TREATMENT-RESISTANT IMMUNE THROMBOCYTOPENIC PURPURA ASSOCIATED WITH LDN USE IN A PATIENT WITH MS

Naltrexone is an opiate µ-receptor antagonist approved for treatment of alcohol dependency at a recommended dose of 50 mg/day. The mechanism of action is proposed to be mediated by a normalization of endogenous endorphin levels. Despite few and inconclusive studies (casuistic or open-label), a naltrexone dose of 3–5 mg/day (low-dose naltrexone, LDN) has been suggested to ameliorate a wide range of diseases, especially Crohn disease1 and multiple sclerosis (MS).2 Its effect in these conditions is mainly advocated by patients via both formal and informal networks. Reluctance among physicians to prescribe naltrexone has encouraged illegal import. Based on safety data from initial marketing studies conducted in the 1970s, naltrexone has generally been considered safe, with no serious side effects or adverse events. We present a case of therapy-resistant immune thrombocytopenic purpura (ITP) after LDN use in a patient with MS.

Case presentation. A 64-year-old Caucasian woman was diagnosed with relapsing-remitting MS in 2012 after an episode of optic neuritis. Her diagnosis was confirmed by MRI fulfilling the MacDonald criteria and by the detection of oligoclonal bands in CSF. Routine blood work at the time of diagnosis and 3 years prior to diagnosis showed nothing abnormal but a mild thrombocytopenia (110 × 10^9/L; figure). Follow-up MRI showed new and enlarging lesions, and she was prescribed immunomodulatory treatment with interferon β. However, due to needle phobia, not previously recognized by the patient or the treating physician, she never started interferon β therapy. Instead, encouraged by positive reports from peers on the effect of LDN in MS, she acquired naltrexone from an Internet provider located in Israel and started treating herself with 3 mg/day. She reported a transient positive effect on her MS-related fatigue.

After 3 to 4 months of LDN treatment, she noticed spontaneous formation of bruises and gum bleeding. After 4 months, she experienced profuse gum bleeding during a dental examination. Routine blood count at her general physician’s lab showed a thrombocyte count (TPC) below 5 × 10^9/L (reference range 165–387 × 10^9/L) without effects on the other blood cells, and she was referred to the Department of Hematology at Haukeland University Hospital. Liver function tests were normal. Other rheumatologic, autoimmune, or infectious causes of low TPC were not detected, and bone marrow smears were consistent with ITP. Due to the temporal relationship between LDN treatment and emergence of ITP symptoms, LDN was considered the most probable cause of her ITP and the medication was discontinued after the diagnosis was made. After a short initial response to prednisolone therapy, TPC again fell below 5 × 10^9/L. Neither IV immunoglobulins nor rituximab in combination with steroids had more than a transient effect on TPC. She underwent splenectomy in May 2014, after which platelet counts stabilized at around 136 × 10^9/L (figure).

Discussion. ITP is defined as thrombocytopenia (low platelet count) with normal bone marrow findings and the absence of other causes of thrombocytopenia. It is generally considered an autoimmune disease, but the exact cause is unknown. Naltrexone, used against opioid dependency, was found to increase the risk of developing ITP in the initial marketing studies.3 LDN is usually taken at a dosage of about 10% of what was given in the initial studies. Some studies have been conducted on the effect of LDN in MS, with varying results, but without reporting any serious adverse events.4,5 However, since the use of LDN is largely off-label, no large safety studies have been conducted. Advocates of LDN use have typically pointed out that this medication is completely safe and without any serious side effects. As this case presentation demonstrates, LDN could possibly predispose to an increased risk of developing ITP. Our patient had platelets at about 120 × 10^9/L for at least 3 years prior to diagnosis with ITP, and it is possible that the association with LDN use is spurious. However, since naltrexone use is known to increase the risk of ITP, it is biologically plausible that LDN has either triggered her ITP or aggravated a subclinical disease. This case demonstrates a possible safety concern
regarding LDN treatment and the importance of thorough review of patients’ on- and off-label drug use.

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