Dermatologic Extrahepatic Manifestations of Hepatitis C

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Abstract

Hepatitis C virus (HCV) affects millions of people worldwide, and an estimated 3.2 million people in the United States. HCV is a hepatotropic and lymphotropic virus that causes not only liver disease, but also a significant number of extrahepatic manifestations (EHMs). Up to 74% of patients affected by HCV will have HCV-related EHMs of some severity in their lifetime. The EHMs vary from simple cutaneous palpable purpura to complex lymphoproliferative disorders, including lymphomas and immune-complex deposit diseases causing local and/or systemic complications. Mixed cryoglobulinemia (MC) is manifested by multiple systemic organ involvement, mainly skin, kidney, peripheral nerves, and salivary glands, and less frequently causes widespread vasculitis and malignant lymphoma. MC affects up to 3% of HCV-infected patients with cryoglobulinemia of clinical significance, i.e. >6%. Severe disease requires immunosuppressive or plasma exchange therapy. HCV prevalence in the United States in patients with porphyria cutanea tarda (PCT) was reported to be 66%, much higher than that in general population. Therefore, all patients with PCT should be screened for HCV. The skin rash of PCT varies from large blisters to small vesicles and/or milia on the hands. Skin manifestations due to PCT usually respond to anti-HCV treatment together with reducing skin sun exposure, avoiding triggers, having routine phlebotomy (especially for people with chronic iron overload states), and using chloroquine. Lichen planus (LP), which typically affects both the skin and oral mucosa is a chronic inflammatory disease of squamous cell origin affecting about 1% of the worldwide population. The prevalence of HCV in patients with LP varies based on geographic location. We review here the basic pathophysiology, clinical features, and management of dermatologic manifestations of HCV.

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Introduction

Hepatitis C virus (HCV) affects approximately 130 to 170 million people worldwide, with an estimated 3.2 million people infected in the United States.1–3 Globally, approximately 350,000 deaths each year are attributed to liver disease.2,2 Hepatitis C was discovered in 1989 and was shown to be a single stranded ribonucleic acid (RNA) virus belonging to a flaviviridae group. HCV is a hepatotropic and lymphotropic virus that causes not only hepatic manifestations, but also a significant number of extra-hepatic manifestations (EHMs). Approximately 74% of patients with hepatitis C will have HCV-related EHMs of some severity in their lifetime.4 The development of various EHMs by HCV likely involves autoimmune mechanisms, as evidenced by the appearance of autoimmun features, such as palpable purpura, complex lymphoproliferative disorders (e.g. lymphomas), and immune-complex deposit diseases that cause local and/or systemic complications.4,5 Among the EHMs, dermatologic manifestations significantly add to morbidity and the overall cost burden on the health care system.6 This article focuses on the basic pathophysiology, clinical features, and management of the most common dermatologic manifestations of HCV.

Classification of EHMs associated with HCV

Classification of EHMs associated with HCV is based on the available literature (Table 1).

Mixed cryoglobulinemia (MC)

Introduction

MC is the most common dermatologic EHM and was first described in 1966.7

Pathophysiology

MC is a systemic vasculitis caused by the deposition of circulating immune complexes in the small vessels. It is manifested by multiple organ involvement, mainly skin, kidney, peripheral nerves, and salivary glands, and less frequently causes widespread vasculitis and malignant lymphoma.1 The vasculitis is associated with the presence of serum cryoglobulins (CG) which are insoluble at temperatures below 37 °C, but can dissolve by warming are responsible for MC (Fig. 1).

Classification

According to Brouet et al.,8 CG can be classified based on the type of immune complex deposition. Type I is purely

Keywords: Dermatologic manifestations; Extra-hepatic manifestations; Hepatitis C virus; Cryoglobulinemia; Porphyria cutanea tarda; Lichen planus.

Abbreviations: AASLD, American Association for the Study of Liver Disease; CG, cryoglobulins; DAA, directly acting antiviral; EHMs, extra-hepatic manifestations; HCV, hepatitis C virus; HFE, High Iron Fe; IFN, interferon; Ig, immunoglobulin; LP, lichen planus; MC, mixed cryoglobulinemia; MCS, MC syndrome; MPGN, membranoproliferative glomerulonephritis; PCT, porphyria cutanea tarda; RF, rheumatoid factor; RNA, ribonucleic acid; SVR, sustained virologic response; UROD, uroporphyringen decarboxylase.

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monoclonal components. Type II is a mixture of polyclonal immunoglobulin IgG and monoclonal IgM, while Type III is a mixture of polyclonal IgG and polyclonal IgM. In MC, the IgM consists of autoantibodies with rheumatoid factor (RF) activity. Type II CG accounts for 50–60% of CGs, and Type III CG account for about 40–50% of CGs.1

Clinical features

The most common symptoms of MC syndrome (MCS) are weakness, arthralgias, and purpura, also called Meltzer’s stria.9 The dermatologic manifestations of MC can vary from simple cutaneous palpable purpura (Fig. 2A) to complex confluent lesions (Fig. 2B), including ulcerations of the skin (Fig. 2C).

Laboratory findings

Ninety percent of patients with MC have anti-HCV AB.10 The presence of serum mixed CGs, high RF levels, and reduced complement C4 levels are all consistent with MC. It may be difficult to detect CGs because of their thermolability and the variability of the rate in which CGs are responsible for vasculitic damage.7,11

Treatment and discussion

The frequency of cryoglobulinemia in HCV-positive patients varies. This variability might be related to the duration of HCV infection and the stage of liver fibrosis.7 Among all causes of MC, 80% of MC is due to HCV, and it is usually associated with Type II CG,12 less commonly with Type III, and rarely with Type I. Patients with chronic hepatitis C and CG can present at various stages of liver fibrosis, but a higher incidence of cirrhosis and stage of fibrosis has been reported in patients with significant cryoglobulinemia (>6%).1,13 Twenty percent of patients with HCV have MC, but most had <6% CG, which is not clinically significant. Only 3% of HCV patients had clinically significant MCS with >6% levels of CG.7 The mean cryocrit levels in HCV patients was 2%.14 Renal involvement due to MCS and age >60 at the time of diagnosis have been linked to worse outcomes.15 Renal involvement due to cryoglobulinemia and HCV tends to present with membranoproliferative glomerulonephritis (MPGN).12,16 Severe, life threatening complications were observed in up to 10% of the patients with MC, and the mortality ranged from 20-80%. Mortality increased with systemic involvement, especially when there was evidence of pulmonary hemorrhage, gastrointestinal ischemia, or cardiac or central nervous system involvement.7,15,17,18

An algorithm for the treatment for MCS associated with HCV is depicted in Fig. 3. In the past, MC due to HCV was treated with standard anti-HCV therapy plus ribavirin. However, severe disease may require immunosuppressive or plasma exchange therapy. Because of the lymphotropic nature4 of HCV, MC affects B cell lymphocytes in peripheral blood, liver infiltrates, and bone marrow. Rituximab (monoclonal antibody against CD20-expressing B cells) has been used in refractory MCS with or without multisystem involvement.
The strategy is to decrease the expansion of B cells, a likely source of CG that failed to respond to standard anti-HCV treatment.5 Because interferon (IFN) may stimulate the immune system, thereby causing further damage in severe or advanced cases of MCS, IFN-free anti-HCV regimens are recommended.19 A thorough review of the literature has failed to identify immunohistochemistry studies regarding the change in skin lesion status after IFN therapy. However, a recent study by Saadoun et al. showed that the use of combination pegylated IFNα with ribavirin and a protease inhibitor seemed to be highly effective in treating HCV-related MC.20 Dermatologic manifestations of MC respond to the treatment and clearance of HCV.21 Patients who relapse after initial clearance of HCV usually have relapses in their vasculitis that parallel the return of viremia.20

In summary, treatment of HCV should not be deferred due to the presence of MC and vasculitis.22 Because of the involvement of HCV in the development of MC, treatment of HCV alone generally improves mild to moderate MC. Because of the potential for exacerbating MC, IFN-based regimens are not recommended as anti-HCV treatment. When an IFN-based anti-HCV regimen is required, concurrent immunosuppression may be necessary. In severe MC with multisystem involvement, immunosuppressive therapy is recommended with rituximab-based regimens with or without plasmapheresis prior to initiation of anti-HCV therapy.

Porphyria cutanea tarda (PCT)

Introduction

Porphyrias are a group of rare inherited or acquired disorders of enzymes that normally participate in the production of heme and porphyrins. Porphyrias generally manifest with either neurological or skin problems or occasionally both. The term ‘porphyria’ is of Greek origin and meaning purplish pigment, relating to the color of porphyrins.

Classification

There are many types of porphyrias, including PCT, hereditary coproporphyria, and variegate porphyria. PCT is the most common type of porphyria and is further categorized in two different subtypes: familial and sporadic.23 In the familial form, the enzyme defect is present in hepatocytes and other cells, such as erythrocytes; whereas in the sporadic (more...
common) form, the enzyme activity is decreased to 50%, affecting predominantly hepatocytes.24

**Pathophysiology**

There are three possible mechanisms by which PCT may develop. The basic defect in PCT is either decreased or absent activity of the enzyme uroporphyrinogen decarboxylase (UROD). Since UROD normally converts uroporphyrinogen into coproporphyrinogen in the heme-biosynthetic pathway, disruption of this pathway may cause PCT.14,15 The second mechanism is an increase in the rate or tendency of uroporphyrinogen to be oxidized to uroporphyrin. The rate of oxidation is influenced by many factors, including chronic hepatitis C, long-term alcohol intake, cytochrome P-4501A2, metabolically active iron within the hepatocytes, and estrogens. This mechanism is likely the key link among PCT, chronic liver disease, and iron overload states. The third mechanism is an increase in activity of hepatic 5-aminolevulinic acid synthase, the first and normally the rate-controlling enzyme of heme synthesis. An increase in activity of this enzyme can result in increased levels of uroporphyrinogen in hepatocytes. Oxidation of uroporphyrinogen to uroporphyrin and other nonporphyrin products can inhibit UROD.9,25

HCV increases iron stores, which in turn inhibit UROD. This occurs when the activity of UROD decreases to <25% of normal levels and results in an accumulation of uroporphyrin and other carboxylated porphyrins in various organs, including the skin and liver. This increase of porphyrins predisposes skin to a photochemical reaction upon exposure to sunlight and causes the typical skin manifestations of PCT. Iron also increases the oxidation of uroporphyrinogen to uroporphomethene, which decreases UROD by competitive inhibition. In the presence of uroporphyrin, photo-activation of the complement system leads to activation of mast cells and proteases, and this causing dermal-epidermal splitting (Fig. 4).25,26

**Clinical features**

PCT is frequently found in middle-aged men who have a history of chronic liver disease either due to chronic iron overload states, heavy alcohol abuse, or HCV. The earliest manifestation of PCT is a skin rash that varies from large blisters, small vesicles, and/or milia on the dorsal aspect of the hands (Fig. 5).27 Other manifestations include increased skin fragility, hypertrichosis (especially involving lateral aspect of the face), chronic hyper/hypopigmentation, chloracne, sclerodermoid changes, dystrophic calcifications with ulcerations, scarring, alopecia, and onycholysis. Patients can also manifest stigmata of advanced liver disease and cirrhosis if the latter is present.

**Treatment**

Treatment consists of controlling sun exposure of the skin, avoiding triggers, getting routine phlebotomy (especially in cases of chronic iron overload), and using chloroquine. Patients with PCT should stop alcohol consumption and estrogen and iron supplementation. For patients with underlying HCV manifesting with PCT, treating the underlying cause with IFN-based anti-HCV treatment has been reported to result in resolution of PCT lesions. Currently, however, there are no clinical trials or data available on whether treatment of HCV infection and achievement of sustained virologic response (SVR) with directly acting antiviral (DAA) agents improves PCT. The cornerstone of the treatment still largely depends on routine phlebotomy and controlling the triggers. Based on the American Association for the Study of Liver Disease (AASLD) guidelines for HCV treatment, PCT is class IIB, level C rating, and newer DAAs are recommended for this condition. Based on the guidelines, achievement of an SVR with DAAs should result in resolution of PCT-related dermatological manifestations.

**Discussion**

In a meta-analysis by Gisbert et al. that included 50 studies (2,167 patients), the mean prevalence of HCV antibodies and positive HCV RNA in patients with PCT was estimated to be 47% (95%CI =45-49%) and 50% (95%CI =47-53%), respectively. Furthermore, when considering only case-control studies in which anti-HCV antibodies and/or HCV RNA by polymerase chain reaction were performed, a tight
association was found between PCT and HCV infection (OR = 82). The prevalence of HCV infection was significantly higher in the sporadic form of PCT (57%, 95% CI = 57-75%) than in the familial form (26%, 95% CI = 17-34%). HCV prevalence in patients in the United States with PCT was reported to be approximately 66%, much higher than that reported in the general population. Therefore, all patients with PCT should be screened for HCV. However, among the HCV-infected population, the overall prevalence of PCT was estimated to be less than 5% and as low as 1%. Since the chance of having HCV infection is higher in patients with PCT, all patients with PCT should be screened for HCV infection. The prevalence of HFE gene mutations in North American patients with PCT is high. It is, therefore, recommended to screen for HFE mutations and HCV infection in patients who present with PCT.

To summarize, patients who are diagnosed with PCT should be screened for HCV because of the association between the two. Only 1-5% of patients with HCV will have evidence of PCT. The skin lesions of PCT respond to standard anti-HCV and PCT therapies.

**Lichen planus**

**Introduction**

Lichen planus (LP) is a chronic inflammatory disease that affects skin and mucous membranes of squamous cell origin. LP affects about 1% of the general population, and it is a common disorder affecting stratified squamous epithelia, particularly of the oral cavity.

**Pathophysiology**

Although the exact pathophysiology of LP remains unknown, one potential mechanism is induction of keratinocyte apoptosis by cytotoxic CD8+T cells, stimulated by unidentified triggers or autoimmune antibodies. The histopathological features of dermal LP lesions exhibit an irregular sawtooth appearance suggestive of epidermal hyperplasia (Fig. 6), deep compact hyperkeratosis with wedge-shaped thickening of the granular cell layer, and a dense band-like T-cell infiltrate obscuring the dermal-epidermal junction.

![Fig. 6. Histopathology of lichen planus. Black solid arrow showing a saw tooth pattern of epidermal hyperplasia, dashed arrow showing band of T cell infiltrate.](image)

**Clinical features**

LP typically affects women between the ages of 30 and 60. Lesions in the mucous membrane usually present as white lace-like Wickham’s striae, which are found in the lateral buccal mucosa, involving sometimes the lips, gingivae, and tongue (Fig. 7A). On the skin, LP usually presents as purplish, flat-topped papules and intensely pruritic, polygonal lesions on the flexor aspect of wrists and forearms, extensor aspect of hands and ankles, lumbar region, shins, and genital area (Fig. 7B).

**Treatment**

Symptoms of LP are usually treated with either topical or injectable steroids. To date, data on the response to IFN treatment are inconsistent, as both improvement and exacerbation of symptoms have been reported. There are strong data demonstrating that treatment of underlying HCV with anti-HCV treatment, especially IFN-based regimens, may not lead to regression of all LP lesions. At this time, there are no convincing data on the response of LP to DAAs. Other treatment options have been reported, such as topical calcineurin inhibitors, phototherapy, systemic retinoids, and methotrexate. For pruritus due to the dermal lesions, oral antihistamines have been reported to provide symptomatic relief.

**Discussion**

A potential connection between hepatitis viruses and LP is shown by the frequent association between LP and chronic liver disease. Given the odds ratio and prevalence figures in various countries, it has been estimated that the prevalence of HCV in patients with LP varies from 4% in Europe to 24% in the Middle East. With the advent of sensitive HCV diagnostic tests, a rise in the number of case reports, cohort, and controlled studies occurred, further supporting a link between LP and HCV. A study compared the prevalence of HCV antibodies in 263 patients with oral LP to a control group of 100 patients who were receiving routine dental care. Twenty-nine percent of those with LP tested positive for HCV antibodies, compared to only 3% in the control group. However, others have shown that 2% of 127 HCV-positive patients had oral LP, and only one in 24 consecutive patients with LP was anti-HCV seropositive. A recent study suggested a weak or no association between HCV infection and LP. The possible explanation for this significant difference could be the low endemicity...
of HCV in India and differences in HCV genotype (3a and 3b are more common in India, while genotypes 1 and 2 are more common in America and Europe).42

In summary, HCV infection contributes to almost 2% of all LP cases, but routine HCV testing of all LP patients is not indicated.43 The main reason for identifying and treating LP lesions is to prevent malignant transformation of the lesions.31 There has been evidence to suggest that low levels of vitamin D in patients with chronic HCV infections may lead to a higher incidence of EHMs.44

Conclusions

HCV causes numerous hepatic as well EHMs. Treating the underlying HCV with anti-HCV therapy, ribavirin with or without interferon has led to significant improvements in some EHMs.

Conflict of interest

None.

Author contributions

Writing the article (BD), conceiving the idea and editing the manuscript (GYW).

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