Lupus Outbreak and Covid-19 Pneumonia-Case Report

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Abstract
Coronavirus disease 2019 (COVID-19) is a respiratory infection that can cause mild symptoms or even death, to patients who suffer from it. It affects all population groups without distinction. Systemic Lupus Erythematosus (SLE) is a chronic and fluctuating autoimmune disease. One of the goals of the treatment is to avoid flare-ups and thereby reduce mortality. Their innate alterations in immunity, added to the use of immunosuppressive drugs to control the disease and prevent outbreaks, makes them more vulnerable to develop severe symptoms in the course of SARS-CoV-2 infection. We present the case of a patient with SLE infected by SARS-CoV-2 with a lupus flare during hospitalization, entailing a diagnostic and therapeutic challenge.

Key words: sle, covid-19, lupus-flare, sars-cov-2

Introduction
It has been more than a year (March 11, 2020) since the World Health Organization (WHO) declares the coronavirus disease 2019 (COVID-19) as a pandemic [1]. There are still many questions about severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2). In its mild form, infected patients develop well-tolerated respiratory, gastrointestinal, and neurological symptoms, but in its severe form they can developed septic shock and multiple organ failure [2-4]. The most affected populations are the elderly and especially those with comorbidities. The immune response plays an important role against viral diseases, both as antiviral defense and in disease progression. Systemic Lupus Erythematosus (SLE) is a chronic, autoimmune and multisystemic disease with a fluctuating clinical course. It mainly affects women, being more prevalent African and Hispanic [5]. Research has been conducted prior to the pandemic showing that patients with SLE are at increased risk of infection, a risk that is greater if patients have higher levels of disease activity, the use of corticosteroids on a regular basis, or who have previously received treatment with cyclophosphamide or rituximab [6,7]. However, some studies suggest that immunosuppression secondary to SLE management may be beneficial by reducing the immune system’s response to COVID-19 [8-9], while other studies report that intense immunosuppression does not allow an effective antiviral response to the exposure to COVID-19, triggering a more severe form of the disease [10,11]. The COVID-19 global rheumatology alliance reported one of the largest series of 600 Covid-19 patients, with various underlying rheumatological disorders from 40 countries. Although the study did not report on the incidence of Covid-19 in these patients, although describes that a glucocorticoid dose greater than 10 mg per day was associated with a higher chance of hospitalization [12].

Despite the various available treatment options at present, only 20-30% of patients with SLE have an inactive or active chronic disease [13,14], the rest experiment flare-ups periods that can cause permanent organic damage with increase of morbidity, mortality, and health costs [15-19].

The consensus definition of lupus flare refers to a measurable increase in the disease activity involving new or worse clinical signs and symptoms and/or laboratory markers. This should be considered clinically significant by the physician and generally leads to evaluate a treatment change or upscale [20]. Of the Instruments available and used to date to assess disease activity, the SLE Disease Activity Index (SLEDAI) with modifications (SLEDAI 2K, SELENA-SLEDAI) [21] and the British Isles Lupus Assessment Group 2004 (BILAG 2004) [22] are the most widely used in both clinical practice and research studies. SLEDAI reflects disease activity in the last 30 days and has a significant prognosis value of mortality in the next 6 months [23]. Among the serological
markers, the increase in the titers of the anti-dsDNA double-chain antibody, and the low levels of complement are part of the evaluation.

The European League against Rheumatism (EULAR) guide, within its recommendations establishes that outbreaks can be treated according to the severity of organ involvement by adjusting ongoing therapies (glucocorticoids, immunomodulatory agents) to higher doses, changing, or adding new therapies. The goal of preventing flare-ups is to avoid permanent organ damage and thereby improve the prognosis [24].

Strict control and monitoring are a fundamental part to reduce outbreaks in patients with SLE; however, the coronavirus pandemic has limited controls in this type of patients given that exposing them to public and hospital circulation leads to an increased risk of contagion. This situation generated a higher risk of poor adherence, greater presence of decompensations, outbreaks or worsening and the possibility of coexisting a flare and coronavirus infection.

The following report is a clinical case of a patient with lupus outbreak and SARS-COV-2 infection.

**Case Presentation**

A 23-year-old female patient with history of systemic lupus erythematosus (SLE) approximately diagnosed 3 years ago with lupus nephritis and HBP.

Usual medication: cyclosporine 100 mg/day, mycophenolate 250 mg every 8 hours, enalapril 10 mg day, ASA (acetylsalicylic acid) 100 mg/day, Omeprazole, Furosemide 40 mg / day.

She consulted in a peripheral hospital due to clinical presentation characterized by persistent non-productive cough associated with dyspnea on moderate efforts. She was admitted for COVID-19 suspicion and treatment according to the Covid-19 Protocol was initiated. Faced with the refusal of the diagnosis by relatives, voluntary discharge is requested and continued with outpatient medical treatment. At home, she continued with persistent cough, fever, adds anosmia, ageusia and progression of dyspnea. On 01/14/2021, after 1 month of clinical symptoms, she re-entered the peripheral hospital and later was transferred to El Salvador’s National Hospital.


Laboratory admission: Hematocrit: 39.2%, Hemoglobin: 13.4 gr/dl, Platelets: 358,000, White blood cells: 7,600, Neutrophils: 93.9%, Lymphocytes: 3.30%, Urea: 39.1 mg/dl, Creatinine: 0.68 mg/dl, Sodium: 139 meq/l, Potassium: 5.3 meq/l, AST UI/l: 115, ALT: 36.9 UI/l, LD: 662 UI/l, CRP = 8.1 mg/l, Ferritin: 2065 ng/ml, IL6: 33.8, Fibrinogen: 349 mg/dl, DD: 2824.98 mg/l. She was admitted to the intensive care unit and started with ventilatory support with a reservoir mask, achieving SATO2: 95%. The COVID-19 Protocol continues. A chest X-ray is performed (Fig.1) which shows bilateral infiltrates left predominantly, and bilateral blunting of costophrenic angles. Arterial blood gas: PH: 7.38, PCO2: 33.5, PO2: 65.9, HCO3: 20.1.

**Figure 1:** Chest X-ray

Twenty four hours after admission the patient presented drowsiness, delirium, radiographic infiltrates progression, hypoxemia, increased breathing effort and arterial hypertension. Oxygen support device was change to a high-flow cannula and set with the following parameters: 50 lts/min flow and fio2 50%, improving saturation to 96%. Thoracic (Fig. 2,3,4) and abdominal (Fig. 5,6) CT scan without contrast was performed, it showed: multiple ground glass opacities in both lung fields, which compromise more than 75% of each one, in addition there is subpleural septal thickening, bilateral basal subpleural bands, and some areas of subpleural honeycombing changes in the apex. In addition, pericardial effusion with a thickness of up to 3.2 cm extends to the pericardial recesses. Basal bilateral laminar pleural effusion was also observed. In the mediastinum, some inflammatory nodes are identified. As an incidental finding, the presence of free perihepatic, perisplenic fluid, was an incidental finding in upper abdomen. Both kidneys enlarged, swollen, with bilateral pararenal laminar fluid. Soft tissues and bone structures present no abnormalities.
**Figure 1:** Thoracic CT scan without contrast

**Figure 2:** Thoracic CT scan without contrast

**Figure 3:** Abdominal CT scan without contrast
Clinical finding seven days after admission suggested deep vein thrombosis in right inferior limb and venous doppler scan shows right femoral and saphenous vein thrombosis. Due to COVID-19 protocol the patient received anticoagulation treatment with enoxaparin 1 mg/kg/day, dose was increased to 1 mg/kg every 12 hours, and antiplatelet agents were add.

Given the CT scan findings associated with arterial hypertension and altered level of consciousness, in addition to COVID-19 pneumonia, clinical signs of lupus outbreak are suspected. SLEDAI scale is performed: 20 (Organic brain syndrome: 8, Urinary casts: 4, Proteinuria: 4, Pleurisy: 2, Pericarditis: 2), corresponding to high activity or very high activity. A general urine test and 24 hrs urine collection test informed: Creatinine clearance: 56 ml / min, proteins in 24 hrs: 6.31 gr / 24hrs.

Given these findings with high suspicion of a lupus outbreak and without evidence of active infection (negative cultures), it was decided to start pulse corticosteroid therapy at a dose of 1 gram / day for 3 days.

The patient evolves favorably with remission of arterial hypertension, drowsiness, headache, delirium, without progression of respiratory failure. Corticosteroids were tapered down until 10 mg per day is achieved, then immunosuppressive treatment is started.

During hospitalization the patient presented asymmetric edema of the left upper limb. Due to deep venous thrombosis clinical suspicion, a vascular doppler was performed, which informed left upper limb cephalic vein thrombosis. Anticoagulation with oral anticoagulants was prescribed.

**Discussion**

The present case report represents a great medical dilemma, due to diagnostic approach and therapeutic decisions complexity of patients with chronic multisystemic diseases admitted to critical care units. Patients with SLE in an advanced-stage renal failure represent a medical challenge itself, in this case report severe COVID-19 pneumonia obscure the medical diagnosis. Despite the substantial bilateral involvement of the lung parenchyma, the latent risk of mechanical ventilation gradually decreased over the hospitalization. However, we noticed clinical manifestations (altered level of consciousness, fever, joint pain, arterial hypertension), laboratory and imaging (proteinuria, increased inflammatory parameters, serositis) that could not be fully attributed to COVID-19, which generated a reasonable clinical uncertainty of a Lupus Outbreak.

To corroborate our clinical suspicion all the methods available in our institution were performed, to prescribe corticosteroid pulses therapy because of the risk that this therapy entails. The use of corticosteroids is justified in the lupus flare, but in COVID-19 there was no evidence available of the benefit of corticosteroids until the publication of the RECOVERY study and to date the METCOVID trial did not show benefit in mortality to 28 days in those under 60 years of age.

In the present clinical case, it was pondered the lupus outbreak as the principal cause of the patient's health deterioration, for which the decision was made to perform corticosteroid pulses therapy, achieving remission of symptoms, reduction of inflammatory parameters and clinical
improvement significant. Based on the clinical evolution, we can interpret the benefit of the therapy with high-dose corticosteroids pulsed in relation to the lupus flare, but we do not know if it could have contributed to limit the COVID 19 cytokine storm [27].

**Conclusion and Final Considerations**

Patients with autoimmune and immunosuppressed diseases always present a great challenge when they are admitted to the critical care unit. We have witnessed that COVID-19 generates a different impact in each population group and to date we still have many questions about how its approach should be in certain patients. The clinical suspicion, the sum of variables, the multidisciplinary intervention in decision-making together with the best available evidence is an adequate strategy in the fight against this terrible disease.

**Abbreviations**

WHO: World Health Organization
SARS-CoV-2: Severe acute respiratory syndrome coronavirus 2
SLE: Systemic Lupus Erythematosus
SELENA: Systemic Lupus Erythematosus Disease Activity Index
SELENA 2K: Systemic lupus erythematosus disease activity index 2000
BILAG: British Isles Lupus Assessment group
ANTI-dsDNA: Anti-double stranded DNA
EULAR: European Alliance of Associations for Rheumatology
AST: Aspartate transaminase.
ALT: Alanine transaminase
LD: L-lactate dehydrogenase
CRP: C-reactive protein
IL6: Interleukin 6
NA: sodium
K: Potassium

**References**


