Heart Involvement in a Woman Treated with Hydroxychloroquine for Systemic Lupus Erythematosus Revealing Fabry Disease

To the Editor:

Chloroquine (CQ) and hydroxychloroquine (HCQ) are widely used in the longterm treatment of connective tissue disease (CTD) and are usually considered safe. However, these therapies may cause serious adverse events, including cardiac toxicity. The cardiomyopathy induced by an antimalarial drug might mimic the cardiac involvement of Fabry disease (FD), a genetic storage disorder that causes a deficiency of a lysosomal enzyme, alpha galactosidase A (GLA). Indeed, CQ and HCQ provoke a dysfunction in the lysosomal enzymes, leading to the impairment of intracellular degradation processes in conjunction with the accumulation of pathological metabolic products. The cardiac presentation of these 2 diseases is close regarding the clinical symptoms as well as the imaging and pathological findings. We report the case of a patient, treated longterm with HCQ, with cardiac disorders revealing FD.

A 61-year-old woman was diagnosed 20 years ago for a CTD with dermatological disorders: alopecia, skin rash, and joint pain. The antinuclear antibodies (ANA) were positive at 1/100. Anti-SSa/Ro and anti-dsDNA antibodies were negative. The skin biopsy showed moderate lymphocytic infiltrate. Systemic lupus erythematosus (SLE) was diagnosed and HCQ (200 mg twice daily) was started in combination with low doses of prednisone (10 mg per day), which led to clinical improvement. Seventeen years later, she presented sudden neurologic disorders with dizziness, left arm weakness, and facial paralysis. The brain magnetic resonance imaging (MRI) showed a recent ischemic stroke of the right middle cerebral artery. In addition, we retrospectively found that 2 of the patient’s brothers were dead of an early sudden cardiac ischemia while under 50 years old. Our definitive diagnosis was cardiomyopathy attributable to FD, probably promoted by the longterm use of HCQ (total cumulative dose of 2480 g).

FD is associated with cardiovascular disorders (myocardial ischemia and brain strokes) and renal impairment. Only 70% of women with the mutation of the GLA gene have manifestations of FD. A large number of women are not diagnosed. The pathology of myocardial tissue is extremely close between FD and HCQ toxicity. In their study, Roos et al. compared both cardiac disorders. They observed the same clinical presentation and the same pathological features with myelin figures and glycogen accumulation in both biopsies. However, they concluded that curvilinear bodies were only present in CQ toxicity. Here, some curvilinear bodies were described, likely meaning that HCQ is partially involved in the cardiac failure. In the literature, to our knowledge, 23 patients with HCQ or CQ cardiotoxicity are reported. They were frequently women and the duration of treatment was between 9 and 35 years. Regarding heart presentation, almost all the patients had left ventricular hypertrophy. The other cardiac events were conduction disorders, rhythm disorders, congestive heart failure, ischemia with hypokinesia, and diastolic dysfunction. Several patients presented chest pain, increasing dyspnea, and elevated cardiac enzymes. Treatment withdrawal could have beneficial effects in some patients with recovery of heart function, but sometimes the development was unfavorable with irreversible damage, death, or heart transplant. FD was screened in only 7 patients (6 negative and 1 already known positive), whereas the pathologic features could correspond to FD. Only 1 case reported on SLE treated with HCQ.
associated with FD. The endomyocardial biopsy showed only myelin figures. Despite HCQ withdrawal and enzyme replacement therapy, this patient continued to have vascular complication with a myocardial infarction and stroke.

Our case highlights HCQ cardiotoxicity and the risk of misdiagnosed FD, especially in women because of nonspecific symptoms. We must also point out the need to realize cardiac biopsies in atypical cardiac involvement in SLE.

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