ()] Lipoic Acid and Insulin Autoimmune Syndrome. Diabetes Care, Y Ishida , T Ohara , Y Okuno
 ⁹⁶ , T Ito , Y Hirota , K Furukawa . 2007. 30 p. .
- 97 [Yu et al. ()] 'Rituximab Selectively Suppresses Specific Islet Antibodies'. L Yu , K Herold , H Krause-Steinrauf
 98 , P Mcgee , B Bundy , A Pugliese . *Diabetes* 2011. 60 (10) p. .
- 99 [Yaturu et al. ()] 'Severe autoimmune hypoglycemia with insulin antibodies necessitating plasmapheresis'. S
 100 Yaturu , C Deprisco , A Lurie . *Endocrine Practice* 2004. 10 (1) p. .