Diabetes and Atypical Autoimmune Polyglandular Syndrome Type 3: A Paediatric Case Report

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ABSTRACT

Objective: Type 1 Diabetes (T1DM) is an autoimmune disease. Children affected by T1DM are at higher risk of other autoimmune comorbidities and the combination of two or more organ specific disorders should define the diagnosis of Autoimmune Polyglandular Syndrome (APLS).

Case Report: Here we report the case of a young girl affected by an atypical APLS type 3. She had a diagnosis of T1DM at 4 years old, celiac disease at 6 years and autoimmune thyroiditis at 7 years. Her personal history was complicated by necrotizing glomerulonephritis ANCA-associated with important effects on glycaemic control and other autoimmune comorbidities.

Conclusion: we analyse a multifaceted case, in a very young girl, characterized by the coexistence of different autoimmune diseases influencing endocrine functions and metabolic control, during the different phases of a complex therapy.

Introduction

Type 1 Diabetes (T1DM) is an autoimmune, multifactorial, disease caused by beta cells destruction induced by immune response [1]. The pathophysiology of T1DM is still unclear but genetic predisposition is a landmark of this condition. HLA and non-HLA genes are involved in the risk of autoimmunity onset and many of these genes are shared by different autoimmune settings; thus, children affected by T1DM are at increased risk of other autoimmune comorbidities [2,3]. In particular, T1DM could be part of autoimmune polyglandular syndrome cluster.

In 1980 Neufled and Blizzard proposed a classification of the autoimmune polyendocrine syndromes (APS) Table 1 [4]. These conditions are characterized by lack of self-tolerance and by induction of immune responses directed against self-structures with the coexistence, in the same patient, of more than one autoimmune disease [5]. The term polyendocrine is not completely appropriate, because several non-endocrine scenarios could be part of the clinical picture of these autoimmune syndromes [5,6]. Here, we report the case of a girl affected by APS type 3 and the course of the different diseases’ diagnosis in her young life.

Table 1: Neufled and Blizzard’s classification of APS [4].

| Classification of APS | Description |
|-----------------------|-------------|
| Type 1                | Chronic Candidiasis, Chronic Hypoparathyroidism, Addison’s disease (at least two present) |
| Type 2                | Addison’s disease (always present) and thyroid autoimmune diseases and/or Type 1 Diabetes |
| Type 3                | Thyroid autoimmune diseases associated with other autoimmune condition (excluding Addison’s disease and/or hypoparathyroidism) |
| Type 4                | Combination of organ specific autoimmune diseases non included in previous groups |
Case Report

MF, female, was born at 38 weeks by vaginal delivery. Pregnancy was complicated by placenta disruption at 18 weeks. Birth weight was appropriate for gestational age. Perinatal period was uncomplicated. Familial history was negative for particular medical condition and consanguinity. At 4 years of age, MF had diagnosis of T1DM complicated by ketoacidosis (pH 7.2, HCO3- 11.7 mmol/L, HbA1c163 mmol/mol). Multi-daily injections therapy, according to basal bolus insulin regimen, was started with a good metabolic control. At 6 years of age, MF complained diarrhea and abdominal pain. Celiac screening was positive; endoscopy confirmed the diagnosis of celiac disease with type 3 histology according Marsh-Oberhuber classification. Gluten Free Diet (GFD) was then switched on with a good compliance.A transient phase of hyperthyroidism was recorded when MF was 7 years old: TSH was 0.007 mcUI/mL, FT4 2.3 ng/dL, anti-thyroglobulin antibodies (ATG) were positive 211.4 UI/mL (normal value < 29), thyrotropin receptor antibodies (TrAbs) and anti-thyroid peroxidase antibodies (ATP) were negative. Ultrasonography showed signs of thyroiditis. A new autoimmune diagnosis was made autoimmune thyroiditis. She didn’t need treatment until 8 years of age, when MF presented clinical signs of hypothyroidism confirmed by laboratory findings (TSH 21.93 mcUI/mL, FT4 0.65 ng/dL) and so levothyroxine was started with at an initial dose of 0.5 mcg/kg/die. She had a second important symptomatic phase of hyperthyroidism at age of 10 years, under thyroxine therapy, and beta blocker drug (propranolol) was used. Blood examinations confirmed TrAbs negativity with TSH 0.02 mcUI/mL and FT4 3.09 ng/dL. Thyroxine treatment was stopped for few weeks. After few months, MF complained arthralgia and swelling of the hands with clinical signs of arthritis. Laboratory evaluation showed a slight increase of Anti-Nuclear Antibodies (ANA 1:80, homogeneous pattern) with negativity of inflammation indexes. Articular ultrasound was performed with normal imaging. HLA typing was HLA DRB1 *04 *16, HLA-A *03 *30, HLA-B *14 *49, HLA-C *07 *08, DQB1 *03 *05, HLA-DQA1 *01:02, HLA-DQA1 *03:03.

Treatment with Nonsteroidal anti-inflammatory drug was started for 3 weeks with good clinical response, but the symptomatology relapsed, after the withdrawal. A new complete evaluation was performed at Paediatric Rheumatologic Hospital with diagnosis of “Transient arthralgia and subcutaneous edema” in spite of an increased level of ANA titre (ANA 1:320, homogeneous pattern). During the follow up period, MF showed also persistent microalbuminuria although good glycaemic control. Proteinuria in 24 hours was normal in repeated controls, until the appearance of pathologic levels (800 mg/24h) deserving further investigations. The new findings were positivity for anti-neutrophil cytoplasmic antibodies (ANCA): subtype anti Myeloperoxidase (MPO), anti-proteinase 3 negative. In May 2016 renal biopsy was performed and histology defined the diagnosis of necrotizing glomerulonephritis ANCA-associated. Optic microscopy showed focal and segmental mesangial hypercellularity, one small area of fibrinoid necrosis and one cellular crescent. Tubules and interstitium were normal. Immunofluorescence showed IgG, IgM and C3 mesangial and parietal traces. The diagnosis was renal-limited vasculitis; MF was 11 years old.

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Figure 1: Thyroid autoimmunity course during the follow up (ATG anti thyroglobulin antibodies, ATPO anti thyroid peroxidase antibodies, x axis: timing): * Autoimmune Thyroiditis diagnosis, + arthralgia, # Glomerulonephritis diagnosis.
Discussion

Type 1 Diabetes is an autoimmune disease in which genetic and environmental factors interact to define not only determinism of its pathophysiology but also the susceptibility to other autoimmune conditions [3].

Often the cluster of autoimmune diseases could be present in the same person and so the diagnosis of autoimmune polyglandular syndrome could be made [9]. In our patient, Autoimmune Polyglandular Syndrome type 3 was diagnosed, but there are some concerns about. Neufled and Blizzard classification of APS was completed by Betterle in 2002 (Table 2); in particular, according to the latter, our clinical case could not be placed in a defined condition: MF had autoimmune diabetes, autoimmune thyroiditis, celiac disease and vasculitis, so she could be diagnosed as APS type 3a, 3b or 3d [4,10]. Eisenbarth et al in 2004 underlined the usefulness of a simpler classification for these autoimmune conditions based on genetic background, age at onset and clinical settings to avoid mistakes and doubts [5]. However, beyond all type of classification, our patient’s history was characterized by an uncommon association in her clinical picture.

Table 2: Type 3 APS according to Betterle classification: the diagnosis requires the presence of autoimmune thyroid disease with other autoimmune conditions [9].

| Endocrine Glands | Gastrointestinal Tract and Liver | Skin, Muscles, Nervous system, Hematologic system | Autoimmune Rheumatic Disorders Vasculitis |
|------------------|---------------------------------|------------------------------------------------|----------------------------------------|
| Type 1 DM Hirata’s syndrome Adeno-hypophysitis Neuro-hypophysitis Chronic hypoparathyroidism | Chronic atrophic gastritis Ulcerative colitis Pernicious anemia Celiac Disease Other inflammatory bowel syndrome Primary biliary cirrhosis Autoimmune hepatitis Sclerosing cholangiatis Autoimmune pancreatitis | Vitiligo Alopecia Bullous Skin diseases Myasthenia gravis Stiff Person Syndrome Multiple Sclerosis Autoimmune anemia Autoimmune thrombocytopenia | Systemic Lupus Erythematosus Discoid Lupus Erythematous Rheumatoid arthritis Mixed connective tissue disease Sero-negative arthritis Systemic sclerosis Dermato/polymyositis Vasculitis |
| 3 A               | 3 B                             | 3 C                                             | 3 D                                   |

ANCA have been detected more frequently in patients with a history of certain autoimmune diseases. Such an association may simply reflect an underlying tendency to immune system dysfunction in these patients and the finding of positive ANCA serology does not reliably herald the development of ANCA-associated vasculitis. In our patient ANCA was searched in the autoimmune panel after the detection of pathologic proteinuria in urine, despite of a good glycaemic control, and correlate with histologic findings at renal biopsy. Few cases of ANCA vasculitis in autoimmune polyendocrinopathy syndrome have been described and this is the first case of renal vasculitis in a patient affected by Autoimmune Polyglandular Syndrome type 3, to our knowledge. Recent evidences underline the importance of HLA genotype in the susceptibility to MPA: gene variants appear to alter immune cell functionality, becoming key factors in the pathogenesis of ANCA-associated vasculitis [11]. In particular, the patient HLA genotype could explain her susceptibility to ANCA associated vasculitis [7]. MF is now adolescent and her clinical history is, probably similar to an iceberg in which we can observe only the visible part, we don’t know the submerged one at this time and we don’t know what else could happen in her next future.

In particular, one important teaching from this case is the influence and complex interactions of autoimmunity on endocrine functionality. At first, the strange instability of thyroid function was one prominent feature in our patient. Transient phases of hypothyroidism and hyperthyroidism were recorded: each phase was accompanied by modification in thyroid autoimmunity subset, as showed in Figure 1, and every phase of hypo/hyperthyroidism was preceded by hyper/hypothyroidism, as to indicate an autoimmune attack to the gland. Probably, the autoimmune system, the therapies used, and the stress could be the reading key of these complex immune-endocrine interactions. These might lead to the lack of recovery equilibrium typical of autoimmune thyroiditis course [12]. The effects of autoimmune comorbidities on glycaemic control has also been experimented: hypoglycaemia and hyperglycaemia were clearly complications observed in our patient management [13]. Last but not least, moreover, this lesson: Nephropathy in Diabetic Children is not only «Diabetic». Renal biopsy played a fundamental role in the diagnosis, prognosis and in our therapeutic decisions [14]. Our case report highlights the importance of a strict clinical and laboratory monitoring in these complex pediatric patients, in a multidisciplinary approach, giving the right attention to the appearance of new signs or symptoms, even if they could be related to the first diagnosis.

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Conflict of Interests

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