Dermatological Side Effects in a Patient with Hepatitis C Infection During Treatment with Pegylated Interferon/Ribavirin+Telaprevir: A Case Report

Hepatit C Enfeksiyonu Olan Hastada Pegile Interferon/Ribavirin+Telaprevir Tedavisi Sırasında Görülen Dermatolojik Yan Etki: Olgu Sunumu

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ABSTRACT

Chronic hepatitis C (HCV) and pegylated interferon (peg-IFN)/ribavirin (RBV) treatment dermatological side effects are well known. New direct-acting antivirals have led to significant improvements in sustained virologic response rates, but several have led to an increase in dermatological side effects versus peg-IFN/RBV alone. A 55-year-old man was administered peg-IFN alpha-2a 180 mcg/one times a week, RBV 1200 mg/day and telaprevir 2250 mg/day with the diagnosis of chronic hepatitis C. Around the 8th week of the treatment the patient presented with itching eruptions on his face, neck, anterior wall of the and the forearm. A twelve administration of the combination therapy, the patient admitted to our hospital itching, rash, edema in whole body and facial edema and peeling in his facial skin. As a result, with the advent of the new direct-acting antivirals, dermatological manifestations will be seen more frequently so patients should be monitored closely in terms of dermatological side effects. (Viral Hepatitis Journal 2014; 20(2): 81-84)

Key words: Rash, hepatitis C, pegylated interferon, ribavirin, telaprevir

ÖZET


Anahtar Kelimeler: Döküntü, hepatit, pegile interferon, ribavirin, telaprevir

Introduction

The worldwide incidence of hepatitis C virus (HCV) infection is between 1% and 5%. In our country the incidence of HCV infection was reported to be 1%-2.4%. These ratios vary between 0.05%-51.6% in different populations (1).

The standard treatment for chronic hepatitis C infection is based on a combination of pegylated interferon (peg-IFN) and ribavirin (RBV). This treatment is associated with a variety of mild or severe adverse events, such as flu-like symptoms, rash, anemia, trombocytopenia, neutropenia and depression (2). The incidence of dermatological adverse events associated with peg-IFN alpha 2a or alfa 2b/RBV during treatment for hepatitis C infection is >10% (3). Sometimes it can be difficult to distinguish cutaneous signs and symptoms of hepatitis C from interferon treatment-related adverse events (4).

New direct-acting antivirals have led to significant improvements in sustained virologic response rates, but several have led to an increase in dermatological side effects versus peg-IFN/RBV alone. It has been reported that dermatological adverse events are seen in approximately half of patients receiving peg-IFN/RBV+telaprevir treatment (5,6). Here, we report a case of dermatological adverse event that occurred during peg-IFN/RBV+telaprevir treatment.

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Case

A 55-year-old male patient admitted to our outpatient clinic in 2009 after he was found to be anti HCV (+) during his routine blood tests. In 2009, his tests revealed: anti-HCV (+), HBsAg (-), anti-HBc IgG (+), anti-HBs (+), anti-HAV total (+), anti-HIV (-), aspartate transaminase (AST): 48 IU/mL, alanine transaminase (ALT): 73 IU/mL. His protein/albumin ratio was 7.1/3.6 g/dL, alpha fetoprotein levels were normal, HCV RNA: 1.23X10^8 IU/mL and genotype was 1b. Hepatobiliary ultrasonography (US) revealed grade 1 hepatosteatosis and a minimal thickening in liver paranchyma. A liver biopsy revealed: a HAI score of 6-18 and fibrosis stage of 5-6. The patient was prescribed peg-IFN alpha 2a 180 mcg/once a week + RBV 1200 mg/day. During follow-up, no significant hematological side effect was observed but he was started on mirtazapin due to a psychiatric adverse event (depression). The treatment was discontinued after 6 months because of partial response. The follow-up data of the patient in 2009 is shown in (Table 1).

During the follow-up period in 2010, treatment with peg-IFN alpha 2b 120mcg/once a week + RBV 1200 mg/day was readministered after his blood tests revealed HCV RNA: 8.01x10^6 IU/mL, ALT: 97 IU/mL, AST: 48 IU/mL. He was prescribed an antidepressant (escitalopram) after his psychiatric evaluation before treatment. Although we obtained a response to the treatment, the treatment was discontinued because the patient was not compliant with the treatment, did not agree to continue using the drugs and also had major depression. The follow up data of the patient in 2010 is presented in (Table 1).

In 2013, the patient was started on peg-IFN alpha 2a 180 mcg/once a week + RBV 1200 mg/day + telaprevir 2250mg/day after his test results revealed HCV RNA: 7.41x10^6 IU/mL, ALT: 30 IU/mL, AST: 30 IU/mL. In the fourth week of this treatment, the following values were obtained: HCV RNA: negative, ALT: 22 IU/mL and AST: 26 IU/mL. Around the 8th week of the treatment the patient presented with itchy rashes on his face, neck, and anterior wall of the forearm (Figure 1). Upon consultation with the dermatology clinics he was initiated urea solution and methylprednisolone aceponate solution. Treatment for HCV was continued without interruption. After 12th administration of the combination therapy, the patient was admitted to our hospital with the complaints of itching, rash, edema generalized to the whole body and facial edema and peeling skin on his face (Figure 2). His physical examination revealed body temperature of 36.7°C, blood pressure of 110/70 mmHg, and heart rate of 80/ min. There were no lymphadenopathy and oral mucosal lesions. All other system evaluations revealed normal findings except for dermatologic lesions. The patient was hospitalized and intravenous hydration was started together with antihistamines and he also continued on his previous lotions to be applied on whole body skin. His blood tests were negative for HCV RNA. Then peg-IFN/RBV+telaprevir treatment was interrupted for a week. Four days after treatment discontinuation, the rashes improved, edema and peeling diminished. One week peg-IFN/RBV treatment without telaprevir was readministered. At 24th week of this treatment the patient was negative for HCV RNA and the treatment was stopped. The followup data for the patient in 2013 is presented in (Table 1).

Discussion

HCV infection causes a variety of dermatological reactions in addition to liver inflammation and fibrosis (7-9). During HCV infection cryoglobulin increases with a ratio of 40%-84% and in 15% of patients cryoglobulenic vasculitis develops. The most common viral cause of porfiria cutanea tarda is HCV. Also an increased incidence of lichen planus is reported in patients infected with HCV compared to those who are not infected. Despite absence of adequate evidence, cutaneous polyarteritis nodosa, psoriasis, urticaria, erythema multiforme can be associated with HCV (7-9).
Mixed cryoglobulinemia, porphyria cutanea tarda, lichen planus and, although without sufficient evidence, cutaneous polyarteritis nodosa, psoriasis, urticaria and erythema multiforme can be associated with HCV infection. Localized reactions associated with the use of peg-IFN and RBV are erythematous or egzematous dermatitis, psoriasis, localized alopecia, skin ulceration and necrosis, local infections at sites of injection, and localized allergic reactions (4,10-12). Generalized reactions include increase in chronic inflammatory diseases of the skin, such as alopecia/abnormal hair growth, dry skin, dermatitis, itching, psoriasis, lichen planus and also immune-related inflammatory diseases and autoimmune diseases like psoriasis and sarcoidosis (4,10,13-15).

When evaluating the treatment of chronic hepatitis C over 22 years, we can observe that the addition of protease inhibitors (telaprevir, boceprevir) to peg-IFN/RBV treatment resulted in a sustained viral response rate of 80% of. In patients infected with HCV genotype 1, the addition of a recently approved protease inhibitor telaprevir to peg-IFN/RBV resulted in sustained viral response rates of approaching 75% for treatment-naive patients and 80% for relapsing patients after initial peg-IFN/RBV. Another protease inhibitor boceprevir resulted in similarly high response rates of 68% and 75% for treatment naive patients and relapsing patients, respectively (16,17).

Protease inhibitors (telaprevir, boceprevir) can provide clinicians with shorter durations of treatment while ensuring higher response rates. Nevertheless, they are associated with more adverse events and drug interactions (11).

While eruptions are seen in 33% of those treated with peg-IFN/RBV this rates are approaching to 50% (90% of these are mild-moderate, 50% occurred in the first 4 weeks) in patients receiving peg-IFN/RBV+telaprevir. Less than 10% of those patients with eruptions progressed to a more severe stage, 0.4% progressed to DRESS (a drug eruption associated with eosinophilia and eruptions progressed to a more severe stage). In our case, telaprevir treatment was discontinued since skin eruption was effecting >50% of body surface area. It is suggested to internalize the patient, discontinue the treatment and then permanent discontinuation of the treatment involving <50% of body surface area; dermatologic consultation, interruption and then permanent discontinuation of the treatment in case of no improvement if it effects >50% of body surface area (18).

In conclusion, since eruptions occur in nearly half of patients receiving telaprevir+peg-IFN/RBV, it is crucial that these patients are followed up for dermatological adverse events. Patients should be advised before the initiation of the treatment to limit exposure to sun/heat and to regularly apply lotions after shower that would prevent dry and itchy skin.

### Conflict of interest: None declared.

### References


### Table 1. Follow-up data during treatment

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