Depressive disorders co-existing with Addison–Biermer anemia – case report

Mark Jean Just1
Mariusz Kozakiewicz2

1Department of General and Endocrinological Surgery, Piekary Medical Centre, Piekary Śląskie, 2Department of Food Chemistry Collegium Medicum in Bydgoszcz, Nicolaus Copernicus University, Toruń, Poland

Background: Anemia is a disease that can co-exist with depression, other mental disorders, or somatic diseases. Anemia can imitate symptoms of depression, while depression symptoms can mask concurring symptoms of anemia.

Case presentation: I am presenting a case of a 48-year-old woman with Addison–Biermer anemia, with co-existing mood disorders. The clinical analysis of the presented patient’s history indicates diagnostic problems and a need for a detailed analysis of drug-related complications that occurred during previous treatment, eg, in the form of neuroleptic malignant syndrome.

Conclusion: The presented case report contains valuable guidelines that can be of assistance in diagnostics and treatment of patients treated for mental disorders, who are also diagnosed with somatic diseases.

Keywords: anemia, autoimmune diseases, depression, neuroleptic malignant syndrome

Introduction

Pernicious anemia, also called Addison–Biermer anemia, is an autoimmune disease most often occurring in people with A blood type, and caused by vitamin B12 absorption disturbances (Figure 1). In the initial clinical presentation, symptoms of this anemia, including apathy, psychomotor retardation, lack of energy, fatigability, or excessive sleepiness, can indicate a possibility of concurring depression.1,2 Furthermore, symptoms of depression can overlap with somatic symptoms of anemia, further complicating the clinical presentation, and therefore, the treatment of underlying disease, or even persisting when anemia symptoms have resolved. Anemia was identified in 10% of patients with depression. Patients particularly susceptible to stress and suffering from anxious personality disorders concomitant with somatic diseases belong to a risk group prone to depressive disorders and anemia. Depression can develop against a background of a chronic stress reaction related to the somatic disease.3 Chronic stress can also lead to development of conversion disorders, with partial or complete loss of correct integration of past memories with a sense of own identity.4,5 Increased conversion or depression symptoms in patients with Addison–Biermer anemia can represent a difficult diagnostic and treatment challenge to hematology specialists, general practitioners, and psychiatrists. Treatment of depression and conversion symptoms should combine pharma- and psychotherapy.6 For severe depression, treatment with anti-depressive drugs alone may be insufficient, requiring additional inclusion of atypical neuroleptics.7 It is worth noting that in people suffering from depression and with a positive history of somatic diseases, such as Addison–Biermer anemia or endocrine diseases, the neuroleptic malignant syndrome (NMS) is more likely to occur following neuroleptic treatment, than in people without somatic diseases.
Case presentation

I am presenting a clinical case of a 48-year-old patient, with A blood type, admitted to the surgical ward due to anemia. It is known from the interview that the patient was previously pharmacologically treated for hypothyroidism, but has not taken medications for several years. The patient felt weak, had concentration problems, and reduced tolerance of physical exercise for approximately 1 month. In physical examination, pale skin and mucous membranes were found.

The patient underwent the following examinations: ultrasound scan of the abdominal cavity, gastroscopy, colonoscopy, computed tomography of the abdominal cavity, and ultrasound scan of the thyroid. The results showed no changes. Laboratory tests conducted on admission day revealed: white blood corpuscles 4.56 K/µL; lymphocytes 2.41; red blood corpuscles 3.00 M/µL; hemoglobin 8.74 g/dL; mean corpuscular volume 81.8 fl; mean corpuscular hemoglobin 28.7 pg; mean corpuscular hemoglobin concentration 35.7 g/dL; platelets 158 K/µL; vitamin B₁₂ level 120 pg/mL (reference value 180–900 pg/mL); iron level 160 µg/dL (reference value 60–160 µg/dL). Thyroid hormone levels for free thyroid hormone – triiodothyronine, free thyroid hormone – thyr oxine, and thyroid-stimulating hormone levels were within the reference values.

Only in the tenth day of her hospitalization, the patient disclosed that in the past she had undergone psychiatric treatment for depression and conversion disorders. She was afraid of mental disease-related stigma. Five years before current hospitalization, the patient had had NMS due to treatment with olanzapine for psychotic depression. During psychiatric consultation, the patient admitted she had not taken neuroleptics for about a year. She decided to discontinue the therapy. For the last 3 months, the patient has complained about deteriorating psychological condition revealing symptoms such as: low mood, apathy, anhedonia, anxiety, memory and concentration problems, sleep disturbances, and suicidal thoughts. The patient was diagnosed with severe depression without psychotic symptoms and with Addison–Biermer anemia. Depression diagnosis was based on clinical assessment of the patient and followed by performing Beck Depression Inventory in which the patient received 26 scores. Pharmacological treatment was commenced, with venlafaxine,
Depressive disorders co-existing with anemia

Vessel interventions. Anemia is an important mortality and long-term risk of death in coronary disease patients. Heart infarction with ST segment elevation, and with acute coronary syndromes without a stable form of that disease, heart infarction with ST segment elevation, and with acute coronary syndromes without a stable form of that disease, heart infarction with ST segment elevation, and with acute coronary syndromes without a stable form of that disease, heart infarction with ST segment elevation, and with acute coronary syndromes without a stable form of that disease, heart infarction with ST segment elevation, and with acute coronary syndromes without a stable form of that disease, heart infarction with ST segment elevation, and with acute coronary syndromes without a stable form of that disease, heart infarction with ST segment elevation, and with acute coronary syndromes without a stable form of that disease.

Anemia is associated with higher short- and long-term risk of death in coronary disease patients with a stable form of that disease, heart infarction with ST segment elevation, and with acute coronary syndromes without ST segment elevations, as well as in patients after coronary vessel interventions. Anemia is an important mortality and morbidity risk factor in chronic kidney failure. A decrease in hemoglobin (Hb) by 1 g/dL is associated with 14%–18% increase in mortality in a dialysis patient population, while Hb level decrease below 8 g/dL doubles the risk of death versus patients with Hb of 10–11 g/dL.

In depression, concurrence of somatic diseases, such as Addison–Biermer anemia or thyroid hormone disturbances can delay therapy, for example, using electroconvulsive therapy in treatment of severe depression. In depression, concurrence of somatic diseases, such as Addison–Biermer anemia or thyroid hormone disturbances can delay therapy, for example, using electroconvulsive therapy in treatment of severe depression. In depression, concurrence of somatic diseases, such as Addison–Biermer anemia or thyroid hormone disturbances can delay therapy, for example, using electroconvulsive therapy in treatment of severe depression. In depression, concurrence of somatic diseases, such as Addison–Biermer anemia or thyroid hormone disturbances can delay therapy, for example, using electroconvulsive therapy in treatment of severe depression. In depression, concurrence of somatic diseases, such as Addison–Biermer anemia or thyroid hormone disturbances can delay therapy, for example, using electroconvulsive therapy in treatment of severe depression.

In treatment of depression, delayed use of electroshocks can result in exacerbation of depression and contribute to the increased risk of suicide. Moreover, in patients treated with atypical neuroleptics there is a risk of NMS. According to the Diagnostic and Statistical Manual of Mental Disorders, fourth edition (DSM-IV), it is possible to diagnose NMS when muscle rigidity and fever are found with at least two of the following symptoms concurring: diaphoresis, dysphagia, incontinence, changes in level of consciousness, mutism, tachycardia, increased or labile blood pressure, leukocytosis, and increased CK levels, indicating muscle injury. Other criteria proposed by Levenson include “major symptoms” (fever, muscle rigidity, and increased CK levels) and “minor symptoms” (tachycardia, variable blood pressure, tachypnea, changes in level of consciousness, diaphoresis, and leukocytosis). The NMS diagnosis is justified in cases with all “major symptoms” present, or when two of them are accompanied by at least two “minor symptoms”. NMS can also develop in patients with positive history of somatic diseases, such as thyroid disorders, anemia, and neurological diseases, or in patients without chronic diseases.

Furthermore, patients who suffered from NMS caused by olanzapine are predisposed to NMS in the future also after administration of other drugs affecting the dopaminergic system. Metabolic changes secondary to hypothyroidism or Addison–Biermer disease, particularly in the dopaminergic pathways within the central nervous system, can also predispose to the development of NMS.

Discussion

When suffering from somatic symptoms of anemia or symptoms resulting from a thyroid function disorder, patients usually visit their general practitioner, hematologist, endocrinologist, or surgeon. The most important task of the doctor is to establish whether some or all of presented symptoms are of an organic origin, or not. The organic cause requires a clinical intervention, eg, in the form of pharmacological or surgical treatment. However, it should be remembered that the initial organic disease may be accompanied by concurrent secondary mental disorders. It is recommended to perform screening laboratory tests in patients with depression, excluding anemia and hypothyroidism, such as: complete blood count, serum iron level, total iron binding capacity, unsaturated iron binding capacity, serum vitamin B12 level, and thyroid-stimulating hormone. In patients belonging to a risk group susceptible to depression, it is recommended to determine oxidative stress markers, eg, chemokines, cytokines. These markers may constitute an early predictor of affective disorders. When the organic cause is excluded, or when concurrent symptoms of mental disorders are observed, then the basic medical procedure involves psychiatric consultation for correct diagnosis and treatment initiation.

Addison–Biermer anemia can co-exist with other somatic diseases or accompany mental disorders (depression, anxiety disorders, conversion disorders).

It is a strong predictor of adverse clinical incidents related to cardiac ischemia. Anemia is associated with higher short- and long-term risk of death in coronary disease patients with a stable form of that disease, heart infarction with ST segment elevation, and with acute coronary syndromes without ST segment elevations, as well as in patients after coronary vessel interventions. Anemia is an important mortality and morbidity risk factor in chronic kidney failure. A decrease in hemoglobin (Hb) by 1 g/dL is associated with 14%–18% increase in mortality in a dialysis patient population, while Hb level decrease below 8 g/dL doubles the risk of death versus patients with Hb of 10–11 g/dL.

In treatment of depression, delayed use of electroshocks can result in exacerbation of depression and contribute to the increased risk of suicide. Moreover, in patients treated with atypical neuroleptics there is a risk of NMS. According to the Diagnostic and Statistical Manual of Mental Disorders, fourth edition (DSM-IV), it is possible to diagnose NMS when muscle rigidity and fever are found with at least two of the following symptoms concurring: diaphoresis, dysphagia, incontinence, changes in level of consciousness, mutism, tachycardia, increased or labile blood pressure, leukocytosis, and increased CK levels, indicating muscle injury. Other criteria proposed by Levenson include “major symptoms” (fever, muscle rigidity, and increased CK levels) and “minor symptoms” (tachycardia, variable blood pressure, tachypnea, changes in level of consciousness, diaphoresis, and leukocytosis). The NMS diagnosis is justified in cases with all “major symptoms” present, or when two of them are accompanied by at least two “minor symptoms”.

NMS can also develop in patients with positive history of somatic diseases, such as thyroid disorders, anemia, and neurological diseases, or in patients without chronic diseases.

Furthermore, patients who suffered from NMS caused by olanzapine are predisposed to NMS in the future also after administration of other drugs affecting the dopaminergic system. Metabolic changes secondary to hypothyroidism or Addison–Biermer disease, particularly in the dopaminergic pathways within the central nervous system, can also predispose to the development of NMS.

Conclusion

In treatment of depressive disorders, a possibility of concurrence of somatic disorders in the same patient should also be considered. Taking into consideration the fact that mental disorders often represent prodromal signs of various somatic diseases, it is necessary to remember that additional tests and evaluation of the somatic status are of particular importance in psychiatric patients. It is worth noting that these patients focus on their emotional experience and ignore their somatic problems.
Acknowledgments
The work was carried out at Department of General Surgery, Municipal Hospital in Piekary Slaskie, Poland. Professional language editing has been performed by native speaker.

Author contributions
Both authors contributed toward data analysis, drafting and revising the paper and agree to be accountable for all aspects of the work.

Disclosure
The authors report no conflicts of interest in this work.

References