Autism spectrum disorder and beta thalassemia minor: A genetic link?

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Received: 18th July 2017, Revised: 18th August 2017, Accepted: 17th September 2017


Autism spectrum disorder (ASD) is characterized by persistent deficits in social interaction and communication, and by restricted and repetitive patterns of behaviors and interests. Beta-thalassemia minor (BTM) is a common genetic blood disorder in Turkey. BTM is a single-gene disease that causes a decrease in beta globin production. We describe a girl aged 4 years and 4 months referred to our department due to speech delay, inability to establish social communication and overactivity. She was diagnosed with ASD according to DSM-5 criteria and Beta-thalassemia minor. Although there have been case reports of BTM with the some psychiatric conditions, to the best of our knowledge there are none concerning comorbid ASD and BTM. The aim of this report is to describe a possible genetic association between ASD and BTM since they have a common link associated with chromosome 11.

Key words: autism spectrum disorder, beta thalassemia minor, genetic, child.

Autism spectrum disorder (ASD) is a neuropsychiatric disorder characterized by persistent deficits in social interaction and communication, and by restricted and repetitive patterns of behaviors and interests.¹

Beta-thalassemia minor (BTM) is a common genetic blood disorder in Mediterranean countries, such as Turkey. BTM is a single-gene disease that causes a decrease in beta globin production and is frequently asymptomatic. Since it can cause mild microcytic hypochromic anemia it is frequently identified after blood count is performed for other reasons.²,³

At laboratory testing, hemoglobin values are slightly low, with a decrease in mean erythrocyte volume (MCV <80 fL) and mean erythrocyte hemoglobin (MCH) levels. In contrast to iron deficiency, erythrocyte numbers are normal or increased. At hemoglobin electrophoresis, hemoglobin A2 (>3.5%) and/or hemoglobin F (2–10%) are mildly elevated. Beta-thalassemia minor is a common and mostly symptomless disease and does not require treatment.⁴

Chromosome 11 has been implicated as being associated with psychiatric disorders such as attention deficit hyperactivity disorder (ADHD), Tourette syndrome, schizophrenia, and bipolar disorder.⁵-⁷ These psychiatric disorders are more common in patients with autism compared to the normal population. Genetics also play an important role in autism. The hemoglobin beta gene is located at 11p15.5. There have been reports suggesting that ASD is associated with the 11p15.5 location.⁸

Although there have been case reports of BTM with the above psychiatric conditions, to the best of our knowledge there are none concerning comorbid ASD and BTM. The aim of this report is to describe a possible genetic association between ASD and BTM since they have a common link associated with chromosome 11.

Case Report

A girl aged 4 years and 4 months was brought to our clinic by her family due to speech delay, inability to establish social communication and overactivity. At psychiatric evaluation, the patient made limited eye contact, did not look when called by name, did not obey verbal commands, did not engage in non-verbal behavior such as gestures or facial expressions,
exhibited inadequate emotional and social interaction and did not share her interests and successes. She was not toilet trained and was able to form only simple two-word sentences. She did not play games with age-appropriate themes, exhibited stereotypic behavior such as clapping and was intensely interested in spinning objects (such as a washing machine agitator). Her developmental level was assessed using the Ankara Developmental Screening Inventory (ADSI). Her age equivalence was 19 months at the global development level, 17 months at the linguistic-cognitive level, 22 months in terms of fine motor skills, 23 months in terms of gross motor skills and 20 months in terms of socialization/self-care. The Turkish versions of the Autism Behavior Checklist (ABC) and the Childhood Autism Rating Scale (CARS) were used to evaluate the severity of her autistic symptoms. Her total scores were 88 on the ABC and 46 on the CARS (indicating severe autism). Blood biochemistry tests, urinalysis, audiometry, thyroid function tests, brain magnetic resonance imagery, electroencephalography and abdominal ultrasound were all normal. Blood tests values were HGB: 9.564 g/dl (normal range 12-18.1 g/dl), MCV: 57.08 fL (normal range 80-97 fL) MCH: 14.84 pg (normal range 25-33.5 pg), and RBC: 6.444 10^6/mm³ (normal range 4.06-4.69 10^6/mm³). Serum iron, unsaturated iron binding capacity (UIBC), ferritin, B12 and B9 levels were normal. BTM was diagnosed on the basis of an HBA2 value of 4.8 (normal range <3%) and an HBF value of 0.6 (normal range <1%) at hemoglobin electrophoresis.

The patient was diagnosed with ASD and global developmental delay according to DSM-5, and with BTM. She was referred for special education.

An informed consent was received from the patient’s father

Discussion

We present a case of a child diagnosed with ASD, a neurodevelopmental disorder, and also with BTM. It is possible that having BTM was an incidental finding or that this disorder had played a contributory role in the development of autism. Multiple genetic factors and genetic-environmental interactions are implicated in the etiology of autism.

Chromosome regions and genes associated with autism have been identified through genome screening, association analysis and candidate gene analyses. Associations have been identified in different chromosomal regions, with eight being particularly important (2, 3, 7, 11, 15, 17, 22 and X chromosomes). Chromosome 11 is thought to be associated with several psychiatric disorders, including impulse control disorders, bipolar disorder, ADHD and Tourette’s syndrome. ADHD and tic disorder are more common in ASD compared to the normal population. There are reports a positive association between autism and the locus containing the gene for HRAS-1 (c-Harvey-ras-1). The HRAS-1 and beta globin genes are located at 11p15.5. No studies to date have investigated the prevalence of beta thalassemia in ASD patients. The presence of chromosome 11, one of the genes linked to autism, in the region containing the ß-globin gene, responsible for BTM, suggests that there may well be a genetic association between the two disorders. The limitation of this manuscript is that no chromosomal analysis was performed. Our scan of the literature revealed no reports of a probable genetic association between thalassemia minor and autism. To the best of our knowledge, this case study is the first to discuss such a possible link. Further studies are now needed to support this study.

REFERENCES