Clinical Practice and Epidemiology in Mental Health

Case report

**Obsessive compulsive disorder comorbidity in DBA**
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Abstract

Diamond-Blackfan Anemia (DBA) is a congenital erythroid aplasia characterized as a normochromic macrocytic anemia with a selective deficiency in red blood cell precursors in otherwise normocellular bone marrow. DBA is known to be associated with mental retardation and learning disabilities. Although comorbidities with other psychiatric conditions have not been reported in the existing literature, we report in this paper a case of a DBA patient with previously undiagnosed comorbidity of obsessive compulsive disorder (OCD), successfully treated with sertraline 200 mg/day and valproic acid 600 mg/day. This case of comorbid presentation has clinical, therapeutic and pathophysiological implications. Given the difficulty of distinguishing among mental retardation, learning disabilities and OCD and the importance of precocious diagnosis in treating OCD especially since there are treatment methods interfering with anemia symptoms, physicians should adapt an adequate screening tool treating a child with DBA and comorbid mental disorder.

Introduction

Diamond-Blackfan Anemia (DBA) is a congenital disease, characterized by a defective erythroid progenitor maturation and is associated with physical malformations.

Majority of cases are sporadic and dominant with 10% of the patients demonstrating recessive inheritance.

Mutations in the gene encoding for ribosomal protein RPS19 (DBA1) have been found in 25% of patients with either the dominant or the sporadic traits [1,2].

It is noteworthy that these mutations are associated with mental retardation as well as with learning disabilities in DBA patients [1,3].

Somatic abnormalities have been found in 47% of the patients registered with the DBA Registry of North America [4,5].

Associated physical anomalies and growth retardation are common and outstanding even in patients with multifactorial etiology such as long term steroid treatment [3].

The combination of clinical and molecular findings suggests a contiguous gene syndrome with a gene focus for mental retardation and skeletal malformations.

Repetitive and stereotyped behaviors are as common as mental retardation and in some cases their manifestations reach the threshold for diagnosis of obsessive compulsive disorder.
disorder (OCD) (according to the Diagnostic Statistical Manual of Mental Disease IV edition Text Revised DSM-IV TR criteria) [6].

In the following case we present a DBA patient with comorbidity of OCD. This case has tremendous significance due to the demonstration of the clinical and the pathophysiologic as well as therapeutic implications, involved in the assessment of behavioral abnormalities in DBA.

**Case presentation**

L. is a 22 year old Italian male, diagnosed with DBA at the age of two. Since being diagnosed with DBA, L. has been treated with monthly blood transfusions and subcutaneous injections of deferoxamine mesylate. Years later, he developed iatrogenic hepatitis due to multiple blood transfusions. Despite attending a special education program for children with learning disabilities, the patient has experienced major difficulties in carrying out daily activities since the age of six. He showed attention deficit at school, social isolation and, since the age of 12, verbal and motor repetitive behaviors, apparently cyclically worsening during mood instability episodes.

L. was reluctant to speak about his repetitive behaviors. L.'s parents attributed their child's behaviours to the developmental disabilities. A standard psychiatric diagnosis was not reached, no treatment was established during childhood.

L.'s grandfather was diagnosed with OCD (checking compulsions) in comorbidity with an Impulsive Control Disorder (Interruptional Explosive Disorders), his grandmother was depressed and alcohol addict.

At the age of sixteen the patient had an episode of herpetic encephalitis with symptoms of delirium and therefore, he was treated for two years with carbamazepine. One year later he was diagnosed with polyendocrinopathy of hypothyroidism, hypoparathyroidism and hypogonadism. The encephalitis process had no consequences. Neuroradiologic studies were performed and revealed white matter hypodensities in the right temporal lobe and in the ventricular choroid plexus, asymmetric sphenoid sinus and hypoplastic pituitary gland. Sellar region and parasellar structures appeared in a regular pattern. No anomalies in encephalic parenchyma were demonstrated after contrast medium injection.

The patient was assessed 12 weeks after the administration of the YBOCS and demonstrated an improved total score of 15, which corresponds to a mild form of OCD.

The patient is in the lower normal range of height (164 cm) and IQ (87). He complained of impulsive sexual and aggressive thoughts that were intrusive, repetitive and distressing. He also complained of compulsive behaviors and rituals, such as hoarding, arranging, ordering, preoccupations with symmetry, exactness, rewriting and doubting. Interrupting the patient while carrying out his rituals lead to violence. The patient had moderate insight of his illness.

At the age of twenty one repetitive behaviors increased in frequency to a level that required psychiatric attention and pharmacological management. Trying to address both mood symptoms and repetitive behaviors, a treatment with low doses of olanzapine and venlafaxine was established, with no improvement in symptoms and a strong deterioration of patient's anemia. L. was hospitalized and a more thorough psychiatric assessment was conducted. He fulfilled the DSM-IV TR criteria for OCD and diagnosis was established by the Structural Clinical Interview for DSM-IV Axis I Disorders (SCID-I) [7]. In order to determine the severity level of obsessive compulsive symptoms, the Yale Brown Obsessive Compulsive scale (YBOCS) [8], a clinician rated 10 items scale, each rated from 0 (no symptoms) to 4 (extremely severe symptoms), was performed on him and revealed a score of 28, which corresponds to a severe form. A treatment with sertraline 200 mg/day (addressing OCD symptoms) and valproic acid 600 mg/day (with the aim of reducing the impulsive features linked to obsessions and according to its efficacy reported in treatment of DBA) [9] has been started.

In the meantime an MRI exam was done as well and it showed low signal areas due to accumulation of paramagnetic substances in the right temporal lobe and in the ventricular choroid plexus, asymmetric sphenoid sinus and hypoplastic pituitary gland. Sellar region and parasellar structures appeared in a regular pattern. No anomalies in encephalic parenchyma were demonstrated after contrast medium injection.

The patient was assessed 12 weeks after the administration of the YBOCS and demonstrated an improved total score of 15, which corresponds to a mild form of OCD with a reduction of more than 45% of the symptoms.

**Discussion**

For the first time we have described DBA with comorbid OCD. The above described case could demonstrate heuristically valuable clinical, therapeutic and pathophysiological implications if more DBA patients with comorbid OCD would be screened by hematologists and therefore deserves further discussion.

There is a great importance in the assessment of obsessive-compulsive symptoms in DBA patients with mental or behavioral disturbances. Since OCD often goes undiagnosed in the presence of more pervasive disturbances [10], our report assumes a “Caveat” value.

Distinguishing between mental retardation, learning disabilities, Asperger Syndrome and OCD can be challenging, especially when treating children. Precocious diagnosis of
OCD can make a tremendous difference in terms of evolutionary trajectory and improved life quality of patients and their families.

Pediatricians should bear in mind the possibility of OCD when treating DBA children with behavioral learning disabilities even in the absence of other malformations.

It has been shown that when adequate screening tools were adopted in clinical disciplines other than psychiatry (for instance in dermatology and immunology), a larger than expected number of undiagnosed OCD patients was revealed [11].

Also, OCD is potentially linked to brain iron accumulation in DBA. Studies done in animals demonstrated that brain iron accumulation leads to damage of neuronal dopaminergic function.

Intraneuronal injection in rats have shown a detrimental effect on dopamine (DA) release and concentration in the caudate putamen (CPu) as well as selective decrease of striatal dopamine (95%), 3,4-dihydroxyphenylacetic acid serotogenic activity (82%), and homovanillic acid (45%) with related behavioral changes, characterized by increased repetitive and compulsive behaviors.

Thus, hemosiderosis might be contributing to psychiatric symptoms in DBA patients [12]. In fact, OCD symptoms may be linked to hemosiderin deposition in the brain and the pituitary gland, just as hypopituitarism has been shown to be linked to hemosiderin deposition in the pituitary as it was hypothesized previously by Berdel [13,14].

Moreover, since OCD has been related also to multiple regions of cortical thinning [15], the MRI imaging in our case of paramagnetic substance accumulation in the right temporal lobe, ventricular plexus and the hypoplastic pituitary gland is suggestive of the importance of neuroimaging assessment and recognition of complications caused by iron deposition due to long term blood transfusions in the management of DBA.

However a conclusion can not be drawn without a verification through larger studies on populations that have undergone blood transfusions at a young age.

Presenting this case report we have shown that OCD symptoms are treatable in DBA as effectively as in other conditions such as mental retardation [16] and Down Syndrome [17].

Despite the fact that some psychiatric medications have shown a worsening of the symptoms of anemia, SSRIs and valproate have been extremely beneficial and safe.

Authors' contributions
SP established the treatment of the patient, conceived the case-report and drafted the manuscript, SM drafted the manuscript, SB drafted the manuscript, MM drafted the manuscript, AI collected information about the case and drafted the manuscript, EH drafted the manuscript. All authors read and approved the final manuscript.

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References
1. Camagnoli MF, Garelli E, Quarello P, Carando A, Varotto S, Nobili B, Longoni D, Pecile V, Zecca M, Dufour C, Ramegni U, Dianzani I: Molecular basis of Diamond-Blackfan anemia: new findings from the Italian registry and a review of the literature. Haematologica 2004, 89(4):480-9.
2. Gazda H, Lipton JM, Willing TN, Ball S, Niemeyer CM, Tchernia G, Mohandas N, Daly MJ, Ploszynska A, Orfali KA, Vlachos A, Glader BE, Rościszewski W, Ohara A, Baker D, Pospisilova D, Webber A, Viskochil DH, Nathan DG, Beggs AH, Sieff CA: Evidence for linkage of familial Diamond-Blackfan anemia to chromosome 8p23.3-p22 and for non-19q non-8p disease. Blood 2001, 97:2145-50.
3. Tendler D, Gustavsson P, Elinder G, Eklöf O, Gordon L, Mandel A, Dahl N: A microdeletion in 19q13.2 associated with mental retardation, skeletal malformations, and Diamond-Blackfan anemia suggests a novel contiguous gene syndrome. J Med Genet 2000, 37(2):128-31.
4. Vlachos A, Klein GW, Lipton JM: The Diamond Blackfan Anemia Registry: tool for investigating the epidemiology and biology of Diamond-Blackfan anemia. J Pediatr Hematol Oncol 2001, 23(6):377-82.
5. Lipton JM, Arsic A, Zyskind E, Zyskind I, Vlachos A: Improving clinical care and elucidating the pathophysiology of Diamond Blackfan anemia: an update from the Diamond Blackfan Anemia Registry. Pediatr Blood Cancer 46(5):558-64. 2006 May I
6. Diagnostic and Statistical Manual of Mental Disorders. 4th edition. Washington, DC: American Psychiatric Association; 1994.
7. First MB, Spitzer RL, Gibbon M, Williams JBW: Structured Clinical Interview for DSM-IV Axis I Disorders, Clinician Version (SCID-CV). Washington DC: American Psychiatric Press, Inc. 1996.
8. Goodman W, Price H, Rasmussen S, Blake JB: The Yale Brown Obsessive Compulsive Scale (Y-BOCS): Part I. Development, use, and reliability, Archives of General Psychiatry. 1989, 46:1006-1011.
9. Jabr FI, Aoun E, Azar C, Taher A: Diamond-Blackfan anemia responding to valproic acid. Blood 104(10):3415. 2004 Nov 15
10. Carcani-Rathwell I, Rabe-Hasketh S, Santosh PJ: Repetitive and stereotyped behaviours in pervasive developmental disorders. J Child Psychol Psychiatry 2006, 47(6):573-81.
11. Leyfer OT, Folstein SE, Bacalman S, Davis NO, Dinh E, Morgan J, Tager-Flusberg H, Lainhart JE: Comorbid psychiatric disorders in children with autism: interview development and rates of disorders. J Autism Dev Disord 2006, 36(7):849-61.
12. Wexemann W, Blaschke S, Solbach M, Grote C, Clement HW, Riederer P: Intraneuronal injected iron progressively reduces striatal dopamine metabolism. J Neural Transm Park Dis Dement Sect 1994, 8(3):209-14.
13. Berdel D, Romahn A, Burmeister W: Pluriglandular insufficiency due to transfusion haemosiderosis in Blackfan-Diamond anemia. Klin Padiatr 1980, 192(1):91-4.
14. Sparacino G, Banco A, Midiri M, Iaia A: MR imaging technique for the diagnosis of pituitary iron overload in patients with transfusion-dependent beta-thalassemia major. AJNR Am J Neuroradiol 1998, 19(10):1905-7.
15. Shin YW, Yoo SY, Lee JK, Ha TH, Lee KJ, Lee JM, Kim IY, Kim SL, Kwon JS: Cortical thinning in obsessive compulsive disorder. Hum Brain Mapp 2007, 28(1):128-35.
16. Bodfish JW, Madison JT: Diagnosis and fluoxetine treatment of compulsive behavior disorder of adults with mental retardation. Am J Ment Retard 1993, 98(3):360-7.

17. Sutor B, Hansen MR, Black JL: Obsessive compulsive disorder treatment in patients with Down syndrome: a case series. Downs Syndr Res Pract 2006, 10(1):1-3.