Human hepatitis viruses-associated cutaneous and systemic vasculitis

Chrong-Reen Wang, Hung-Wen Tsai

ORCID number: Chrong-Reen Wang 0000-0001-9881-7024; Hung-Wen Tsai 0000-0001-9223-2535.

Author contributions: Wang CR designed the report and wrote the paper; Wang CR and Tsai HW collected and analyzed the clinical data.

Conflict-of-interest statement: The authors declare no conflicts of interest.

Open-Access: This article is an open-access article that was selected by an in-house editor and fully peer-reviewed by external reviewers. It is distributed in accordance with the Creative Commons Attribution NonCommercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited and the use is non-commercial. See: http://creativecommons.org/Licenses/by-nc/4.0/

Manuscript source: Invited manuscript

Specialty type: Gastroenterology and hepatology

Country/Territory of origin: Taiwan

Peer-review report’s scientific

Abstract

Human hepatitis viruses (HHVs) include hepatitis A virus, hepatitis B virus (HBV), hepatitis C virus (HCV), hepatitis delta virus, and hepatitis E virus and can cause liver inflammation in their common human host. Usually, HHV is rapidly cleared by the immune system, following acute HHV invasion. The morbidities associated with hepatitis A virus and hepatitis E virus infection occur shortly after their intrusion, in the acute stage. Nevertheless, the viral infectious process can persist for a long period of time, especially in HBV and HCV infection, leading to chronic hepatitis and further progressing to hepatic cirrhosis and liver cancer. HHV infection brings about complications in other organs, and both acute and chronic hepatitis have been associated with clinical presentations outside the liver. Vascular involvement with cutaneous and systemic vasculitis is a well-known extrahepatic presentation; moreover, there is growing evidence for a possible causal relationship between viral pathogens and vasculitis. Except for hepatitis delta virus, other HHVs have participated in the etiopathogenesis of cutaneous and systemic vasculitis via different mechanisms, including direct viral invasion of vascular endothelial cells, immune complex-mediated vessel wall damage, and autoimmune responses with stimulation of autoreactive B-cells and impaired regulatory T-cells. Cryoglobulinemic vasculitis and polyarteritis nodosa are recognized for their association with chronic HHV infection. Although therapeutic guidelines for HHV-associated vasculitis have not yet been established, antiviral therapy should be initiated in HBV and HCV-related systemic vasculitis in addition to the use of corticosteroids. Plasma exchange and/or combined cyclophosphamide and corticosteroid therapy can be considered in patients with severe life-threatening vasculitis manifestations.

Key Words: Human hepatitis viruses; Hepatitis B virus; Hepatitis C virus; Cryoglobulinemic vasculitis; Polyarteritis nodosa; Antiviral therapy
INTRODUCTION

Hepatocellular injury caused by acute or chronic inflammation of the liver is derived from adverse factors, including alcohol consumption, autoimmune response, drug use, steatosis, and viruses[1]. Hepatotropic viruses are known for their role in the etiology of viral hepatitis, and, from the public health perspective, the disease is associated with heavy health burden and higher annual mortality[4]. In addition to uncommon herpes viruses-related hepatitis due to Epstein-Barr virus, cytomegalovirus, and herpes simplex virus infection[4], human hepatitis viruses (HHVs) are the most common causes of acute and chronic hepatitis in their human host[4]. The discovery of HHV starts with clinical description of disease, antigen/antibody reaction establishment, virus-like particles visualization, and finally viral genomes sequencing[4]. Early experiments illustrated two types, infectious hepatitis and serum hepatitis[4]. Australia antigens in serum hepatitis yielded the detection of 42-nm Dane particles[24]. Subsequently, deoxyribonucleic acid (DNA) sequencing was performed and hepatitis B virus (HBV) was found[4]. Later, virus-like particles were observed in infectious hepatitis, and complementary DNA (cDNA) from hepatitis A virus (HAV) was sequenced[23-24]. Hepatitis C virus (HCV) was identified in non-A, non-B hepatitis patients receiving multiple transfusions[25], transmission to chimpanzees[26], and cDNA library screens[4]. Hepatitis delta virus (HDV), a HBV-associated virus[27], is a separate defective virus requiring the help of HBV for its infection[28]. After elucidating the hepatitis E virus (HEV) cDNA[29], an endemic non-A, non-B hepatitis[30], there are five members of HHVs[3].

Typically following HHV invasion into humans, there is rapid clearance of viruses by the host defense system, with a self-limiting disease course[4]. The morbidities associated with HAV and HEV infection, commonly transmitted via the fecal-oral route, occur shortly after their intrusion in the acute stage[18-19]. The infectious processes can persist for a long period of time in HBV, HCV, and HDV infection, progressing to chronic hepatitis and leading to liver fibrosis, irreversible cirrhosis, and hepatocellular carcinoma[25-29]. Table 1 summarizes and compares the common characteristics of the five HHV members[21-24].

EXTRAHEPATIC MANIFESTATIONS OF HUMAN HEPATITIS VIRUSES INFECTION

Although HHVs primarily affect hepatocytes, they can cause extrahepatic manifestations, with damage to other organs[19-24]. Both acute and chronic forms of viral hepatitis are associated with various clinical presentations outside of the liver, which may precede or follow hepatic dysfunction. The crucial pathogenic mechanism is caused by the immune responses against viral pathogens, with the deposition of immune complexes (IC) in targeted tissues.
### Table 1 Common characteristics and comparative features in five members of human hepatitis viruses

|                | HAV | HBV | HCV | HDV | HEV |
|----------------|-----|-----|-----|-----|-----|
| **Family; Genus** | Picornaviridae Hepatitisvirus | Hepadnaviridae Orthohepadnavirus | Flaviviridae Hepacivirus | Deltaviridae Deltavirus | Hepeviridae Orthohepeviruses |
| **Discovery** | 1983 Highest in AH | 1979 Highest in CH | 1989 Second in CH | 1986 Less in CH | 1990 Less in CH |
| **Frequency** | | | | | |
| **Genome** | Linear ssRNA; 7500 nt | Circular dsDNA; 3200 nt | Linear ssRNA; 9600 nt | Circular ssRNA; 1700 nt | Linear ssRNA; 7200 nt |
| **Length** | 27-32 nm | Quasi-envelope; Unknown | 42 nm | Envelope; HTPC, HSP | 36-43 nm; Envelope; NTCP, HSP |
| **Envelope** | Unknown | Quasi-envelope; HTPC, HSP | Unknown | HTPC, HSP | Unknown |
| **Diameter** | 55-65 nm | Envelope; CD81, SR-BI | 30-34 nm | Quasi-envelope; Unknown | |
| **Receptor** | HSP | NTCP, HSP | 55-65 nm | Envelope; CD81, SR-BI | |
| **Incubation; Age** | 14-30 d; Children; Acute | 30-180 d; Any; Acute/chronic | 14-180 d; Any; Acute/chronic | 14-160 d; Any; Acute/chronic | 14-70 d; Adults; Acute |
| **Course** | Enteric, sexual, vertical | Parental, sexual, vertical | Parental, sexual, vertical | Parental, sexual, vertical | Enteric, vertical, parenteral |
| **Route of spread** | | | | | |
| **Viruses** | Anti-HAV IgM | Anti-HBc IgM, HBs antigen, DNA by PCR | Anti-HCV IgG, RNA by PCR | Anti-HDV IgM, RNA by PCR | Anti-HEV IgM, RNA by PCR |
| **Vaccine; Post-exp; Av agent** | Available; Ig/vaccine; Not available | Available; Ig/vaccine; NA, IFN-a | Not available; Not effective; DAA agent | HBV vaccine; Not available; IFN-a | Available in China; Not available; Not available |
| **FH; LC/HCC; Prognosis** | Very rare; Nil; Full recovery | Very rare; Yes; Chronic carrier | Extremely rare; Yes; High carrier rate | Yes; Yes; Chronic carrier | Yes; Nil; Full recovery |

1 Co-infection/superinfection with hepatitis B virus.
2 By cloning and sequencing.
3 Partial dsRNA in hepatitis B virus.
4 Diameter of virion.
5 Quasi-envelope with internal protein rather than viral glycoprotein.
6 Commonly affected age.
7 Progress to chronicity in 80% superinfection/5% co-infection hepatitis B virus.
8 Up to 20% in superinfection with hepatitis B virus and pregnant women.
9 Poorer in hepatitis B virus superinfection with hepatitis delta virus and pregnant women with hepatitis E virus. AH: Acute hepatitis; Av: Antiviral; CH: Chronic hepatitis; DAA: Direct-acting antiviral; DNA: Deoxyribonucleic acid; ds: Double-stranded; Post-exp: Post-exposure; FH: Fulminant hepatitis; HBc: Hepatitis B virus core antigen; HBs: Hepatitis B virus surface antigen; HCC: Hepatocellular carcinoma; HSP: Heparan sulfate proteoglycans; IFN: Interferon; Ig: Immunoglobulin; LC: Liver cirrhosis; NA: Nucleoside/nucleotide analogues; nt: Nucleotide; NTCP: Sodium taurocholate co-transporting polypeptide; PCR: Polymerase chain reaction; RNA: Ribonucleic acid; SR-BI: Scavenger receptor class B type I; ss: Single-stranded.

Extrahepatic manifestations with evanescent rash and transient arthralgia that develop in acute HAV infection are rare. In the protracted clinical course with cholestasis or relapsing disease, cutaneous vasculitis and arthritis occur with a predilection for lower extremities, and usually there is spontaneous recovery after the clearance of HAV. Other infrequently observed presentations include glomerulonephritis, myocarditis, thrombocytopenia, and neurological complications like Guillain-Barre syndrome. Extrahepatic infection can be asymptomatic or manifest as fulminant hepatitis. Neurological involvement is the most frequently encountered extrahepatic disorder, followed by hematological and gastrointestinal manifestations, possibly caused by autoimmune responses related to molecular mimicry. Neuralgic amyotrophy and Guillain-Barre syndrome are two common neuromuscular disorders, and other uncommon presentations include mononeuritis multiplex, meningoradiculitis, and encephalitis. Thrombocytopenia is a frequently identified hematological disorder, and pancreatitis is the most common gastrointestinal abnormality. Myocarditis and glomerulonephritis are rarely observed complications. Although there is evidence to support the treatment of HEV-related neurological manifestations with corticosteroids, the currently recommended therapy is plasma exchange and immune modulation with intravenous immunoglobulin. Although extrahepatic manifestations exist with acute infection of all HHVs, such presentations are more commonly identified in HBV, with up to 20% occurrence. Acute HBV infection is often subclinical and asymptomatic in around two-thirds of cases. In the pre-icteric prodromal phase, there is serum sickness-like illness with arthritis and dermatitis due to the deposition of circulating IC composed of HBV surface (HBs) antigen and further activation of complements. These manifestations are usually transient and are resolved after the onset of jaundice. Guillain-Barre syndrome has been reported to be associated with acute HBV infection, and both...
plasma exchange and intravenous immunoglobulin are effective treatments,
implying that there is autoimmune-mediated damage to the myelin sheath of peripheral nerves.

Owing to the availability of viral markers and our understanding of pathogenic pathways, extrahepatic manifestations of chronic HBV infection have been well-elucidated for a while\[^{76}\]. The etiopathogenesis outside the liver in the chronic phase involves the deposition of IC comprised of HBs and/or HBe antigens, followed by the local activation of complement cascades and the recruitment of inflammatory cells\[^{61}\]. Notably, higher viral load or persistent infection can promote the production of IC, leading to deposition at small or medium-sized arteries\[^{62}\]. In addition, viral replication has been demonstrated in the endothelium of targeted vessels\[^{63}\]. Taken together, both mechanisms suggest that inhibition of viral replication, either spontaneously or under antiviral therapy, can reduce extrahepatic manifestations. Up to one-fifth of victims with chronic HBV infection have morbidities outside the liver\[^{64}\], and these are comprised of arthritis, glomerulonephritis, uveitis, peripheral neuropathy, Raynaud phenomenon, Sjögren syndrome, cutaneous vasculitis, and systemic vasculitis, including polyarteritis nodosa and cryoglobulinemic vasculitis\[^{65-67}\]. Since the administration of immunosuppressive agents increases the risk of additional hepatic HBV replication with worsening liver disease\[^{68}\], the treatment of HBV-associated glomerulonephritis and vasculitis is mainly based on antiviral agents, interferon (IFN)-α or nucleoside/nucleotide analogues (NAs)\[^{69-71}\].

In most cases, symptomatic manifestations are uncommon during acute HCV infection\[^{72}\]. Extrahepatic presentations include arthralgia, myalgia, and rash\[^{73}\]. Patients with HCV infection usually progress to chronic stage without recovery\[^{74-76}\]. Nevertheless, antiviral therapy is highly effective for the acute infection, resulting in the clearance of HCV with sustained virological response (SVR)\[^{77}\]. Persistent HCV infection is a leading cause of chronic liver disease\[^{78}\]. Although essentially curable with direct-acting antiviral (DAA) therapy, only a portion of patients are diagnosed. Notably, extrahepatic manifestations occur in up to three-quarter of victims with chronic HCV infection\[^{79}\], and cryoglobulinemia is the most frequently encountered presentation (in 40% to 60% of infected patients)\[^{80}\]. Direct viral invasion and deposits of soluble IC are two pathogenic processes involved in the development of disease outside the liver\[^{81-83}\]. The clinical presentations include arthralgia, myalgia, glomerulonephritis, Raynaud phenomenon, Sjögren’s syndrome, Hashimoto’s thyroiditis, Graves’ disease, ulcerative keratitis, peripheral neuropathy, and cryoglobulinemia vasculitis\[^{84-86}\]. Occasionally, extrahepatic-related autoimmune comorbidities such as cryoglobulinemia vasculitis can lead to the diagnosis of HCV infection\[^{87}\]. Sustained eradication of HCV by IFN-α or DAA agents has shown to be beneficial for outcomes following these manifestations\[^{88,89}\]. Notably, a prospective study with 9895 HCV-infected cases receiving DAA medications revealed that viral clearance is responsible for a significant decrease in HCV extrahepatic mortality\[^{90}\].

**VASCULITIS MANIFESTATION IN HUMAN HEPATITIS VIRUSES INFECTION**

There is growing evidence demonstrating a causal relationship between viral pathogens and vasculitis. Except for HDV\[^{91,92}\], other HHVs have been shown to participate in the etiopathogenesis of cutaneous and systemic vasculitis via different mechanisms, including direct viral invasion of vascular endothelium, IC-mediated vessel wall damage, and autoimmune responses with stimulation of autoreactive B-cells and impaired regulatory T cells\[^{93-95}\].

Cutaneous leukocytoclastic vasculitis (CLV) rarely occurs during acute HAV infection, and it may resolve after the regression of hepatitis\[^{96-98}\]. In chronic HAV infection, CLV is rarely observed, with an 1% incidence\[^{99}\]. Hbs antigen has been identified in affected skin lesions\[^{100}\]. Notably, IFN-α therapy is effective against HBV-associated CLV\[^{80}\]. The histopathological picture of HHV-triggered CLV shows relatively less eosinophils and lymphocytes compared with those in drug-induced CLV\[^{101}\].

Acute HAV or HEV infection can induce Henoch-Schönlein purpura (referred to as HSP)\[^{102-106}\], a systemic vasculitis caused by the widespread deposition of circulating immunoglobulin (Ig)A IC in small-sized blood vessels of skin, joint, lung, kidney, testis, gastrointestinal tract, and nervous system\[^{107}\]. HAV- and HEV-related HSP usually have a spontaneous recovery. Owing to the defective liver catabolism of IgA IC with further tissue deposits, HSP occurs in HBV or HCV chronic liver diseases,
usually requiring antiviral and corticosteroid therapy\[^{94-96}\].

Vascular involvement with systemic vasculitis in HHV infection is a recognized extrahepatic morbidity with potential life-threatening organ dysfunction\[^{97,98}\]. Among them, cryoglobulinemic vasculitis and polyarteritis nodosa are well-known vascular comorbidities in HHV infection\[^{99,100}\].

### HUMAN HEPATITIS VIRUSES-ASSOCIATED CRYOGLOBULINEMIC VASCULITIS

Cryoglobulinemic vasculitis is caused by IC-mediated inflammation of small-sized blood vessels and is accompanied by the activation of complements\[^{101}\]. Cryoglobulins are circulating antibodies that precipitate \textit{in vitro} at temperatures less than \textit{37 °C} and dissolve after rewarming, and these are either Ig or a mixture of Ig and complement components\[^{102}\]. Individuals with cryoglobulinemia may remain asymptomatic without clinical abnormalities\[^{103,104}\]. A typical triad of purpura, arthralgia, and weakness associated with organic dysfunction and elevated levels of rheumatoid factor (RF) was defined as cryoglobulinemic disease in 1966 by Meltzer \textit{et al}\[^{105}\]. Based on the composition of Ig, cryoglobulinemia can be classified into type I with monoclonal Ig (usually IgM), type II with polyclonal IgG and monoclonal IgM RF, and type III with polyclonal IgG and polyclonal IgM RF\[^{106,107}\]. Type I is an infrequently encountered subgroup and is only found in hematological malignancies\[^{108}\], whereas mixed types II and III can be associated with viral hepatitis caused by HBV, most commonly observed with HCV, and rarely identified with HAV infection\[^{96,109,110}\]. Despite the presence of cryoglobulins in up to 60% of chronic HCV patients, cryoglobulinemic vasculitis only appears in around 10% of cases\[^{111}\]. In chronic HHV infection, envelope glycoproteins can help the virus enter intrahepatic and circulating B lymphocytes to produce Ig with RF activity, resulting in the generation of cryoglobulins and formation of IC with vascular wall deposition\[^{102,104}\].

Recurrent palpable purpura with the characteristic histopathological finding of leukocytoclastic vasculitis is the most common clinical manifestation\[^{112}\]. Other cutaneous presentations are ischemic ulcers, digital gangrene, and Raynaud’s phenomenon. In Figure 1, the presence of leukocytoclastic vasculitis in biopsied skin lesions from a HCV-infected patient with cryoglobulinemia-associated purpura is shown. In addition, the presence of Meltzer triad was identified in most patients at the onset of disease\[^{113}\] and skin, joint, kidney, and peripheral nerve are frequently affected organs in cryoglobulinemic vasculitis\[^{86,106}\]. Renal involvement with membranoproliferative glomerulonephritis as the most common finding is associated with significant morbidity and even mortality\[^{109}\]. HCV core protein and Ig are major components of IC and are distributed along the capillary walls of glomeruli\[^{114}\]. Peripheral neuropathy with mononeuritis multiplex is commonly observed, which can be the first clinical manifestation\[^{115}\]. Other rarely involved organs include the central nervous system, gastrointestinal tract, myocardium, and lung, and their involvement leads to the clinical presentations of impaired cognitive function, intestinal ischemia, hypertrophic cardiomyopathy, and diffuse alveolar hemorrhage\[^{86,116,117}\].

Sporadic cases with cryoglobulinemic vasculitis, either in adults or children, have been observed during acute HAV infection\[^{94,118-120,121}\]. These patients had biopsy-proven CLV plus arthritis or glomerulonephritis, and the use of corticosteroids were required to control these complications.

Based on clinical observations, the Meltzer triad had been regarded as an IC-mediated disease secondary to the HBV infection\[^{122}\]. Since then, HBs antigen in biopsied vessel wall and HBs antigen/anti-HBs in circulating cryoprecipitates have been identified in cryoglobulinemic vasculitides\[^{123}\]. A retrospective study from 913 cases with cryoglobulinemia demonstrated a HBs antigen positivity of 5.8\%\[^{124}\], whereas analyzed data from 231 patients with mixed cryoglobulinemia only revealed an HBV infection rate of 1.8\%\[^{125}\]. Nevertheless, HBV is among three common infectious etiologies in patients with cryoglobulinemia\[^{126}\]. This virus is predominantly associated with type II cryoglobulinemia, and its common extrahepatic manifestations are purpura, arthralgia, and neuropathy\[^{127}\]. Furthermore, compared to adults, children with symptomatic cryoglobulinemia have a lower occurrence of HBV infection but higher frequencies of articular and cutaneous involvement\[^{128}\]. Notably, the presence of HBs antigen represents one of the main independent predictors of mortality in cryoglobulinemic vasculitis\[^{129}\].

Accumulated evidence indicates that the use of NAs for viral suppression in HBV-related cryoglobulinemic vasculitis can cause the disappearance of cryoglobulins,
normalize liver function, and significantly clinically improve most patients with mild disease, indicating a role for HBV replication in the pathogenesis\cite{70,123}. Antiviral therapy with NAs should be a life-long prescription, and discontinuation is considered only after persistent HBs antigen loss with the seroconversion and undetectable HBV DNA\cite{70,123}. Despite an observation with ineffective responses\cite{70}, IFN-α can be an alternative therapeutic modality\cite{124}. The use of corticosteroids without NA therapy is ineffective for suppression of HBV viremia, resulting in refractory or relapsing disease\cite{125}. Nevertheless, only a portion of patients with severe disease, including peripheral neuropathy and renal involvement, can achieve clinical regression under NA therapy\cite{70,123,128}. CD20-positive B-cells are expanded and activated in mixed cryoglobulinemia, and they participate in the production of cryoglobulins\cite{129}. Clinical remission has been reported in patients with glomerulonephritis treated with NA in combination with rituximab, a monoclonal antibody against CD20 on the surface of B-cells\cite{70,128}. Considering the risk of viral reactivation in patients with positive HBs or an occult infection with negative HBs and positive anti-HBc, prophylaxis with NAs should be initiated before the rituximab treatment\cite{70,124,129}. Because of a potentially fatal complication, its use should be avoided during an active flare of HBV infection\cite{70,129}. Prescription of this biologic as a second-line agent can be considered in patients with severe disease refractory to NA therapy\cite{70,123}.

Among the underlying diseases in cryoglobulinemia, HCV infection is the most common cause, with a 73% positive frequency of anti-HCV and a 86% occurrence of HCV ribonucleic acid\cite{70,123,130,131}. Nevertheless, only a small portion of cryoglobulinemia with HCV infection develops significant vasculitis manifestations\cite{132}. Similar to HBV infection, HCV is mainly associated with type II cryoglobulinemia\cite{133}. The interaction of HCV envelope proteins with CD81 surface receptor can stimulate B-cells to expand clonally and produce monoclonal IgM RF that binds polyclonal anti-HCV core antigen, which suggests that cryoglobulinemia is due to host immune responses against HCV infection\cite{134,135}. Although the HCV-related mixed cryoglobulinemia is a benign B-cell lymphoproliferative condition, chronic viral stimulation on the immune system leads to the selection of abnormal clones\cite{136}. An epidemiological survey elucidated a link between HCV infection and B-cell non-Hodgkin’s lymphoma (NHL)\cite{137,138}. A large-scale cohort with 146394 HCV-infected patients demonstrated a more than 20% increased risk of this malignancy\cite{139,140}. Furthermore, the overall risk of B-cell NHL in mixed cryoglobulinemia patients was estimated to be 35 times higher than that in the general population\cite{141}. Collectively, these observations have indicated a pathogenic role of HCV in B-cell NHL.

Since HCV-related cryoglobulinemic vasculitis is an antigen-driven process and its activity usually correlates with viremia, the most effective treatment is the eradication of underlying viral infection\cite{142}. Anti-HCV therapy can follow the existing guideline as antiviral drug\cite{143}. DAA agents alone induce SVR with less adverse events and are more effective than the combined IFN-α and ribavirin regimen in cryoglobulinemic vasculitis\cite{144}. A prospective multicenter study carried out in these patients, all achieved SVR with a 90% complete clinical response after DAA therapy for 12 wk or more\cite{145}. Altogether, patients with HCV-associated cryoglobulinemic vasculitis had higher SVR (74%-100%) and clinical remission (61%-100%) rates after receiving DAA medications\cite{146}. Higher complete response rates (75%-100%) were observed in cutaneous and musculoskeletal

Figure 1 Cryoglobulinemic vasculitis. The small vessels show neutrophilic inflammation, with fibrinoid necrosis and fragmented neutrophil nuclei (black arrows). Hematoxylin and eosin staining, 400 ×.
presentations, while lower responses (30%-70%) were identified in peripheral nerve and renal involvement[16]. In addition, DAA therapy is beneficial for HCV-associated B-cell malignancy, resulting in higher SVR and lymphoproliferative disease response rates in indolent NHL patients[17].

In addition to optimized DAA agents, corticosteroids in combination with cyclophosphamide are also considered as first-line therapy in severe fulminant manifestations, such as intestinal necrotizing vasculitis, rapid progressive glomerulonephritis, and diffuse alveolar hemorrhage[18,19]. Plasma exchange with warm apheresis solution to avoid cryoglobulin precipitation is an adjunct treatment that is useful in life-threatening disease by removing circulating cryoglobulins to interrupt the IC-mediated pathogenesis[20]. An earlier investigation showed that the addition of rituximab to the combined IFN-α and ribavirin regimen can enhance the clearance of cryoglobulins and shorten the time to clinical remission[21]. A subsequent randomized controlled trial in patients with severe disease revealed that rituximab monotherapy is as effective as conventional immunosuppressive treatment[22]. Despite the promise of rituximab as a therapeutic agent[23], there is a substantial risk for IC formation between this biologic and RF-positive IgM to exacerbate the vasculitis activity in HCV-associated type II cryoglobulinemic vasculitis[24].

HUMAN HEPATITIS VIRUSES-ASSOCIATED POLYARTERITIS NODOSA

Polyarteritis nodosa is a rare disease, with an annual incidence ranging from 0 to 2 cases per million population[25]. It is a necrotizing vasculitis affecting small- and medium-sized arteries, with systemic involvement but usually sparing the lungs. A skin-restricted form involving the area below the knees can progress to the systemic form, suggesting the same entity for both forms[26,27]. Most patients with polyarteritis nodosa belong to the idiopathic type with autoimmune-mediated mechanisms[28,29]. The secondary type is often observed in viral diseases, like cytomegalovirus, human immunodeficiency virus, and HHV infections. The association between HBV and polyarteritis nodosa was first recognized in 1970[30,31], and since then a substantial portion of cases have been identified after HBV infection. Later on, owing to the introduction of the vaccination protocol, the occurrence of HBV-related polyarteritis nodosa has gradually disappeared from clinical practice[32]. Although HCV positivity has been observed in this disorder, it is not a dominant etiological factor[33].

The occlusion and rupture of inflamed arteries can produce ischemia and hemorrhage in various organs and tissues, including skin, joint, kidney, testis, gastrointestinal tract, and peripheral nerve[34,35]. The most frequently involved organ system is the skin, presenting manifestations of palpable purpura, livedoid lesions, subcutaneous nodules, and necrotic ulcers[36]. *Figure 2* demonstrates the histopathological findings of transmural necrotizing arteritis with neutrophilic infiltration and fibrinoid necrosis in a cutaneous biopsy specimen from a patient with polyarteritis nodosa-associated nodules (Case No. 2 in Table 2). Mononeuritis multiplex and symmetrical polyneuropathy are common neurological complications[37]. Gastrointestinal manifestations are usually associated with significant morbidity and can manifest as an acute surgical abdomen[38]. Renal involvement without glomerulonephritis is related to infarction or hemorrhage caused by the rupture of renal microaneurysms[39]. Although rare as an initial presentation, orchitis due to testicular ischemia is a characteristic finding of polyarteritis nodosa[40].

As supported by clinical evidence, the idiopathic type benefits from combined corticosteroids and cyclophosphamide therapy, which induce remission in severe disease with organ dysfunction; whereas, mild manifestations can be treated with corticosteroids alone[41,42]. After the completion of cyclophosphamide therapy, azathioprine or methotrexate can be prescribed as a remission-maintenance agent[34]. Currently, small-molecule targeted drugs and biologics have been used in refractory patients naïve to HBV infection with successful outcomes, including Janus kinase inhibitor, B-cell targeted agent, tumor necrosis factor blocker, and interleukin-6 blocker[43-46]. The clinical data and medication profiles with biologics in polyarteritis nodosa patients recently diagnosed by authors are shown in Table 2. Although poor prognosis in earlier years is due to a delayed diagnosis[97], the overall outcome has improved to a 5-year survival rate of 80%. Outcomes are significantly worsened in association with HBV infection, age above 65 years, new-onset hypertension, renal impairment with high creatinine levels, gastrointestinal involvement requiring surgery, and peripheral nerve involvement[98].

There are no available reports related to the association of polyarteritis nodosa with...
Table 2 Clinical and medication profiles in five polyarteritis nodosa patients naïve to human hepatitis virus infection

| No | Age/sex | Initial symptom | Diagnosis period | BVAS/FFS | HHV status | Follow-up manifestation | Therapy | Outcome                  |
|----|---------|-----------------|------------------|----------|------------|------------------------|---------|--------------------------|
| 1  | 23/F    | Arthralgia, rash | 2 mo             | 6/0      | Negative   | Joint, PN, skin        | Az, Cs  | Survival, remission      |
| 2  | 29/F    | Arthralgia, fever, rash | 1 yr             | 11/0     | Negative   | FN, GI, joint, PN, skin | Az, Cs  | Survival, remission      |
| 3  | 38/F    | Fever, rash     | 1 yr 5 mo        | 14/0     | Negative   | Joint, PN, skin        | Az, Cs, Cy | Survival, remission     |
| 4  | 39/M    | Rash            | 3 yr 10 mo       | 18/1     | Negative   | Kidney, skin           | Az, Cs, RTX | Survival, chronic renal insufficiency |
| 5  | 42/M    | Arthralgia, rash | 6 mo             | 7/0      | Negative   | Joint, PN, skin, kidney, testis | ADA, Az, Cs, Cy | Survival, remission |

1Enrollment from 2012 to 2019.
2Time period from initial symptoms to the established diagnosis.
3Calculation at the disease diagnosis.
4Human hepatitis virus status, including examinations for hepatitis A virus, hepatitis B virus, hepatitis C virus, and hepatitis delta virus.
5Erythematous nodosum-like skin lesions. ADA: Adalimumab; Az: Azathioprine; BVAS: Birmingham vasculitis activity score; Cs: Corticosteroids, Cy: Cyclophosphamide; F: Female; FFS: Four-factor score, including age above 65 years, cardiac symptoms, gastrointestinal involvement, and renal insufficiency; FN: Foot necrosis; GI: Gastrointestinal; HHV: Human hepatitis virus; M: Male; PAN: Polyarteritis nodosa; PN: Peripheral neuropathy; RTX: Rituximab.

Hepatitis A virus (HAV). Nevertheless, published cases with cutaneous or renal manifestations have shown histopathological evidence of medium-sized blood vessels’ involvement compatible with the diagnosis of polyarteritis nodosa [162,163].

Simultaneous development of mixed cryoglobulinemia and polyarteritis nodosa has been documented in patients with coinfection of HCV and HBV [164,165]. In a clinical survey on polyarteritis nodosa, the positive frequency of anti-HCV was 20% (5% with detectable HCV ribonucleic acid and they were more likely to have cutaneous manifestation) [166]. In a large cohort with 161 cases of HCV-related vasculitis (19% polyarteritis nodosa and 81% cryoglobulinemic vasculitis), there were more acute and severe clinical presentations in the former group, including constitutional symptoms, new-onset hypertension, gastrointestinal tract involvement, and mononeuritis multiplex [167]. Despite no differences in the 5-year survival rates, there was a higher complete remission rate for polyarteritis nodosa than for cryoglobulinemic vasculitis. The therapeutic guidelines for HCV-related polyarteritis nodosa have not been established yet. Reported cases have received various combinations of corticosteroids, cyclophosphamide, and antiviral agents. B-cell targeted therapy has been applied to patients with HCV-related polyarteritis nodosa [167,168]. A higher clinical relapse rate was observed for rituximab treatment than for a combined regimen with IFN-α and ribavirin in this disease [167].

During the 1970s to 1980s, HBV was a major cause of polyarteritis nodosa, with...
nearly half of cases having this infection, but the frequency has decreased due to improved blood safety and a viral vaccine campaign since early the 1990s, indicating HBV as a causal etiology\(^{[169]}\). Although histopathological studies have rarely confirmed the presence of HBV antigens in the vessel wall\(^{[170]}\), the pathogenesis of HBV-related type is related to the deposits of IC, different from the idiopathic disease\(^{[169]}\). When comparing HBV- with HCV-associated polyarteritis nodosa\(^{[157,158]}\), the former group has more general symptoms and orchitis, less cutaneous manifestation and new-onset hypertension, a lower survival rate, and a much shorter average period from viral infection to vasculitis development, 7 mo vs 20 years. In the HBV-related type, most cases have higher viral replication and HBV DNA levels, leading to the persistent presence of circulating IC\(^{[169]}\).

The prevalence of HBs antigen in the general Taiwan population used to be near 20%, but the HBV seropositive rate in children has deceased dramatically from 11% to less than 1% after the initiation of the national vaccination program in 1984\(^{[171]}\). Despite a high inpatient prevalence with more than 10 cases per 100000 discharges in the United States\(^{[172]}\), polyarteritis nodosa has been very rarely encountered by practicing physicians in Taiwan. In particular, even with more than 2 million infected patients, there is no reported association between HBV and polyarteritis nodosa\(^{[173]}\). Perinatal mother-to-infant HBV transmission is the most important factor responsible for a high carriage rate of HBs antigen in Taiwan\(^{[174]}\). Vertical transmission in infants is asymptomatic until adulthood and associated with a greater risk of chronic infection, suggesting that HBV infection causes an immunotolerant status in children who are less prone to develop full immune responses\(^{[175]}\). Parenteral HBV infection during adult life can induce polyarteritis nodosa within 1 year after viral infection, indicating an early post-infectious disease via this transmission route\(^{[169]}\). In Table 3, polyarteritis nodosa from various larger-number case series with different HBV-associated frequencies is compared\(^{[148,153,176-184]}\). Patients reported from Taiwan have no HBV infection, younger female predominance, and more cutaneous or testicular manifestation than those from other series with a HBV association.

After initial therapy with corticosteroids to reduce acute vascular inflammation, the therapeutic approach in HBV-associated polyarteritis nodosa is to clear circulating IC and suppress HBV replication by plasma exchange and antiviral agents with NAs or IFN-α\(^{[186]}\). Since the addition of cyclophosphamide to corticosteroids has shown to benefit patients presenting with poor prognostic factors\(^{[187]}\), this regimen can be considered in patients with severe disease. Notably, plasma exchange has not been shown to be beneficial in polyarteritis nodosa patients without HBV infection\(^{[156]}\). Furthermore, antiviral agents can be used concurrently with corticosteroids or combined cyclophosphamide and corticosteroids therapy\(^{[197,199]}\). Contradictory to the idiopathic disease, relapses have rarely been observed in HBV-related type, especially when viral replication has ceased and seroconversion from HBe antigen to antibody has been achieved after antiviral therapy\(^{[146,197]}\).

**CONCLUSION**

Although HHVs primarily affect hepatocytes, they can also cause complications in other organs, and both acute and chronic viral hepatitis is associated with clinical presentations outside the liver. Vascular involvement with cutaneous and systemic vasculitis is a well-known extrahepatic morbidity. There is growing evidence suggesting a causal relationship between viral pathogens and vasculitis. Except for HDV, other HHVs have participated in the etiopathogenesis of cutaneous and systemic vasculitis via different mechanisms, including direct viral invasion of vascular endothelial cells, IC-mediated vessel wall damage, and autoimmune responses with stimulation of autoreactive B-cells and impaired regulatory T cells. Cryoglobulinemic vasculitis and polyarteritis nodosa are recognized for their association with chronic HHV infection. Therapeutic guidelines for HHV-associated vasculitis have not been established yet. Antiviral therapy should be initiated in HBV and HCV-related systemic vasculitis in addition to the use of corticosteroids. Plasma exchange and/or combined cyclophosphamide and corticosteroid therapy can be considered in patients with severe life-threatening vasculitis manifestations.
Table 3 Comparisons of clinical and outcome profiles in polyarteritis nodosa from various case series with different hepatitis B virus-infected frequencies

| Source of cases   | No/age F% | HBV status | Fever | AM | Skin | PN  | GI  | Renal | TestisM% | Therapy | Death |
|-------------------|-----------|------------|-------|----|------|-----|-----|-------|----------|---------|-------|
| Taiwan            | 12/31, 67%| 0%         | 67%   | 58%| 75%  | 42% | 50% | 75%   | 50%      | Az, Cs, Cy, Bio | 33%    |
| United Kingdom    | 17/49, 24%| 31%        | 76%   | 77%| 65%  | 59% | 65% | 77%   | NA       | Cs, Is  | 38%   |
| United States     | 53/54, 34%| 11%        | 31%   | 55%| 58%  | 60% | 25% | 66%   | NA       | Cs, Cy  | 42%   |
| Canada            | 45/54, 47%| 19%        | 63%   | 51%| 44%  | 51% | 53% | 44%   | 4%       | Cs, Cy  | 53%   |
| South Korea       | 27/47, 37%| 56%        | 52%   | 30%| 44%  | 63% | 48% | 48%   | 24%      | Az, Cs, Cy, Is | 15%    |
| France            | 348/51,37%| 35%        | 64%   | 59%| 50%  | 74% | 38% | 51%   | 17%      | Av, Cs, Cy, Is | 25%    |
| India             | 27/38, 26%| 26%        | 52%   | 37%| 37%  | 82% | 30% | 59%   | 30%      | Av, Az, Cs, Cy | 11%    |

1Mean age at disease diagnosis. AM: Articulomuscular; Az: Azathioprine; Av: Antiviral agent; Bio: Biologics; Cs: Corticosteroids; Cy: Cyclophosphamide; F: Female; GI: Gastrointestinal; Is: Immunosuppressive agent; M: Male; NA: Not available; PN: Peripheral neuropathy.

ACKNOWLEDGEMENTS

The authors are indebted to Dr. Wu IC, Division of Gastroenterology and Hepatology, for his valuable comments on HHV-related clinical manifestations, and to other physicians at the National Cheng Kung University Hospital involved in the diagnosis and management of reported patients. The Institutional Review Board of National Cheng Kung University Hospital approved this study (No. B-ER-105-108).

REFERENCES

1 Arroyo V, Moreau R, Jalan R. Acute-on-Chronic Liver Failure. N Engl J Med 2020; 382: 2137-2145 [PMID: 32459224 DOI: 10.1056/NEJMrna1914900]
2 Razavi H. Global Epidemiology of Viral Hepatitis. Gastroenterol Clin North Am 2020; 49: 179-189 [PMID: 32389357 DOI: 10.1016/j.gct.2020.01.001]
3 Noor A, Panwala A, Forouhar F, Wu GY. Hepatitis caused by herpes viruses: A review. J Dig Dis 2018; 19: 446-455 [PMID: 29923691 DOI: 10.1111/1751-2980.12640]
4 Rasche A, Sander AL, Corman VM, Drexler JF. Evolutionary biology of human hepatitis viruses. J Hepatol 2019; 70: 501-520 [PMID: 30472320 DOI: 10.1016/j.jhep.2018.11.010]
5 Khuroo MS, Sofi AA. The Discovery of Hepatitis Viruses: Agents and Disease. J Clin Exp Hepatol 2020; 10: 391-401 [PMID: 32655240 DOI: 10.1016/j.jceh.2020.04.006]
6 Krugman S, Giles JP, Hammond J. Infectious hepatitis. Evidence for two distinctive clinical, epidemiological, and immunological types of infection. JAMA 1967; 200: 365-373 [PMID: 4164595 DOI: 10.1001/jama.200.5.365]
7 BLUMBERG BS, ALTER HJ, VISNICH S. A "NEW" ANTIGEN IN LEUKEMIA SERA. JAMA 1965; 191: 541-546 [PMID: 14239025 DOI: 10.1001/jama.1965.0308007025007]
8 Dane DS, Cameron CH, Briggs M. Virus-like particles in serum of patients with Australia-antigen-associated hepatitis. Lancet 1970; 1: 695-698 [PMID: 4190997 DOI: 10.1016/s0140-6736(70)90296-6]
9 Burrell CJ, Mackay P, Greensway PJ, Hofschneider PH, Murray K. Expression in Escherichia coli of hepatitis B virus DNA sequences cloned in plasmid pBR322. Nature 1979; 279: 43-47 [PMID: 377993 DOI: 10.1038/279043a0]
10 Koff RS, Feinstone SM, Kapikian AZ, Purcell RH. Hepatitis A: detection by immune electron microscopy of a virus like antigen associated with acute illness [Science 1973;182:1026-1028]. J Hepatol 2002; 37: 2-6 [PMID: 12076855 DOI: 10.1016/s0168-8278(02)00169-1]
11 Ticehurst JR, Racaniello VR, Baroudy BM, Baltimore D, Purcell RH, Feinstone SM. Molecular cloning and characterization of hepatitis A virus cDNA. Proc Nail Acad Sci USA 1983; 80: 5885-5889 [PMID: 6310601 DOI: 10.1073/pnas.80.19.5885]
12 Alter HJ, Holland PV, Morrow AG, Purcell RH, Feinstone SM, Moritsugu Y. Clinical and serological analysis of transfusion-associated hepatitis. Lancet 1975; 2: 838-841 [PMID: 53329 DOI: 10.1016/s0140-6736(75)90234-2]
13 Alter HJ, Purcell RH, Holland PV, Popper H. Transmissible agent in non-A, non-B hepatitis. Lancet 1978; 1: 459-463 [PMID: 76017 DOI: 10.1016/s0140-6736(78)90131-9]
Myocarditis—A Case Report with Literature Review.

Allen O, Pediatr immunoglobulin M and complement associated with hepatitis A virus infection.

Dan M, with persisting cholestatic hepatitis A virus infection.

Ilan Y, cryoglobulinemia associated with relapsing hepatitis A virus infection.

Spring Harb Perspect Med

Shin EC

1089-1099 [PMID: DOI: 10.1093/cshperspect/a033431]

Abutaleb A, from the virus responsible for enterically transmitted non-A, non-B hepatitis.

Reyes GR, virus distinct from post-transfusion non-A, non-B type.

Khuroo MS, derived from a blood-borne non-A, non-B viral hepatitis genome.

Choo QL, Hepatitis B virus infection.

Feng Z, Naked Viruses That Aren’t Always Naked: Quasi-Enveloped Agents of Acute Hepatitis.

Taylor JM

Kaplan DE, Extrahepatic manifestations of chronic hepatitis B.

Reddy KR. Extrahepatic manifestations of chronic viral hepatitis.

World Health Organization

Immunother Dis

DOI:

10.1016/j.vaccine.2020.01.002

Aslan AT, Hepatitis B virus: Epidemiology, diagnosis, clinical manifestations, and treatment. World J Gastroenterol 2020; 26: 5543-5560 [PMID: DOI: 10.3748/wjg.v26.i37.5543]

Yuen MF, Chen DS, Dusheiko GM, Janssen HLA, Lau DTY, Locarnini SA, Peters MG, Lai CL. Hepatitis B virus infection. Nat Rev Dis Primers 2018; 4: 18035 [PMID: DOI: 10.1038/s41573-018.35]

Kaplan DE. Hepatitis C Virus. Ann Intern Med 2020; 173: ITC33-ITC48 [PMID: DOI: 10.7326/AITC202009010]

Gilmann C, Chronic hepatitis delta: A state-of-the-art review and new therapies. World J Gastroenterol 2019; 25: 4580-4597 [PMID: DOI: 10.3748/wjg.v25.i32.4580]

Taylor JM. Infection by Hepatitis Delta Virus. Viruses 2020; 12 [PMID: DOI: 10.3390/v12060648]

Feng Z, Hirai-Yuki A, McKnight KL, Lemon SM. Naked Viruses That Aren’t Always Naked: Quasi-Enveloped Agents of Acute Hepatitis. Annu Rev Virol 2014; 1: 539-560 [PMID: DOI: 10.1146/annurev-virology-031413-085359]

Cheung A, Kwo P. Viral Hepatitis Other than A, B, and C: Evaluation and Management. Clin Liver Dis 2020; 24: 405-419 [PMID: DOI: 10.1016/j.cld.2020.04.008]

Cao Y, Bing Z, Guan S, Zhang Z, Wang X. Development of new hepatitis E vaccines. Hum Vaccin Immunother 2018; 14: 2254-2262 [PMID: DOI: 10.1080/21645515.2018.1469591]

World Health Organization. Hepatitis 2020. https://www.who.int/health-topics/hepatitis?tab=1

Schiff ER. Atypical clinical manifestations of hepatitis A. Vaccine 1992; 10 Suppl 1: S18-S20 [PMID: DOI: 1475999]

Pyrsopoulos NT, Reddy KR. Extrahepatic manifestations of chronic viral hepatitis. Curr Gastroenterol Rep 2001; 3: 71-78 [PMID: DOI: 10.11073/a11894-001-0044-1]

Amarapurkar DN, Amarapurkar AD. Extrahepatic manifestations of viral hepatitis. Ann Hepatol 2002; 1: 192-195 [PMID: 15280806]

Han SH. Extrahepatic manifestations of chronic hepatitis B. Clin Liver Dis 2004; 8: 403-418 [PMID: DOI: 15481347]

Romano C, Cuomo G, Ferrara R, Del Mastro A, Esposito S, Sellitto A, Adinolfi LE. Uncommon immune-mediated extrahepatic manifestations of HCV infection. Expert Rev Clin Immunol 2018; 14: 1089-1099 [PMID: DOI: 30338718]

Shin EC, Jeong SH. Natural History, Clinical Manifestations, and Pathogenesis of Hepatitis A. Cold Spring Harb Perspect Med 2018; 8 [PMID: DOI: 10.1101/cshperspect.a031708]

Inman RD, Hodge M, Johnston ME, Wright J, Heathcote J. Arthritis, vasculitis, and cryoglobulinemia associated with relapsing hepatitis A virus infection. Ann Intern Med 1986; 105: 706-703 [PMID: DOI: 10.1073/pnas.77.10.6124]

Halan Y, Hillman M, Oren R, Zlotogorski A, Shouval D. Vasculitis and cryoglobulinemia associated with persisting cholestatic hepatitis A virus infection. Am J Gastroenterol 1990; 85: 586-587 [PMID: DOI: 2337062]

Dan M, Yaniv R. Cholestatic hepatitis, cutaneous vasculitis, and vascular deposits of immunoglobulin M and complement associated with hepatitis A virus infection. Am J Med 1990; 89: 103-104 [PMID: DOI: 2368780]

Shenoy R, Nair S, Kamath N. Thrombocytopenia in hepatitis A—an atypical presentation. J Trop Pediatr 2004; 50: 241-242 [PMID: DOI: 15357567]

Allen O, Edhi A, Hafeez A, Halalau A. A Very Rare Complication of Hepatitis A Infection: Acute Myocarditis-A Case Report with Literature Review. Case Rep Med 2018; 2018: 3625139 [PMID: DOI: 10.1155/2018/3625139]
Wang CR et al. HHV-associated cutaneous and systemic vasculitis

30302093 DOI: 10.1155/2018/3625139

Stübgen JP. Neuromuscular complications of hepatitis A virus infection and vaccines. J Neurol Sci 2011; 300: 2-8 [PMID: 20920814 DOI: 10.1016/j.jns.2010.09.015]

Dalton HR, Kang S, Izopet J, Kamar N, Iozot P, Peron JM. Neurologic Disorders in Immunocompetent Patients with Autochthonous Acute Hepatitis E. Emerg Infect Dis 2015; 21: 1928-1934 [PMID: 26490255 DOI: 10.3201/eid2111.141789]

Dalton HR, van Eijk JJ, Cintas P, Madden RG, Jones C, Webb GW, Norton B, Pique J, Lutgens S, Devooght-Johnson N, Woolson KL, Baker J, Saunders M, Househam L, Griffiths J, Abravanel F, Iozot J, Kang S, Izopet J, Kamar N, Iozot P, Peron JM. Neurologic Disorders in Immunocompetent Patients with Autochthonous Acute Hepatitis E. Emerg Infect Dis 2015; 21: 1928-1934 [PMID: 26490255 DOI: 10.3201/eid2111.141789]

Despieres LA, Kaplan E, Attarian S, Cohen-Bacrie S, Pelletier J, Guedj J, Motte A, Charrel R, Gerolami R, Colson P. Neurologic disorders and hepatitis E, France, 2010. Emerg Infect Dis 2011; 17: 1510-1512 [PMID: 21801637 DOI: 10.3201/eid708.100208]

Marion O, Abravanel F, Del Bello A, Esposito L, Lhomme S, Puissant-Lubrano B, Alric L, Faguer S, Iozot J, Kang S, Izopet J, Kamar N. Hepatitis E virus-associated cryoglobulinemia in solid-organ-transplant recipients. Liver Int 2018; 38: 2178-2189 [PMID: 29845733 DOI: 10.1111/liv.13894]

Woolson KL, Forbes A, Vine L, Beynon L, McElhinney L, Panayi V, Hunter JG, Madden RG, Glasgow T, Kotecha A, Dalton HC, Mihailescu L, Warshaw U, Hussaini HS, Palmer J, Mclean BN, Haywood B, Bendall RP, Dalton HR. Extra-hepatic manifestations of autochthonous hepatitis E infection. Aliment Pharmacol Ther 2014; 40: 1282-1291 [PMID: 25303615 DOI: 10.1111/apt.12986]

Nayak HK, Kamble NL, Raizada N, Garg S, Daga MK. Acute pancreatitis complicating acute hepatitis E virus infection: a case report and review. Case Reports Hepatol 2013; 2013: 531235 [PMID: 25374721 DOI: 10.1155/2013/531235]

Raj M, Kumar K, Ghoshal UC, Saraswat VA, Aggarwal R, Mohindra S. Acute Hepatitis E-Associated Acute Pancreatitis: A Single Center Experience and Literature Review. Pancreas 2015; 44: 1320-1322 [PMID: 26390412 DOI: 10.1097/MPA.0000000000000402]

Sengupta P, Biswas S, Roy T. Hepatitis E-Induced Acute Myocarditis in an Elderly Woman. Case Rep Gastroenterol 2019; 13: 342-349 [PMID: 31572104 DOI: 10.1159/000051998]

Del Bello A, Guibeva-Frueger C, Josse AG, Rostaing L, Iozot J, Kang S, Iozot P, Peron JM. Successful treatment of hepatitis E virus-associated cryoglobulinemic membranoproliferative glomerulonephritis with ribavirin. Transpl Infect Dis 2015; 17: 279-283 [PMID: 25708383 DOI: 10.1111/tid.12353]

van Eijk JJ, van Alfen N, Berrevoets M, van der Wilt GJ, Pillen S, van Engelen BG. Evaluation of prednisolone treatment in the acute phase of neuralgic amyotrophy: an observational study. J Neurol Neurosurg Psychiatry 2009; 80: 1102-1104 [PMID: 19321467 DOI: 10.1136/jnnp.2008.163386]

Osman C, Jennings R, El-Ghariani K, Pinto A. Plasma exchange in neurological disease. Pract Neurol 2020; 20: 92-99 [PMID: 31300488 DOI: 10.1136/practneurol-2019-002336]

Trepo C, Guéhin L. Polyarteritis nodosa and extrahepatic manifestations of HBV infection: the case against autoimmune intervention in pathogenesis. J Autoimmun 2001; 16: 269-274 [PMID: 11334492 DOI: 10.1006/jaut.2000.0502]

McMahon BJ, Allward WL, Hall DB, Heyward WL, Bender TR, Francis DP, Maynard JE. Acute hepatitis B virus infection: relation of age to the clinical expression of disease and subsequent development of the carrier state. J Infect Dis 1985; 151: 599-603 [PMID: 3973412 DOI: 10.1093/infdis/151.4.599]

Kappus MR, Sterling RK. Extrahepatic manifestations of acute hepatitis B virus infection. Gastroenterol Hepatol (N Y) 2013; 9: 123-126 [PMID: 23983659]

Yimam KK, Merriman RB, Todd Frederick R. A rare case of acute hepatitis B virus infection causing guillain-barré syndrome. Gastroenterol Hepatol (N Y) 2013; 9: 121-123 [PMID: 23983658]

Cacoub P, Terrier B. Hepatitis B-related autoimmune manifestations. Rheum Dis Clin North Am 2009; 35: 125-137 [PMID: 19481001 DOI: 10.1016/j.rdc.2009.03.006]

Cacoub P, Saadoun D, Bourlière M, Khiri H, Martineau A, Benshamou Y, Varasteh M, Pol S, Thibault V, Rotily M, Halfon P. Hepatitis B virus genotypes and extrahepatic manifestations. J Hepatol 2005; 43: 764-770 [PMID: 16087273 DOI: 10.1016/j.jhep.2005.05.029]

Mason A, Theal J, Bain V, Adams E, Perrillo R. Hepatitis B virus replication in damaged
endothelial tissues of patients with extrahepatic disease. *Am J Gastroenterol* 2005; **100**: 972-976 [PMID: 15784404 DOI: 10.1111/j.1572-0241.2005.41308.x]

64 **Levo Y**, Gorevic PD, Kassab HJ, Zucker-Franklin D, Franklin EC. Association between hepatitis B virus and essential mixed cryoglobulinemia. *N Engl J Med* 1977; **296**: 1501-1504 [PMID: 865530 DOI: 10.1056/NEJM197706302962605]

65 **Stübbgen JP**. Neuromuscular disorders associated with hepatitis B virus infection. *J Clin Neuromuscul Dis* 2011; **13**: 26-37 [PMID: 22361623 DOI: 10.1097/NIN.0b013e3181df2b2b]

66 **Guillévin L**, Lhote F, Sauvaget F, Debois P, Rossi F, Levallois D, Pourrat J, Christoforov B, Trépo C. Treatment of polyclonarity nodosa related to hepatitis B virus with interferon-alpha and plasma exchanges. *Ann Rheum Dis* 1994; **53**: 333-334 [PMID: 7912504 DOI: 10.1136/ard.53.3.334]

67 **Guillévin L**, Mahr A, Cohen P, Larroche C, Quereyl V, Loustaud-Ratti V, Imbert B, Hauspater P, Roudier J, Bielefeld P, Petitjean P, Smadja D, French Vasculitis Study Group. Short-term corticosteroids then lamivudine and plasma exchanges to treat hepatitis B virus-related polyarteritis nodosa. *Arthritis Rheum* 2004; **51**: 482-487 [PMID: 15188337 DOI: 10.1002/art.20401]

68 **Tang S**, Lai FM, Lui YH, Tang CS, Kung NN, Ho YW, Chan KW, Leung JC, Lai KN. Lamivudine in hepatitis B-associated membranous nephropathy. *Kidney Int* 2005; **68**: 1750-1758 [PMID: 16164651 DOI: 10.1111.j.1525-1735.2005.00591.x]

69 **Chung DR**, Yang WS, Kim SB, Yu E, Chung YH, Lee Y, Park JS. Treatment of hepatitis B virus associated glomerulonephritis with recombinant human alpha interferon. *Am J Nephrol* 1997; **17**: 112-117 [PMID: 9096440 DOI: 10.1159/000169083]

70 **Mazzaro C**, Dal Maso L, Visentini M, Gatto S, Andreone P, Toffolatti F, Gatti V. Hepatitis B virus-related cryoglobulinemic vasculitis. The role of antiviral nucleo(s)ide analogues: a review. *J Intern Med* 2019; **286**: 290-298 [PMID: 31124596 DOI: 10.1111/joi.12913]

71 **Martinello M**, Hajarizadeh B, Grebely J, Dore GJ, Matthews GV. Management of acute HCV infection in the era of direct-acting antiviral therapy. *Nat Rev Gastroenterol Hepatol* 2018; **15**: 412-424 [PMID: 29773899 DOI: 10.1038/s41575-018-0026-5]

72 **Loomba R**, Rivera MM, McBurney R, Park Y, Haynes-Williams V, Rehermann B, Alter HJ, Herrine SK, Liang TJ, Hoofnagle JH, Heller T. The natural history of acute hepatitis C: analysis of a 126-case prospective, multicenter cohort. *Hepatol* 2019; **70**: 80-92 [PMID: 31124596 DOI: 10.1002/hep.30984]

73 **Morin T**, Pariente A, Lahmek P, Rabaud C, Silvain C, Cadranel JF; Association Nationale des Hépato-gastroentérologues des Hôpitaux généraux (ANGH) Société de Pathologie Infectieuse de France. Neuromuscular disorders associated with hepatitis B virus infection. *Neuromuscul Disord* 2014; **24**: 73-85 [PMID: 24237457 DOI: 10.1016/j.nmd.2013.12.011]

74 **Spearman CW**, Dusheiko GM, Hellard M, Sonderup M. Hepatitis C. *Lancet* 2019; **394**: 1451-1466 [PMID: 31638157 DOI: 10.1016/S0140-6736(19)32320-7]

75 **Cacoub P**, Poynard T, Ghillani P, Charlotte F, Olivi M, Piette JC, Opolon P. Extrahepatic manifestations of chronic hepatitis C. MULTIVIRC Group. Multidepartment Virus C. *Arthritis Rheum* 1999; **42**: 2204-2212 [PMID: 10524695 DOI: 10.1002/1529-0131(199910)42:10<2204::AID-ART23>3.0.CO;2-D]

76 **Kuna L**, Jakub J, Smolc R, Wu GY, Smolic M. HCV Extrahepatic Manifestations. *J Clin Transl Hepatol* 2019; **7**: 172-182 [PMID: 31293918 DOI: 10.14281/JCHT.2018.00049]

77 **Zegans ME**, Aminger W, Chapman C, Gordon SR. Ocular manifestations of hepatitis C virus infection. *Curr Opin Ophthalmol* 2002; **13**: 423-427 [PMID: 12441848 DOI: 10.1097/00055755-200206000-00014]

78 **Petra S**, Craxi A. Extrahepatic Manifestations of Chronic Viral C Hepatitis. *Gastroenterol Clin North Am* 2020; **50**: 347-360 [PMID: 32389367 DOI: 10.1016/j.gtc.2020.01.012]

79 **Comarmond C**, Cacoub P, Saadoun D. Treatment of chronic hepatitis C-associated cryoglobulinemia vasculitis at the era of direct-acting antivirals. *Therap Adv Gastroenterol* 2020; **13**: 175628420942617 [PMID: 32782479 DOI: 10.1177/1756284209426177]

80 **Carrat F**, Fontaine H, Dorival C, Simony M, Diallo A, Hezode C, De Ledinghen V, Larrey D, Haour G, Bronniovick JP, Zoulim F, Asselah T, Marcellin P, Thabut D, Leroy V, Tran A, Habersetzer F, Samuel D, Guyader D, Chazouilleres O, Mathurin P, Metivier S, Alric L, Riachi G, Gournay J, Abergel A, Cales P, Ganne N, Loustaud-Ratti V, D'Alteroche L, Causse X, Geist C, Minello A, Ross I, Guel-Simeon M, Portal I, Raffi F, Bourliere M, Pol S; French ANRS CO22 Hepatery cohort. Clinical outcomes in patients with chronic hepatitis C after direct-acting antiviral treatment: a prospective cohort study. *Lancet* 2019; **393**: 1453-1464 [PMID: 30765123 DOI: 10.1016/S0140-6736(18)32111-1]

81 **Thomas K**, Vassiliopoulos D. Infections and vasculitis. *Curr Opin Rheumatol* 2017; **29**: 17-23 [PMID: 27662570 DOI: 10.1097/BOR.0000000000000348]

82 **Lidar M**, Lipschitz N, Langlevitz P, Shoenfeld Y. The infectious etiology of vasculitis. *Autoimmun rev* 2009; **42**: 432-438 [PMID: 19811260 DOI: 10.1016/j.autrev.2009.08.002]

83 **Vergani D**, Dieli-Vergani G. Autoimmune manifestations in viral hepatitis. *Semim Immunopathol* 2013; **35**: 73-85 [PMID: 23010889 DOI: 10.1007/s12299-012-0328-6]

84 **Cozzani E**, Herzum A, Burlando M, Parodi A. Cutaneous manifestations of HAV, HBV, HCV. *G Ital Dermatol Venereol* 2019 [PMID: 31804053 DOI: 10.23736/S0392-0488.19.06488-5]

85 **Grigorescu I**, Dumitrascu DL. Spontaneous and antiviral-induced cutaneous lesions in chronic hepatitis B virus infection. *World J Gastroenterol* 2014; **20**: 15860-15866 [PMID: 25400473 DOI: 10.1097/CND.0b013e318293c473]
Rustichelli R, Di Belgioioso GB, Comotti C, Quarenghi MI. Multicenter study on hepatitis C virus-related cryoglobulinemic glomerulonephritis. *Am J Kidney Dis* 2007; 49: 69-82 [PMID: 17185147 DOI: 10.1053/j.jkd.2006.09.015]

110 Johnson RJ, Gretchen DR, Yamabe H, Hart J, Facchi CE, Hartwell P, Couser WG, Corey L, Wener MH, Alpers CE. Membranoproliferative glomerulonephritis associated with hepatitis virus infection. *N Engl J Med* 1993; 328: 465-470 [PMID: 7678440 DOI: 10.1056/NEJM199302183280701]

111 Ammendola A, Sampola S, Ambrosone L, Ammendola E, Ciccone G, Migliaresi S, Di Iorio G. Peripheral neuropathy in hepatitis-related mixed cryoglobulinemia: electrophysiologic follow-up study. *Muscle Nerve* 2005; 31: 382-385 [PMID: 15515001 DOI: 10.1002/mus.20184]

112 Casato M, Saadoun D, Marchetti A, Limal N, Piecq C, Pantano P, Galanaud D, Cianci R, Duhauphet P, Piette JC, Fiorilli M, Cacoub P. Central nervous system involvement in hepatitis C virus cryoglobulinemia vasculitis: a multicenter case-control study using magnetic resonance imaging and neuropsychological tests. *J Rheumatol* 2005; 32: 484-488 [PMID: 15742440]

113 Retamoza S, Díaz-Lagares C, Bosch X, Boyé A, Brito-Zerón P, Gómez ME, Yagüe J, Forns X, CID, Ramos-Casals M. Life-Threatening Cryoglobulinemic Vasculitis With Hepatitis C: Clinical Description and Outcome of 279 Patients. *Medicine (Baltimore)* 2013; 92: 273-284 [PMID: 23974248 DOI: 10.1097/MD.0b013e3182e1f771]

114 Cavalli G, Berti A, Fragasso G, De Cobelli F. Hypertrophic cardiomyopathy secondary to hepatitis C virus-related vasculitis. *J Cardiovasc Med (Hagerstown)* 2016; 17 Suppl 2: e156-e157 [PMID: 24979124 DOI: 10.2459/JCM.0000000000000109]

115 Terrier B, Saadoun D, Sène D, Scerra S, Musset L, Cacoub P. Presentation and outcome of gastrointestinal involvement in hepatitis C virus-related systemic vasculitis: a case-control study from a single-centre cohort of 163 patients. *Gut* 2010; 59: 1709-1715 [PMID: 20841367 DOI: 10.1136/gut.2010.218123]

116 Muñoz-Martínez SG, Díaz-Hernández HA, Suárez-Florez D, Sánchez-Ávila JF, Gamboa-Dominguez A, García-Juárez I, Torre A. Atypical manifestations of hepatitis A virus infection. *Rev Gastroenterol Mex* 2018; 83: 134-143 [PMID: 29685743 DOI: 10.1016/j.rgmx.2017.10.004]

117 Nassih H, Bourrahouat A, Sab IA. Hepatitis A Virus Infection Associated with Cryoglobulinemic Vasculitis. *Indian Pediatr* 2020; 57: 71-72 [PMID: 31937705]

118 Levo Y, Gorevic PD, Kassab HJ, Tobias H, Franklin EC. Liver involvement in the syndrome of mixed cryoglobulinemia. *Ann Intern Med* 1977; 87: 287-292 [PMID: 900672 DOI: 10.7326/0003-4819-87-3-287]

119 Gower RG, Sausker WF, Kohler PF, Thorne GE, McIntosh RM. Small vessel vasculitis caused by hepatitis B virus immune complexes. Small vessel vasculitis and HBsAG. *J Allergy Clin Immunol* 1978; 62: 222-228 [PMID: 701656 DOI: 10.1016/0091-6749(78)90211-7]

120 Monti G, Galli M, Invernizzi F, Pioletti P, Saccardo A, Monteverde A, Pietrogrande M, Renoldi P, Bombardiieri S, Bordin G. Cryoglobulinaemias: a multi-centre study of the early clinical and laboratory manifestations of primary and secondary disease. *GISC. Italian Group for the Study of Cryoglobulinaemias.* QJM 1995; 88: 115-126 [PMID: 7704562]

121 Mazarro C, Dal Maso L, Urraro T, Mauro E, Castelvovo L, Casarini P, Monti G, Gattai V, Zignego AL, Pozzato G. Hepatitis B virus related cryoglobulinemic vasculitis: A multicentre open label study from the Gruppo Italiano di Studio delle Criglobulinemia - GISC. *Dig Liver Dis* 2016; 48: 780-784 [PMID: 27106525 DOI: 10.1016/j.dld.2016.03.018]

122 Liu YT, Huang JL, Ou LS, Lin YH, Yu KH, Luo SF, Ho HH, Liu LB, Yeh KW. Comparison of cryoglobulinemia in children and adults. *J Microbiol Immunol Infect* 2013; 46: 59-64 [PMID: 22237397 DOI: 10.1016/j.jmi.2011.12.027]

123 Mazarro C, Dal Maso L, Visintini M, Ermacora A, Tonizzo M, Gattai V, Andreone P. Recent news in the treatment of hepatitis B virus-related cryoglobulinemic vasculitis. *Minerva Med* 2020; 111: 566-572 [PMID: 32573522 DOI: 10.23736/S0026-4806.20.06771-3]

124 European Association for the Study of the Liver, . European Association for the Study of the Liver. EASL. 2017 Clinical Practice Guidelines on the management of hepatitis B virus infection. *J Hepatol* 2017; 67: 390-398 [PMID: 28427875 DOI: 10.1016/j.jhep.2017.03.021]

125 Kamimura H, Setsu T, Kimura N, Yokoo T, Sakamaki A, Kamimura K, Tsuichiya A, Takamura M, Yamagiwa S, Terai S. Renal Impairment in Chronic Hepatitis B: A Review. *Diseases* 2018; 6 [PMID: 29921773 DOI: 10.3390/diseases6020052]

126 Ferri C, Antonelli A, Mascia MT, Sebastiani M, Fallahi P, Ferrari D, Giunti M, Pileri SA, Zignego AL. B-cells and mixed cryoglobulinemia. *Autoimmun Rev* 2007; 7: 114-120 [PMID: 18035328 DOI: 10.1016/j.autrev.2007.02.019]

127 Pasquet F, Combarnous F, Macgregor B, Coppere B, Mausserey C, Ninet J, Hot A. Safety and efficacy of rituximab treatment for vasculitis in hepatitis B virus-associated type II cryoglobulinemia: a case report. *J Med Case Rep* 2012; 6: 39 [PMID: 22284897 DOI: 10.1186/1757-1628-6-39]

128 Terrier B, Marie I, Lacraz A, Belenotti P, Bonnet F, Chiche L, Graffin B, Hot A, Kahn JE, Michel C, Queneteur T, de Saint-Martin L, Hermine O, Léger JM, Mariette X, Senet P, Plaisier E, Cacoub P. Non HCV-related infectious cryoglobulinemia vasculitis: Results from the French nationwide CryoVas survey and systematic review of the literature. *J Autoimmun* 2015; 65: 74-81 [PMID: 26320984 DOI: 10.1016/j.jaut.2015.08.008]

129 Tsutsumi Y, Yamamoto Y, Ino S, Ohigashi H, Shiratori S, Naruse H, Teshima T. Hepatitis B virus
reactivation with a rituximab-containing regimen. World J Hepatol 2015; 7: 2344-2351 [PMID: 26413224 DOI: 10.4254/wjh.v7.i21.2344]

130 Khan ZH, Ilyas K, Ghazanfar H, Khan HH, Hussain Q, Hammad S, Munir A, Asim R. Fatal Fulminant Hepatitis from Rituximab-induced Hepatitis B Reactivation in a Patient with Follicular Lymphoma: A Case Report and a Brief Review of Literature. Cureus 2018; 10: e2257 [PMID: 29725560 DOI: 10.7759/cureus.2257]

131 Ferri C, Greco F, Longombardo G, Palla P, Moretti A, Marzo E, Mazzoni A, Pasero G, Bombardieri S, Highfield P. Association between hepatitis C virus and mixed cryoglobulinaemia [see comment]. Clin Exp Rheumatol 1991; 9: 621-624 [PMID: 1662567]

132 Rosa D, Saletti G, De Gregorio E, Zorat F, Comar C, D'Oro U, Nuti S, Houghton M, Barnaba V, Pozzato G, Aigrignani S. Activation of naïve B lymphocytes via CD81, a pathogenetic mechanism for hepatitis C virus-associated B lymphocyte disorders. Proc Natl Acad Sci 2005; 102: 18544-18549 [PMID: 16339892 DOI: 10.1073/pnas.0509402102]

133 Charles ED, Green RM, Marukian S, Talal AH, Lake-Baakaar GV, Jacobson JM, Rice CM, Dustin LB. Clonal expansion of immunoglobulin M+CD27+ B cells in HCV-associated mixed cryoglobulinemia. Blood 2008; 111: 1344-1356 [PMID: 17942751 DOI: 10.1182/blood-2007-07-101717]

134 Ferri C, La Civita L, Zigkego AL, Pasero G. Viruses and cancers: possible role of hepatitis C virus. Eur J Clin Invest 1997; 27: 711-718 [PMID: 9352239 DOI: 10.1046/j.1365-2362.1997.1790728.x]

135 Mele A, Pulsoni A, Bianco E, Musto P, Szklo A, Sanpaolo MG, Iannitto E, De Renzo A, Martino B, Liso V, Andrizzzi C, Pusterla S, Dore F, Maresca M, Rapicetta M, Marcucci F, Mandelli F, Franceschi S. Hepatitis C virus and B-cell non-Hodgkin lymphomas: an Italian multicenter case-control study. Blood 2003; 102: 996-999 [PMID: 12714514 DOI: 10.1182/blood-2002-10-3230]

136 Giordano TP, Henderson L, Landgren O, Chiao EY, Kramer JR, El-Serag H, Engels EA. Risk of non-Hodgkin lymphoma and lymphoproliferative precursor diseases in US veterans with hepatitis C virus. JAMA 2007; 297: 2010-2017 [PMID: 17488966 DOI: 10.1001/jama.297.18.2010]

137 Monti G, Pioletti P, Saccardo F, Campanini M, Candela M, Cavallero G, De Vita S, Ferri C, Mazzaro C, Migliarese S, Osi E, Pietrogrande M, Galli M, Invernizzi F. Incidence and characteristics of non-Hodgkin lymphomas in a multicenter case file of patients with hepatitis C virus-related symptomatic mixed cryoglobulinemias. Arch Intern Med 2005; 165: 101-105 [PMID: 15642884 DOI: 10.1001/archinte.165.1.101]

138 Bunchorntavakul C, Mitramani R, Reddy KR. Advances in HCV and Cryoglobulinemical Vasculitis in the Era of DAAs: Are We at the End of the Road? J Clin Exp Hepatol 2018; 8: 81-94 [PMID: 29743799 DOI: 10.1016/j.jceh.2017.11.012]

139 Saadoun D, Pol S, Ferfar Y, Alric L, Hezode C, Si Ahmed SN, de Saint Martin L, Comarmond C, Bouyer AS, Musset L, Poynard T, Resche Rigon M, Cacoub P. Efficiency and Safety of Sofosbuvir Plus Daclatasvir for Treatment of HCV-Associated Cryoglobulinemia Vasculitis. Cureus 2017; 9: 22147667

140 Arcaini L, Besson C, Frigenti M, Fontaine H, Goldaniaga M, Casadeo M, Visentini M, Torres HA, Loutstaud-Ratti V, Peveling-Oberhag J, Fabris P, Rossotti R, Zignego AL, Pasero G. Viruses and cancers: possible role of hepatitis C virus. Eur J Cancer Prev 2013; 22: 28219772 DOI: 10.1097/01.jpc.0000438275.63295.22

141 Ramos-Casals M, Zigkego AL, Ferri C, Brito-Zerón P, Retamozo S, Casadeo M, Lampeprecht P, Mangia A, Saadoun D, Tzioufas AG, Younossi ZM, Cacoub P, International Study Group of Extrahepatic Manifestations related to HCV (ISG-EHCV). Evidence-based recommendations on the management of extrahaepatic manifestations of chronic hepatitis C virus infection. J Hepatol 2017; 66: 1282-1299 [PMID: 28219772 DOI: 10.1016/j.jhep.2017.02.010]

142 Saadoun D, Resche Rigon M, Sene D, Terrier B, Karras A, Perard L, Schoindre Y, Coppéré B, Blanc F, Musset L, Prete JC, Rosenzweig M, Cacoub P. Rituximab plus Peg-interferon-alpha/ribavirin compared with Peg-interferon-alpha/ribavirin in hepatitis C-related mixed cryoglobulinemia. Blood 2010; 116: 326-34; quiz 504 [PMID: 20439619 DOI: 10.1182/blood-2009-10-248518]

143 De Vita S, Quartuccio L, Isola M, Mazzaro C, Scaini P, Lensi M, Campanini M, Naclerio C, Tavoni A, Pietrogrande M, Ferri C, Mascia MT, Masolini P, Zabotti A, Maseri M, Roccatello D, Zigkego AL, Pioletti P, Gabriele A, Filipini D, Perrella O, Migliarese S, Galli M, Bombardieri S, Monti G. A randomized controlled trial of rituximab for the treatment of severe cryoglobulinemic vasculitis. Arthritis Rheum 2012; 64: 843-853 [PMID: 22147661 DOI: 10.1002/art.33313]

144 Desbois AC, Comarmond C, Saadoun D, Cacoub P. Cryoglobulinemia vasculitis: how to handle. Curr Opin Rheumatol 2017; 29: 343-347 [PMID: 28368978 DOI: 10.1097/BOR.0000000000000390]

145 Sène D, Ghielmini-Dalbin P, Amoura Z, Musset L, Cacoub P. Rituximab may form a complex with IgM kappa mixed cryoglobulin and induce severe systemic reactions in patients with hepatitis C virus-induced vasculitis. Arthritis Rheum 2009; 60: 3848-3855 [PMID: 19950292 DOI: 10.1002/art.25000]

146 Ishiguro N, Kawashima M. Cutaneous polymyositis nodosa: a report of 16 cases with clinical and histopathological analysis and a review of the published work. J Dermatol 2010; 37: 85-93 [PMID: 20175828 DOI: 10.1111/j.1346-8138.2009.00752.x]
Group. Hepatitis B virus-associated polyarteritis nodosa: clinical characteristics, outcome, and

Saadoun D, Conn DL, Czaja AJ, Wright TL, Brecher ME. Frequency and significance of

Sharma A, Sharma K. Hepatotropic viral infection associated systemic vasculitides-hepatitis B virus

Pagnoux C, Seror R, Henegar C, Mahr A, Cohen P, Le Guern V, Bienvenu B, Mouton L, Guillemin L, French Vasculitis Study Group. Clinical features and outcomes in 348 patients with polyarteritis nodosa: a systematic retrospective study of patients diagnosed between 1963 and 2005 and entered into the French Vasculitis Study Group Database. Arthritis Rheum 2010; 62: 616-626 [PMID: 20112401 DOI: 10.1002/art.27240]

Imboden JB. Involvement of the Peripheral Nervous System in Polyarteritis Nodosa and Antineutrophil Cytoplasmic Antibodies-Associated Vasculitis. Rheum Dis Clin North Am 2017; 43: 533-639 [PMID: 28961248 DOI: 10.1016/j.rdc.2017.06.001]

Pagnoux C, Mahr A, Cohen P, Guillemin L. Presentation and outcome of gastrointestinal involvement in systemic necrotizing vasculitides: analysis of 62 patients with polyarteritis nodosa, microscopic polyangiitis, Wegener granulomatosis, Churg-Strauss syndrome, or rheumatoid arthritis-associated vasculitis. Medicine (Baltimore) 2005; 84: 115-128 [PMID: 15758841 DOI: 10.1097/01.md.0000158825.87055.0b]

Wang CR, Yang CC. Adalimumab therapy in hepatitis B virus-negative polyarteritis nodosa: a case report. Medicine (Baltimore) 2018; 97: e11053 [PMID: 29923992 DOI: 10.1097/MD.0000000000011053]

Ginsberg S, Rosner I, Slobodin G, Rozenbaum M, Kaly L, Boulnam N, Awisat A, Hussein H, Novofovtovski I, Silayy A, Rimar D. Infliximab for the treatment of refractory polyarteritis nodosa. Clin Rheumatol 2019; 38: 2825-2833 [PMID: 30972576 DOI: 10.1007/s10067-019-04474-9]

Krusche M, Ruffner N, Köttler I. Tocilizumab treatment in refractory polyarteritis nodosa: a case report and review of the literature. Rheumatol Int 2019; 39: 337-344 [PMID: 30465270 DOI: 10.1007/s00296-018-4210-2]

Press J, Maslovitz S, Avinaoach I. Cutaneous necrotizing vasculitis associated with hepatitis A virus infection. J Rheumatol 1997; 24: 965-967 [PMID: 9150090]

Candan F, Ayan S, Tas F, Göke E, Elagoz S. Spontaneous renal laceration as the presenting feature of polyarteritis nodosa in a patient with familial Mediterranean fever after hepatitis A infection. Rheumatol Int 2005; 25: 475-477 [PMID: 15765217 DOI: 10.1007/s10067-005-0597-7]

Delia Rossa A, Tavoni A, Lorello S, Casula F, Bombardieri S. HBV and HCV infection, polyarteritis nodosa and mixed cryoglobulinaemia: a case report. Clin Rheumatol 2006; 19: 502-504 [PMID: 11147768 DOI: 10.1007/s1006700700138]

Garcia de La Peña Lefèvre P, Mouton L, Cohen P, Lhote F, Guillemin L. Polyarteritis nodosa and mixed cryoglobulinaemia related to hepatitis B and C virus coinfection. Ann Rheum Dis 2001; 60: 1068-1069 [PMID: 11602482 DOI: 10.1136/ard.60.11.1068]

Carson CW, Conn DL, Craza AJ, Wright TL, Brecher ME. Frequency and significance of antibodies to hepatitis C virus in polyarteritis nodosa. J Rheumatol 1993; 20: 304-309 [PMID: 8097250]

Saadoun D, Terrier B, Sennoun O, Sene D, Maisonneuve T, Musset L, Amoura Z, Rigon MR, Cauchy P. Hepatitis C virus-associated polyarteritis nodosa. Arthritis Care Res (Hoboken) 2011; 63: 427-435 [PMID: 20981809 DOI: 10.1002/acr.20381]

Néel A, Masseau A, Hervier B, Bossard C, Cauchy P, Pagnoux C, Hamidou MA. Life-threatening hepatitis C virus-associated polyarteritis nodosa successfully treated by rituximab. J Clin Rheumatol 2011; 17: 439-441 [PMID: 22089995 DOI: 10.1097/RHU.0b013e31823a58d7]

Guillemin L, Mahr A, Callard P, Godmer P, Pagnoux C, Leray E, Cohen P, French Vasculitis Study Group. Hepatitis B virus-associated polyarteritis nodosa: clinical characteristics, outcome, and
impact of treatment in 115 patients. Medicine (Baltimore) 2005; 84: 313-322 [PMID: 16148731 DOI: 10.1097/01.md.0000180792.80212.5c]

170 Lin CL, Kao JH. Perspectives and control of hepatitis B virus infection in Taiwan. J Formos Med Assoc 2015; 114: 901-909 [PMID: 26184565 DOI: 10.1016/j.jfma.2015.06.003]

171 Ungrasesat P, Koster MJ, Cheungpasitporn W, Wijampreecha K, Thonggrayoon C, Kroner PT. Inpatient burden and association with comorbidities of polyarteritis nodosa: National Inpatient Sample 2014. Semin Arthritis Rheum 2020; 50: 66-70 [PMID: 31362895 DOI: 10.1016/j.semarthritis.2019.07.009]

172 Chen WY, Lin KT, Chuang CY, Chen CY. Clinical studies of polyarteritis nodosa. Taiwan Yi Xue Hui Za Zhi 1977; 76: 982-989 [PMID: 25312]

173 Wang CR, Liu MF, Tsai RT, Chuang CY, Chen CY. Circulating intercellular adhesion molecules-1 and autoantibodies including anti-endothelial cell, anti-cardiolipin, and anti-neutrophil cytoplasm antibodies in patients with vasculitis. Clin Rheumatol 1993; 12: 375-380 [PMID: 8258240 DOI: 10.1007/BF02231583]

174 Tsai WL, Tsai IC, Lee T, Hsieh CW. Polyarteritis nodosa: MDCT as a “one-stop shop” modality for whole-body arterial evaluation. Cardiovasc Intervent Radiol 2008; 31 Suppl 2: S26-S29 [PMID: 17508232 DOI: 10.1007/s00270-007-9059-9]

175 Huang MN, Wu CH. Polyarteritis nodosa and antiphospholipid syndrome causing bilateral renal infarction. J Rheumatol 2009; 36: 197 [PMID: 19208533 DOI: 10.3899/jrheum.080601]

176 Li KJ, Hsieh SC, Toh YC, Yu CL. Clinical images: non-hepatitis B virus-related polyarteritis nodosa presenting with fever and diffuse intra-abdominal microaneurysms. Arthritis Rheum 2011; 63: 5357 [PMID: 21834065 DOI: 10.1002/art.30579]

177 Tsai HC, Liao HT, Tsai CY. Polyarteritis nodosa with intra-hepatic arterial haemorrhage. Liver Int 2020; 40: 2858-2859 [PMID: 32810375 DOI: 10.1111/liv.14641]

178 Lu FT, Ni YH. Elimination of Mother-to-Infant Transmission of Hepatitis B Virus: 35 Years of Experience. Pediatr Gastroenterol Hepatol Nutr 2020; 23: 311-318 [PMID: 32704492 DOI: 10.1523/pghn.2020.23.4.311]

179 Hong M, Bertolletti A. Tolerance and immunity to pathogens in early life: insights from HBV infection. Semin Immunopathol 2017; 39: 643-652 [PMID: 28685270 DOI: 10.1007/s00247-017-0641-1]

180 Travers RL, Allison DJ, Brettle RP, Hughes GR. Polyarteritis nodosa: a clinical and angiographic analysis of 17 cases. Semin Arthritis Rheum 1979; 8: 184-199 [PMID: 34221 DOI: 10.1016/s0049-0172(79)80007-4]

181 Cohen RD, Conn DL, Ilstrup DM. Clinical features, prognosis, and response to treatment in polyarteritis. Mayo Clin Proc 1980; 55: 146-155 [PMID: 6101626]

182 Fortin PR, Larson MG, Watters AK, Yeadon CA, Choquette D, Esdaile JM. Prognostic factors in systemic necrotizing vasculitis of the polyarteritis nodosa group—a review of 45 cases. J Rheumatol 1995; 22: 78-84 [PMID: 7699687]

183 Bae YD, Choi HJ, Lee JC, Park JJ, Lee YJ, Lee EB, Song YW. Clinical features of polyarteritis nodosa in Korea. J Korean Med Sci 2006; 21: 591-595 [PMID: 16891798 DOI: 10.3346/jkms.2006.21.4.591]

184 Sharma A, Pinto B, Dhokia A, Rathia M, Singhal M, Dhir V, Sharma K, Parkash M, Modi M, Vijayvergiya R, Sinha SK, Nada R, Minz RW, Singh S. Polyarteritis nodosa in north India: clinical manifestations and outcomes. Int J Rheum Dis 2017; 20: 390-397 [PMID: 27990777 DOI: 10.1111/1756-185X.12954]

185 Guillemin L, Lhote F, Gayraud M, Cohen P, Jarrousse B, Lortholary O, Thibult N, Casassus P. Prognostic factors in polyarteritis nodosa and Churg-Strauss syndrome. A prospective study in 342 patients. Medicine (Baltimore) 1996; 75: 17-28 [PMID: 8569467 DOI: 10.1097/00005179-199601000-00003]

186 Guillemin L, Lhote F, Cohen P, Jarrousse B, Lortholary O, Généreau T, Léon A, Bussel A. Corticosteroids plus pulse cyclophosphamide and plasma exchanges versus corticosteroids plus pulse cyclophosphamide alone in the treatment of polyarteritis nodosa and Churg-Strauss syndrome patients with factors predicting poor prognosis. A prospective, randomized trial in sixty-two patients. Arthritis Rheum 1995; 38: 1638-1645 [PMID: 7488285 DOI: 10.1002/art.1780381116]
