**A novel variant in ATM gene causes ataxia telangiectasia revealed by whole-exome sequencing**

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**ABSTRACT**

A novel variant in ATM gene (p.Gln1982*) revealed by whole-exome sequencing in a 9-year old boy. He presented with recurrent upper respiratory tract infections, failure to thrive, developmental delay, ataxic gait, and bulbar telangiectasia.

**Case Report. Patient information.** The proband is a 9-year old Saudi boy who attends the pediatric clinic since the age of 3 years with developmental delay and ataxia. He is a product of full-term pregnancy, born by spontaneous vertex delivery with birth weight of 2.60 kg at the 10th centile and a length of 45 cm below the third centile. His parents were first-degree cousins, and he had 2 siblings who died with undiagnosed developmental delay, unsteady gait, and cerebral atrophy. The proband was referred at the age of 8 years to our immunology clinic for further evaluation of recurrent upper respiratory tract infections. He used to get approximately 6-7 episodes a year of upper respiratory tract infections, otitis media and pharyngitis, mostly needed parenteral or oral antibiotics. His response to...
antibiotics was generally slow and occasionally needed a prolonged or a second course of antibiotics. He has no history of chronic diarrhea or skin infection.

**Clinical findings.** His developmental evaluation showed that he was 4–5 years delayed in the intellectual, speech and motor domains. His weight was 17 kg and height was 110 cm, both were below the third centile. Clinical examination revealed ocular telangiectasia and no evidence of skin lesions, lymphadenopathy or hepatosplenomegaly. Neurological examination showed intact cranial nerves, proximal muscles weakness, normal reflexes, and tone. He has impaired cerebellar signs including finger-nose test and wide ataxic gait.

Diagnostic assessment revealed normal while blood count, lymphocyte count, and peripheral blood smear. Serum immunoglobulins analysis showed a normal IgM level at 1.26 g/L (normal=0.38–2.35), IgG level at 10.2 g/L (normal=6.6–16.2) and low IgA=<0.25g/L (normal=0.57–3.18). Brain MRI showed non-specific mild atrophic changes within the cerebellar hemispheres.

**Molecular genetic analysis.** Informed consent was obtained from the parents. The genomic DNA was isolated from the peripheral blood leukocytes of the proband and his parents. Approximately 37 Mb (214,405 exons) of the Consensus Coding Sequences (CCS) were enriched from fragmented genomic DNA by >340,000 probes designed against the human genome (Nextera Rapid Capture Exome, Illumina). The generated library sequenced on an Illumina platform to an average coverage depth 70-100X. An end to end inhouse bioinformatics pipelines in CentoGene laboratory, including base calling, primary filtering of low-quality reads and probable artifacts, and annotation of variants was applied. All disease-causing variants reported in HGMD®, in ClinVar or CentoMD® (class 1) as well as all variants with minor allele frequency (MAF) of less than 1% in ExAC database were considered. Evaluation is focused on exons and intron boundaries +/-20.

**Molecular genetics result.** Genomic DNA sequencing revealed a novel, homozygous variant in ATM gene. It is a nonsense substitution which interrupts the reading frame by a premature stop codon (c.5944 C>T, p.Gln1982*) (Figure 1). This variant has been detected in both parents in a heterozygous state. It is classified as likely pathogenic (class 2) according to the recommendations of the American College of Medical Genetics and Genomics (ACMG).

**Discussion.** Ataxia telangiectasia is a rare but an important cause of ataxia in childhood. Early diagnosis of this disorder helps in tailoring and mostly avoiding complications especially those related to impaired immunity. Ataxia is the classical presentation of A-T which usually begins during toddler age group manifested as impaired coordination. Likewise, children at school age with A-T experience difficulties with reading and writing. Furthermore, patients with A-T may develop a difficulty with involuntary movement at any age, which includes chorea, myoclonic jerks dystonia and other different types of tremors. Our patient followed the clinical path of A-T presenting with ataxia and recurrent upper respiratory tract infections; however, it took many years to be referred to the specialized pediatric immunology clinic to confirm the diagnosis.

Telangiectasia generally becomes apparent by the age of 5–8 years, and involves the bulbar conjunctivae

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**Table 1** - A novel variant in ATM gene causes ataxia telangiectasia revealed by whole-exome sequencing.

| Date       | Relevant past medical history and intervention                                                                 |
|------------|---------------------------------------------------------------------------------------------------------------|
| 16.05.2017 | 8 years old boy who is a product of full term pregnancy, referred to the immunology clinic to rule out any associated with primary immunodeficiency. Patient with history wide ataxic and failure to thrive. |
| Date       | Summaries initial and follow-up visits                                                                      | Diagnostic testing (including dates)                                                                 | Interventions           |
| 2015-2016  | During the whole years of 2016 patient have 6-7 episodes of upper respiratory tract infection                   | No exact date or specific test or other investigation that parent can recall (patient from outside Riyadh). | None                     |
| 2016       | History of diarrhea on and off in year 2016                                                                    | No exact or specific test or other investigation that parent can recall (patient from outside Riyadh)    | None                     |
| 26.04.2017 | History wide ataxic gait                                                                                       | MRI Brain shows mild atrophic changes within the hemispheres                                            | None (follow-up in the clinic) |
| 16.05.2017 | Because of history of recurrent URTI as past medical history                                                 | Blood sent for FBC and differential which showed normal WBCs count and normal diff                     | None                     |
|            | Blood immunoglobulin level: IgM=1.26g/l, IgG=10.2g/l and Ig A=<0.25g/l                                        | Serum immunoglobulin level: IgM=1.26g/l, IgG=10.2g/l and Ig A=<0.25g/l                                 |                          |
| 14.06.2017 | For the above history ataxia telangiectasia need to rolled out                                              | Blood sent or molecular genetics study which showed premature stop codon variant (c.5944C>T, p.Gln1982*) | Genetic counseling for the parents                   |

MRI - magnetic resonance imaging, FBC - Full Blood Count, IgA - immunoglobulin A, IgG - immunoglobulin G, IgM - immunoglobulin M, URTI - upper respiratory tract infection, WBC - White Blood Cell
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Figure 1 • Chromatogram sequences showing homozygous variance in the index patient and heterozygous in parents.

as in our patient. Occasionally telangiectasia can also affect other body areas which are exposed to the sun, especially the face and ears.

About 25% of patients with A-T will have pulmonary manifestations with either a cough or recurrent chest infection, or even bronchiectasis. Similarly, our patient had recurrent upper respiratory tract infection. Patients with A-T are at risk of developing malignancy, which occurs in 25% of all A-T patients during their lifetime. Lymphoma and leukemia are the most common types of malignancy in patients with A-T at age <16–20 years. Our patient is now under continuous surveillance regarding the risk of malignancy. Also, growth failure with short stature and poor weight gain are commonly reported in individuals with A-T as in our patient. The neuro-imaging studies are usually normal in early childhood years. However, as the disease progresses, the magnetic resonance imaging (MRI) studies are likely to show the pathological finding of diffuse and progressive cerebellar changes similar to those seen in our patient.

This study reports a novel variant in ATM gene which results in premature stop codon. Unlike the previously reported variants in ATM genes which were mainly diagnosed by Sanger sequencing, we opted to use whole exome sequencing (WES) in our patient. This is because WES is readily accessible and affordable to our institute compared to Sanger sequencing.

Like our patient, Stankovic et al reported that 71% of patients from the British Isles with A-T had stop codon variants leading to premature termination of the protein. Compared to our patient who presented with a slowly progressive course, other patients with truncated protein tend to have a severe phenotype with an early neurological deficits. On the contrary, other types of variants as missense or splice site tend to have milder phenotype with late onset of symptoms. Despite this observation regarding phenotype/genotype correlation in terms of type of the variants, a clinical heterogeneity among A-T patients remains the role with variable age of onset, severity of cerebellar involvement extent of immunodeficiency and predisposition to cancer. This clinical heterogeneity is illustrated by our patient who has a stop codon variant, yet he had a slowly progressive course with minimal neurological signs and apparent intact immunity and no evidence of malignancy so far.

In conclusion, this study identified a novel pathogenic stop codon variant in ATM gene in a child with the ataxia-telangiectasia. This report highlights the utility of the new generations of genetic testing in pinpointing the molecular etiology of undiagnosed genetic disorders, especially in consanguineous families that previously lost children with similar presentation.

References

1. Syllaba I, Henner K. Contribution a l’indépendance de l’athetose double idiopathique et congenitale. Rev Neurol 1926; 1: 541-62.
2. Boder E, Sedgwick RP. A familial syndrome of progressive cerebellar ataxia, Oculo-cutaneous telangiectasia and frequent pulmonary infection: a preliminary report on seven children, an autopsy, and a case history. Univ Southern Calif Med Bull 1957; 9: 15-28.
3. Nissenkorn A, Levy-Shraga Y, Banet-Levi Y, Lahad A, Sarouk I, Modan-Moses D. Endocrine abnormalities in ataxia telangiectasia: findings from a national cohort. Pediatr Res 2016; 79: 889-894.
4. Stankovic T, Kidd AM, Sutcliffe A, McGuire G M, Robinson P, Weber P, et al. ATM mutations and phenotypes in ataxia-telangiectasia families in the British Isles: expression of mutant ATM and the risk of leukemia, lymphoma, and breast cancer. Am J Hum Genet 1998; 62: 334-345.
5. Shaikh AG, Zee DS, Mandir AS, Lederman HM, Crawford TO. Disorders of upper limb movements in ataxia-telangiectasia. PLoS One 2013; 8: e67042.
6. Rothblum-Oviatt C, Wright J, Lefton-Greif M, McGrath-Morrow S, Crawford TO, Lederman HM. Ataxia telangiectasia: a review. Orphanet J Rare Dis 2016; 11:159.
7. Bhart JM, Bush A, van Gerven M, Nissenkorn A, Renke M, Yarlett L, et al. ERS statement on the multidisciplinary respiratory management of ataxia telangiectasia. Eur Respir Rev 2015; 24: 565-581.
8. Suarez F, Mahlaoui N, Canioni D, Andriamanga C, Dubois C, Brousse N, et al. Incidence, presentation, and prognosis of malignancies in ataxia-telangiectasia: A report from the French National Registry of primary immune deficiencies. J Clin Oncol 2015; 33: 202-208.
9. Voss S, Pietzner J, Hoche F, Taylor AR, Last JI, Schubert R, et al. Growth retardation and growth hormone deficiency in patients with ataxia telangiectasia. J Growth Factors 2014; 32: 123-129.
10. Hoche F, Seidel K, Theis M, Vlaho S, Schubert R, Zielen S, et al. Neurodegeneration in ataxia telangiectasia: what is new? What is evident? Neuropediatrics 2012; 43: 119-129.