

# Journal of Diabetes and Endocrinology

Volume 6 Number 1, January 2015

ISSN 2006-9871



*Academic  
Journals*

## ABOUT JDE

The **Journal of Diabetes and Endocrinology (JDE)** is published monthly (one volume per year) by Academic Journals.

**Journal of Diabetes and Endocrinology (JDE)** is an open access journal that provides rapid publication (monthly) of articles in all areas of the subject such as steroid hormones, clinical chemistry and biochemistry, neuroendocrinology, hypoglycemia in diabetes etc.

The Journal welcomes the submission of manuscripts that meet the general criteria of significance and scientific excellence. Papers will be published shortly after acceptance. All articles published in JDE are peer-reviewed.

## Submission of Manuscript

Submit manuscripts as e-mail attachment to the Editorial Office at: [jde@academicjournals.org](mailto:jde@academicjournals.org). A manuscript number will be mailed to the corresponding author shortly after submission.

The Journal of Diabetes and Endocrinology will only accept manuscripts submitted as e-mail attachments.

Please read the **Instructions for Authors** before submitting your manuscript. The manuscript files should be given the last name of the first author.

## Editors

**Prof. Masayoshi Yamaguchi**

*Division of Endocrinology and Metabolism and Lipids,  
Department of Medicine,  
Emory University School of Medicine,  
1639 Pierce Drive, 1305 WMRB, Atlanta,  
Georgia 30322-0001,  
USA.*

**Dr. Krishna M. Boini**

*Department of Pharmacology and Toxicology,  
Virginia Commonwealth University,  
1220 East Broad Street, MSB II, Room # 3054  
Richmond, VA - 23298.  
USA*

**Dr. Mohamed A. M. Mohamed Dkhil Hamad**

*Zoology Dept. College of Science,  
King Saud University,  
Egypt.*

**Dr. Aliyu Mohammed**

*Department of Human Physiology  
Ahmadu Bello University, Zaria.  
Nigeria*

**Prof. Bhupen Chandra Behera**

*Gayatri College of Pharmacy,  
Gayatri Vihar, Jamadarpali,  
Sambalpur- 768200, Orissa  
India.*

**Dr. Srinivas Nammi**

*Research Academic  
Faculty of Pharmacy  
The University of Sydney  
NSW 2006,  
Australia.*

**Dr. Mamdouh Moawad Ali Hassan**

*Biochemistry Department  
Genetic Engineering & Biotechnology Division  
National Research Center  
El Tahrir St., El Dokki 12622  
Cairo,  
Egypt.*

## Editorial Board

**Dr. Fawad Javed**

*Karolinska Institutet,  
Department of Dental Medicine,  
Huddinge,  
Sweden*

**Prof. Hab Lidia Rudnicka**

*Dept. Dermatology,  
Central Clinical Hospital,  
MSWiA, Warsaw,  
Poland*

**Dr. Aamir (Amer) Jalal Al Mosawi**

*University of Baghdad  
College of Medicine  
Iraq*

**Dr. Abbasi**

*Tehran University of Medical Sciences,  
Tehran,  
Iran*

**Dr. Rehab Fawzy Abdel-Rahman**

*Department of Pharmacology,  
National Research Centre (NRC),  
El- Tahrir St. Dokki, Cairo,  
Egypt.*

**Dr. JIMOH, Ahmed Kayode**

*Titilayo Street, Isale Afo,  
Ikirun Ifelodun LGA  
Kwara State  
Nigeria*

**Dr. Saganuwan Alhaji Saganuwan**

*Department of Veterinary Physiology,  
Pharmacology & Biochemistry,  
University of Agriculture Benue State,  
Nigeria.*

**Dr. Alireza Shirpoor**

*Permanent lecturer of  
Urmia University of Medical Sciences  
Iran*

**Dr. Bassim Atta**

*Prof. Food Chemistry & Analysis  
Faculty of Agriculture  
Tanta University  
Egypt*

**Prof. (Dr.) Bhupen Chandra Behera**

*Gayatri College of Pharmacy,  
Gayatri Vihar, Jamadarpali,  
Sambalpur- 768200, Orissa  
India*

**Dr. Ben-zhi Cai**

*Baojian Road 157#, Harbin,  
150081,  
China.*

**Dr. Chatchalit Rattarasarn**

*Division of Endocrinology and Metabolism  
Department of Medicine  
Ramathibodi hospital, Mahidol university  
Bangkok,  
Thailand 10400*

**Dr. Sónia Catarina Correia**

*Centre for Neuroscience and Cell Biology,  
Department of Zoology,  
University of Coimbra,  
3004-517 Coimbra  
Portugal*

**Mohamed A. M. M. Dkhil Hamad**

*faculty of science,  
Helwan University,  
Egypt.*

**Dr. Bakari Adamu Girei**

*Department of Medicine,  
Ahmadu Bello University  
Zaria,  
Nigeria*

**Dr. Aliyu Mohammed**

*Adamawa State,  
Nigeria*

**Ana Isabel Marques Duarte**

*Center for Neuroscience & Cell Biology  
Institute of Biochemistry, Faculty of Medicine (Pólo I)  
University of Coimbra  
3004-504 Coimbra  
Portugal*

# Journal of Diabetes and Endocrinology

Table of Content: Volume 6 Number 1 January 2015

## ARTICLES

### Research Articles

- Pulmonary artery hypertension and type 1 diabetes mellitus with suspected autoimmune polyendocrine syndrome in a pediatric patient** 1  
Ildiko H. Koves and Delphine Yung

*Case Report*

# Pulmonary artery hypertension and type 1 diabetes mellitus with suspected autoimmune polyendocrine syndrome in a pediatric patient

Ildiko H. Koves<sup>1\*</sup> and Delphine Yung<sup>2</sup>

<sup>1</sup>Division of Endocrinology and Diabetes, Seattle Children's Hospital, Seattle WA 98040, USA.

<sup>2</sup>Division of Cardiology, Seattle Children's Hospital, 4800 Sand Point Way NE, USA.

Received 10 November, 2014; Accepted 24 December, 2014

The objective of this study was to describe a constellation of rare pediatric disorders, pulmonary arterial hypertension (PAH) and autoimmune polyendocrine syndromes (APS). This study presents a brief report of a child diagnosed with autoimmune type 1 diabetes mellitus (T1DM) at age 11 and with PAH three years later, when he was re-presented with symptoms of chest pain, lethargy, syncope and vomiting. Following a saline bolus, he became severely hypoxic. Echocardiogram and cardiac catheterization confirmed PAH. Following a literature search of these two concomitant rare conditions, suspicion was raised of a uniform diagnosis of APS; autoimmune regulator (AIRE) gene analysis revealed a heterozygous c1203T>C (p.P401P) mutation on chromosome 21q22. Screening for other autoimmune involvement was negative thus far. Pulmonary AH should be included within the rare components of APS as the independent occurrence of these two rare disorders is highly unlikely in particular in the context of an identified mutation within the AIRE gene.

**Key words:** Autoimmune polyglandular syndrome, pulmonary arterial hypertension, type 1 diabetes mellitus, autoimmune regulator gene, polyglandular endocrinopathy.

## INTRODUCTION

Autoimmune polyendocrine syndromes are rare. There have been approximately 500 patients worldwide. The highest prevalence was found among the Iranian Jewish community (1:9,000), in Sardinia (1:14,400) and in Finland (1:25,000) (Weiler et al., 2012). It is characterized by multiple autoimmune illnesses, most typically Addison disease, hypoparathyroidism, and/or chronic mucocutaneous candidiasis. Type 1 diabetes mellitus

(DM) is infrequent, present in 18% of cases (Eisenbarth and Gottlieb, 2004). Type 1 APS (OMIM#240300) is caused by homozygous, compound heterozygous, or heterozygous mutation in the autoimmune regulator gene (AIRE; 607358) on chromosome 21q22, which encodes a transcription factor.

Pulmonary arterial hypertension is a rare disorder associated with autoimmune illnesses and should be

\*Corresponding author. E-mail: [ildiko.koves@seattlechildrens.org](mailto:ildiko.koves@seattlechildrens.org). Tel: + 1 206 987 5037. Fax: +1 206 987 2720.

Author(s) agree that this article remain permanently open access under the terms of the [Creative Commons Attribution License 4.0 International License](http://creativecommons.org/licenses/by/4.0/)

considered as an autoimmune association. This case report broadens the currently recognized autoimmune components of APS.

## CASE REPORT

An 11 year old male, presented with symptoms consistent with type 1 diabetes mellitus (T1DM) of vomiting, weight loss, lethargy and hyperglycemia. He lived at an altitude of 1600 m in Arvada, Colorado, and three years later he represented chest pain and syncope during vomiting. Echocardiogram suggested pulmonary hypertension. Cardiac catheterization confirmed PAH with mean pulmonary arterial pressure of 77 mmHg in room air, which decreased to 32 mmHg on acute vasodilator testing with oxygen and inhaled nitric oxygen. Due to the severity of his presentation, he was initially managed on intravenous epoprostenol and oral sildenafil. Due to the side effects, he was weaned off intravenous epoprostenol, and started on diltiazem as repeat catheterization showed acute pulmonary vasoreactivity. Nineteen months later, he relocated to Seattle to live at lower altitude.

He is of mixed ancestry; mother of mixed German, Irish and father of African-American background. There was no consanguinity, separated parents. The family history was significant for aunts on both parental side with systemic lupus erythematosus and both maternal grandparents with autoimmune thyroid disease. A half brother on the paternal side was diagnosed with PAH at 9 years and passed away at 14 years of age.

He was born at term with no other significant illnesses. Following T1DM, PAH was diagnosed as possibly associated with hereditary component, living at high altitude and/or autoimmune disease. At presentation to Seattle, height was 172.8 cm (46th), weight 59 kg (46th), body mass index (BMI) 20 (42nd), blood pressure 123/66 mmHg, pulse 66 min<sup>-1</sup>, respiratory rate 18 min<sup>-1</sup>, pulse oximetry in room air 100%. Salient physical examination finding: loud P2, otherwise negative examination.

The initial diagnostic evaluation is summarized as shown in Table 1. He had poor metabolic control of diabetes (HbA1C: 12.1%) due to poor adherence with insulin therapy. Laboratory screening for other autoimmune involvement (thyroid, parathyroid) were negative. Clinically, he had no suggestion of Addison's disease or other mucocutaneous involvement. He continued on basal bolus subcutaneous insulin therapy for T1DM. Fifteen months after relocation to sea level, cardiac catheterization demonstrated almost normal mean pulmonary pressure (23 mmHg) and further decrease with oxygen and inhaled nitric oxide to 19 mmHg. To improve adherence, he is now on tadalafil therapy, 20 mg daily and amlodipine 5 mg twice a day.

He receives regular follow up and clinical screening for other potential associated autoimmune illnesses and

complications of diabetes. Pulmonary pressure and cardiac function remain stable on the aforementioned therapy.

## Genetic assessment

Genetic testing was pursued following literature review and another case identification (Alghamdi et al., 2010). A novel heterozygous mutation was identified in the AIRE gene (21q22.3) at c1203T>C (p.P401P) using genomic DNA sequence analysis with PCR amplification of exons1-14 with automated fluorescence dideoxy sequencing method (DNA Diagnostic Laboratory, Baylor College of Medicine, Huston).

## DISCUSSION

Several recent novel mutations were identified from Scandinavian (Wolff et al., 2007), Arabic (Faiyaz-Ul-Haque et al., 2009), Polish (Stolarski et al., 2006), Slovenian (Podkrajsek et al., 2005) and Indian (Zaidi et al., 2009) populations with racial differences affecting the AIRE gene (gene locus MIM#607358). This patient had a mixed racial northern European and African-American background and his mutation may relate to the latter ancestry. Some APSs have milder phenotypes and both the clinical features and AIRE mutations may be more diverse than previously thought (www.omim.org). Defects in this gene cause the rare autosomal-recessive systemic autoimmune disease known as type 1 APS syndrome previously also termed as autoimmune polyendocrinopathy-candidiasis-ectodermal dystrophy (APECED).

To our knowledge this is the youngest patient described with the two rare disorders and an identifiable AIRE gene mutation. There are a few young adult case reports (Alghamdi et al., 2010; Barrou et al., 1989; Garcia-Hernandez et al., 2006) with the constellation of PAH, T1DM and APS. The timing of autoimmune component of their presentation varied with recognized morbidities from the early 20s to later diagnosis of PAH in their 50s. Patients with type 1 APS with any two of specific autoimmune conditions: mucocutaneous candidiasis, Addison's disease, T1DM or hypoparathyroidism, almost always have AIRE mutations. Recognized AIRE gene mutations cause different autoimmune illnesses including hypothyroidism, pernicious anemia, alopecia, vitiligo, hepatitis, ovarian atrophy, keratitis, Graves disease, myasthenia gravis (McAlpine and Thomson, 1988), enterochromaffin cells hormone secretion leading to malabsorption (Hogenauer et al., 2001).

Our patient without Addison's disease, hypoparathyroidism, enteropathy, evidence of hypogonadism, pernicious anemia, atrophic gastritis and/or mucocutaneous candidiasis did not fit the typical type 1 APS, yet an AIRE

**Table 1.** Diagnostic evaluation.

T1DM and APS 1 (Reference)	PAH
HBA1C: 10-14% (<6%)	Echocardiogram: Severe septal wall flattening; Estimated RV systolic pressure: 115 mmHg
C-peptide < 0.1 ng/ml (0.8-3.1)	Cardiac catheterization: PA pressure; Mean: 77 mmHg
GAD: 93 (<25)	Presentation O <sub>2</sub> sats: 80s
Insulin antibody: 0.143 (<0.013)	Spirometry: FVC: 3.34 L (76% predicted); FEV1: 2.8 (75%); FEV1: FVC 83.8%; TLC: 4.01 L
ICA512A: 0 (<7)	Lung ventilation perfusion scan: normal
TSH: 2.81 mIU/ml (0.5-4.5)	-
Free T4: 1.3 ng/dl (0.8-2)	-
Antithyroglobulin Ab: neg	-
Anti TPO: neg	-
TSH receptor Ab: <1 IU/L (<1.75)	-
PTH related peptide: 0.5 pmol/ml (<2)	-
Total Ca: 9.4 mg/dl (8.7-10.7)	-
ACTH: 13 pg/ml (10-60)	-

HBA1C: Hemoglobin A1C; GAD: Glutamic acid decarboxylase antibody; ICA512: Islet cell antibody 512a; TSH: thyroid stimulating hormone; Free t4: Free thyroxine4; PTH: parathyroid hormone; Ab: antibody; AntiTPO: anti thyroid peroxidase antibody; ACTH: Adrenocorticotrophic hormone; RV BP: Right ventricle blood pressure; PA: pulmonary artery pressure; FVC: forced vital capacity; FEV1: forced expiratory volume in 1 second; TLC: total lung capacity.

AIRE gene mutation was found. A similar scenario has been described by Bhansali et al. (2003) whereby an adolescent with the specific association of PAH and type 1 APS had no clinical evidence of mucocutaneous candidiasis and an adult with fatal PAH had previously recognized type 1 APS (Korniszewski et al., 2003). This case fatality underscores the need to consider PAH in all children with APS.

Pulmonary arterial hypertension is characterized by obliteration of the small vasculature of pulmonary arteries. It is a rare, inhomogeneous disorder and a subset of patients have associated autoimmune morbidity such as Hashimoto thyroiditis, systemic lupus erythematosus, rheumatoid arthritis, mixed connective tissue disease, scleroderma and T1DM (Morse et al., 1992; Badesch et al., 2010).

Type 2 APS and PAH have also been described in an adolescent who developed Hashimoto thyroiditis and T1DM following PAH diagnosis (Alghamdi et al., 2010) and in adults (Garcia-Hernandez et al., 2006) and no AIRE gene mutation was found. This further broadens the clinical and genetic variations and perhaps autoimmune polyendocrinopathy should be considered as a spectrum disorder with phenotypic and genotypic overlap.

## Conclusion

To our knowledge, this is the youngest reported patient with the constellation of these two rare disorders with a novel mutation in the AIRE gene region on chromosome 21q22.3.

In the authors opinion, PAH should be included within

the rare components of APS as the independent occurrence of these two rare disorders is highly unlikely in particular in the context of an identified mutation within the AIRE gene region. More cases are needed to indicate that the concurrence is a true association.

## Conflict of interest

The authors have no relevant conflict of interest to disclose.

## ACKNOWLEDGEMENTS

The authors acknowledge the Denver Children's Hospital Pulmonary Hypertension and Diabetes team for this patient's initial management and Dr. Jerry Zimmerman, Critical Care, Seattle Children's, for providing advice.

## REFERENCES

- Alghamdi MH, Steinrath M, Panagiotopoulos C, Potts JE, Sandor GG (2010). Primary pulmonary arterial hypertension and autoimmune polyendocrine syndrome in a pediatric patient. *Pediatr. Cardiol.* 31(6):872-874.
- Badesch DB, Raskob GE, Elliott CG, Krichman AM, Farber HW, Frost AE, Barst RJ, Benza RL, Liou TG, Turner M, Giles S, Feldkircher K, Miller DP, McGoon MD (2010). Pulmonary arterial hypertension: baseline characteristics from the REVEAL Registry. *Chest* 137(2):376-387.
- Barrou Z, Pehuet-Figoni M, Weber S, Lockhart A, Luton JP (1989). Polyendocrinopathy combined with primary pulmonary arterial hypertension. *Presse Med.* 18(19):963-965.
- Bhansali A, Kotwal N, Suresh V, Murlidharan R, Chattopadhyay A, Mathur K (2003). Polyglandular autoimmune syndrome type 1 without



- chronic mucocutaneous candidiasis in a 16 year-old male. *J. Pediatr. Endocrinol. Metab.* 16(1):103-105.
- Eisenbarth GS, Gottlieb PA (2004). Autoimmune polyendocrine syndromes. *N. Engl. J. Med.* 350(20):2068-2079.
- Faiyaz-UI-Haque M, Bin-Abbas B, Al-Abdullatif A, Abdullah Abalkhail H, Toulimat M, Al-Gazlan S, Almutawa AM, Al-Sagheir A, Peltekova I, Al-Dayel F, Zaidi SH (2009). Novel and recurrent mutations in the AIRE gene of autoimmune polyendocrinopathy syndrome type 1 (APS1) patients. *Clin. Genet.* 76(5):431-440.
- Garcia-Hernandez FJ, Ocana-Medina C, Gonzalez-Leon R, Garrido-Rasco R, Sanchez-Roman J (2006). Autoimmune polyglandular syndrome and pulmonary arterial hypertension. *Eur. Respir. J.* 27(3):657-658.
- Högenauer C, Meyer RL, Netto GJ, Bell D, Little KH, Ferries L, Santa Ana CA, Porter JL, Fordtran JS (2001). Malabsorption due to cholecystokinin deficiency in a patient with autoimmune polyglandular syndrome type I. *N. Engl. J. Med.* 344(4):270-274.
- Korniszewski L, Kurzyna M, Stolarski B, Torbicki A, Smerdel A, Płoski R (2003). Fatal primary pulmonary hypertension in a 30-yr-old female with APECED syndrome. *Eur. Respir. J.* 22(4):709-711.
- McAlpine JK, Thomson JE (1988). Myasthenia gravis and Schmidt syndrome. *Postgrad. Med. J.* 64(756):787-788.
- Morse JH, Barst RJ, Fotino M (1992). Familial pulmonary hypertension: immunogenetic findings in four Caucasian kindreds. *Am. Rev. Respir. Dis.* 145(4 Pt 1):787-792.
- Podkrajsek KT, Bratanic N, Krzisznik C, Battelino T (2005). Autoimmune regulator-1 messenger ribonucleic acid analysis in a novel intronic mutation and two additional novel AIRE gene mutations in a cohort of autoimmune polyendocrinopathy-candidiasis-ectodermal dystrophy patients. *J. Clin. Endocrinol. Metab.* 90(8):4930-4935.
- Stolarski B, Pronicka E, Korniszewski L, Pollak A, Kostrzewa G, Rowińska E, Włodarski P, Skórka A, Gremida M, Krajewski P, Płoski R (2006). Molecular background of polyendocrinopathy-candidiasis-ectodermal dystrophy syndrome in a Polish population: novel AIRE mutations and an estimate of disease prevalence. *Clin. Genet.* 70(4):348-354.
- Weiler FG, Dias-da-Silva MR, Lazaretti-Castro M (2012). Autoimmune polyendocrine syndrome type 1: case report and review of literature. *Arq. Bras. Endocrinol. Metabol.* 56(1):54-66.
- Wolff AS, Erichsen MM, Meager A, Magitta NF, Myhre AG, Bollerslev J, Fougner KJ, Lima K, Knappskog PM, Husebye ES (2007). Autoimmune polyendocrine syndrome type 1 in Norway: phenotypic variation, autoantibodies, and novel mutations in the autoimmune regulator gene. *J. Clin. Endocrinol. Metab.* 92(2):595-603.
- Zaidi G, Sahu RP, Zhang L, George G, Bhavani N, Shah N, Bhatia V, Bhansali A, Jevalikar G, Jayakumar RV, Eisenbarth GS, Bhatia E (2009). Two novel AIRE mutations in autoimmune polyendocrinopathy-candidiasis-ectodermal dystrophy (APECED) among Indians. *Clin. Genet.* 76(5):441-448.

# Journal of Diabetes and Endocrinology

## Related Journals Published by Academic Journals

- *African Journal of Pharmacy and Pharmacology*
- *Journal of Dentistry and Oral Hygiene*
- *International Journal of Nursing and Midwifery*
- *Clinical Reviews and Opinions*
- *Journal of AIDS and HIV Research*
- *International Journal of Nutrition and Metabolism*

**academicJournals**