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**Pleiotropic effects of statins in the diseases of the liver**

Janicko M *et al.* Statins in liver disease

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**Abstract**

Statins are a class of molecules that inhibit HMG CoA reductase. They are usually prescribed as a lipid lowering medication. However, there is accumulating evidence that statins have multiple secondary effects both related and unrelated to their lipid-lowering effect. This narrative review of the literature aims to provide the reader with information from clinical studies related to the effect of statin and statins´ potential use in patients with liver diseases. In patients with advanced liver disease due to any etiology, statins exhibit an antifibrotic effect possibly through the prevention of hepatic sinusoidal microthrombosis. Two randomized controlled trials confirmed that statins decrease hepatic vein pressure gradient in patients with portal hypertension and improve the survival of patients after variceal bleeding. Lower rates of infections were observed in patients with cirrhosis who received statin treatment. Statins decrease the risk of hepatocellular carcinoma (HCC) in patients with advanced liver disease in general but particularly in patients with chronic hepatitis B and C. Statins in patients with chronic hepatitis C likely increase the virological response to the treatment with pegylated interferon and ribavirin and have the potential to decrease the rate of fibrosis. Finally, data from randomized controlled trials also confirmed that the addition of statin prolongs the survival of patients with advanced HCC even more than sorafenib. Statins are a very promising group of drugs especially in patients with liver disease, where therapeutic options can often be limited. Some indications, such as the prevention of re-bleeding from esophageal varices and the palliative treatment of HCC have been proven through randomized controlled trials, while additional indications still need to be confirmed through prospective studies.

**Key words:** Statins; Hepatitis; Cirrhosis; Esophageal varices; Hepatocellular carcinoma

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**Core tip:** The greatest benefit of statins seems to be in patients with advanced liver disease. Observational studies suggest that statins have an antifibrotic effect possibly through the prevention of hepatic sinusoidal microthrombosis, reduce the rate of infections and decrease the risk of hepatocellular carcinoma in all cirrhotics, but particularly in patients with chronic hepatitis B and C. Data from randomized controlled trials confirmed that statins decrease hepatic vein pressure gradient, prevent re-bleeding, and improve the survival of patients after variceal bleeding. Statins also seem to prolong the survival of patients with advanced hepatocellular carcinoma even more than those treated with Sorafenib, which is the current standard of care for these patients.

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**Introduction**

Statins are an inhomogeneous group of molecules that inhibit the activity of hydroxymetylglutaryl-coenzyme A reductase (HMG CoA reductase), a key enzyme in the synthesis of cholesterol. Statins were discovered as a byproduct in the search for new antimicrobial agents. The first statin (mevastatin) was the product of Penicillinum citrinium, but its clinical use was abandoned due to hepatotoxicity[[1](#_ENREF_1)]. Lovastatin was the first clinically successful statin to be used effectively[[2](#_ENREF_2)]. Scandinavian simvastatin survival study (4S) confirmed that statins reduce cardiovascular as well as general mortality in patients with atherosclerosis[[3](#_ENREF_3)].

 Individual molecules from the statin group differ in several important attributes. Lovastatin, simvastatin, fluvastatin and atorvastatin are lipophilic, whereas pravastatin and rosuvastatin are hydrophilic. Pravastatin is metabolized in the liver by sulfation, while lovastatin, simvastatin and atorvastatin are metabolized by cytochrome P-450 3A4. Fluvastatin and rosuvastatin are metabolized partially by cytochrome P-450 2C9[[4](#_ENREF_4)].

 Besides lipid lowering properties statins also exhibit multiple pleiotropic effects, which could be detrimental (*i.e.*, adverse effects) or beneficial. It is unknown whether the pleiotropic effects are directly related to the primary effect of the drug. Statins exhibit various antiatherogenic effects, such as the improvement of endothelial function, antioxidative, antiproliferative and antiinflammatory properties as well as neoangiogenesis[[4](#_ENREF_4),[5](#_ENREF_5)]. Additionally, statins reduce the risk of sudden cardiac death, deep vein thrombosis and fibrosclerotic aortic stenosis, while treatment with statins could influence the regression of left ventricular hypertrophy[[4](#_ENREF_4)]. Statins also exhibit multiple non-cardiovascular effects. For example, a cross-sectional analysis of three hospital databases showed that patients using statin had a 60% lower prevalence of Alzheimer’s disease[[6](#_ENREF_6)]. Besides Alzheimer’s, statins also reduce the risk of other types of dementia[[7](#_ENREF_7)] and have a lower prevalence of vitiligo, osteoporosis, rheumatoid arthritis and sclerosis multiplex[[4](#_ENREF_4)]. Increased risk of type 2 diabetes mellitus could be counted among the negative pleiotropic (adverse) effects of statin therapy[[8](#_ENREF_8)].

 Non-alcoholic steatosis (NAFLD) of the liver is considered a hepatic manifestation of metabolic syndrome[[9](#_ENREF_9)]. While the exact causality is unknown, the significant increase of both subcutaneous and visceral fat along with dyslipoproteinemia in the majority of these patients is closely associated with the accumulation of fat in the liver. In some patients the hepatic fat stimulates an inflammatory response that causes non-alcoholic steatohepatitis that in turn could progress to liver cirrhosis. Besides liver related morbidity and mortality, the presence of NAFLD is a significant and independent risk factor for cardiovascular events[[10](#_ENREF_10)]. Statins are prescribed in these patients to positively influence lipoprotein metabolism. Moreover, increasing evidence suggests that statins improve all aspects of NAFLD. Statins decrease the elevated plasmatic activity of liver enzymes[[11](#_ENREF_11)], while statins in monotherapy and in combination with antioxidants decrease hepatic fat accumulation[[12](#_ENREF_12),[13](#_ENREF_13)]. Prolonged administration of statins could also reduce liver fibrosis[[12](#_ENREF_12)]. Interestingly one study showed that statin therapy longer than two years in obese patients reduces the prevalence of liver steatosis[[14](#_ENREF_14)]. Statin treatment also reduces the risk of cardiovascular mortality, and the risk reduction is significantly greater in patients with elevated liver enzymes[[11](#_ENREF_11)].

 Another well-established indication for statins are cholestatic liver diseases, particularly primary biliary cholangitis, which is commonly associated with elevated total and LDL-cholesterol levels, and statins could partially reverse this negative effect[[15](#_ENREF_15)].

**Statins and chronic viral hepatitis B/C**

Chronic viral hepatitis B and C could progress to liver cirrhosis, which increases the risk of developing hepatocellular cancer (HCC). Chronic viral hepatitis, particularly hepatitis B, could lead to hepatocellular cancer even without cirrhosis[[16](#_ENREF_16),[17](#_ENREF_17)].

 The aim of the treatment of chronic hepatitis C is the elimination of the virus. An undetectable virus 24 weeks after the end of treatment is termed “sustained viral response (SVR)”. Interferon based therapy was the standard of care of chronic hepatitis C patients before direct-acting antivirals became available. The rate of SVR for interferon based therapy depends on multiple factors (IL28B gene polymorphisms, pre-treatment hepatitis C virus (HCV) viral load, HCV reduction dynamics, the degree of fibrosis, *etc*.)[[18](#_ENREF_18)].

***Statins and HCV RNA without concomitant antiviral treatment***

Multiple authors havereported the effect of statin treatment of HCV viral load. An in-vitro study conducted by Ikeda *et al*[[19](#_ENREF_19)] showed that fluvastatin, lovastatin, simvastatin and atorvastatin prevent the replication of HCV RNA, and that this effect is significantly stronger in fluvastatin compared to other statins. *In vivo* studies showed varied results. Forde *et al*[[20](#_ENREF_20)] compared three groups of patients with chronic hepatitis C. Group A consisted of patients with dyslipidemia on statin treatment (without specification) for at least 60 d prior to the HCV RNA quantification, group B included dyslipidemic patients without statin, and group C included patients without dyslipidemia and not on statin treatment. The authors did not report significant differences in HCV RNA levels among these three groups of patients. Fluvastatin dosed 80 mg daily led to the reduction of HCV RNA in 50% of patients, with the highest weekly reduction by 1.75 decadic logarithm. The reduction of HCV RNA occurred in the first four weeks of treatment in 82% patients with viral response. However, after the reduction of the dose the HCV RNA increased in 22% of responders in the following 2-5 wk[[21](#_ENREF_21)]. Another observational study from Romania showed a significant decrease of HCV RNA after treatment with either 40 mg of fluvastatin or 20 mg of lovastatin (mean levels of HCV RNA before treatment 2376074 ± 3427596 IU/mL, and 1321136 ± 1343570 IU/mL after treatment, *p =* 0.001).The administration of both statins was associated with significant reduction of proinflammatory signaling by IL6 and TNF-α, while the fluvastatin group also had lower IL-8 levels[[22](#_ENREF_22)]. On the other hand, a study by Sheridan et al. did not find significant differences in HCV RNA levels between patients treated with 40-80 mg of fluvastatin (+/- ω-3-polyunsaturated fatty acids) and controls after 12 wk of treatment. The main limitation of this study is, that it included 35% of patients that had already been diagnosed with cirrhosis and 45% that were non-responders to PEG IFN treatment[[23](#_ENREF_23)]. Fluvastatin treatment also had a surprisingly negative effect in HCV/HIV coinfected patients, where it led to a mild increase of HCV RNA (HCV RNA before treatment 5.63 ± 0.5 log10 IU/mL *vs* 5.84 ± 0.6 log10 IU/mL after treatment, *p =* 0.001), compared to no change in HCV RNA in the control group[[24](#_ENREF_24)].

 The effect of other statins on HCV RNA has not been proven in any studies. Simvastatin treatment for three months did not affect HCV RNA levels significantly[[25](#_ENREF_25)] and neither did the combination of simvastatin with sertralin[[26](#_ENREF_26)]. Twelve weeks treatment with rosuvastatin titrated to 40 mg daily led to the decrease of HCV RNA higher than one decadic logarithm only in one out of eleven patients[[27](#_ENREF_27)]. A meta-analysis showed a relatively small but significant decrease of HCV RNA (0.2 decadic logarithm decrease, 95%CI: 0.09-0.31, *p <* 0.001) in patients treated with fluvastatin,but lovastatin, simvastatin, atorvastatin and rosuvastatin had no effect on HCV RNA levels[[28](#_ENREF_28)]. These results suggest that standard statin therapy does not have a significant effect on the dynamics of HCV RNA viral load, with the possible exception of fluvastatin.

 Despite the dubious effects of statins on HCV viral load, there is a distinctive antifibrotic effect of this treatment in HCV infected patients. The data comes from a large observational study from Taiwan, performed in 1997-2010 included 226856 patients with chronic hepatitis C. Cirrhosis was present in 34273 patients. The incidence of cirrhosis during the follow-up was significantly higher in patients not taking statins (1311.2 *vs* 445.5 cases per 100000 person-years) Hazard ratios were 0.33 (95%CI: 0.31-0.36), 0.24 (95%CI: 0.22-0.25), and 0.13 (95%CI: 0.12-0.15) when statin users were compared with non-statin users with cumulative defined daily doses (cDDD) of 28-83, 84-365, and greater than 365 respectively[[29](#_ENREF_29)].

***Statins and HCV RNA with concomitant antiviral treatment***

The possibility of improving the treatment efficacy of standard antiviral treatment with the addition of statins has been evaluated with interferon based treatment, which was the standard of care before the development of direct acting antivirals. The efficacy of pegylated interferon with ribavirin was about 50%[[30](#_ENREF_30),[31](#_ENREF_31)]. The addition of a statin effectively enhanced the antiviral effect of this treatment, particularly fluvastatin, exhibiting synergistic inhibitory effect on HCV RNA replication[[19](#_ENREF_19)]. Several studies have explored this synergy in studies *In vivo*. Japanese authors reported the sustained virological response (SVR) rate in patients treated with PEG IFN and ribavirin with the addition of 20 mg fluvastatin to be as high as 67%, however it is important to note that this observational study did not have any control group[[32](#_ENREF_32)]. Another study explored the addition of 20 mg fluvastatin to PEG IFN + ribavirin treatment (46 patients) and compared them to a control group (48 patients). The duration of treatment was 48 wk in patients with complete early viral response (cEVR) and 72 wk in patients without cEVR, but with HCV RNA negativity in 13–36 wk of treatment. There was no difference in cEVR between the statin and control group (50% *vs* 54.2%), but patients with cEVR achieved SVR in the fluvastatin group more frequently than the control group (91.3% *vs* 65.4%, *p =* 0.042). Furthermore, patients in the control group relapsed significantly more often than in the fluvastatin group (39.4% *vs* 14.7% respectively, *p =* 0.027)[[33](#_ENREF_33)]. There is also anecdotal evidence that the addition of pitavastatin and eicosapentaenoic acid to the PEG IFN + ribavirin treatment could increase the rate of achieving SVR[[34](#_ENREF_34)]. These results were confirmed in a meta-analysis of five different studies that included fluvastatin, simvastatin, rosuvastatin and pitavastatin. The addition of statin doubled the chance of SVR (OR = 2.02, 95%CI: 1.38-2.94) as well as rapid and early viral responses[[35](#_ENREF_35)].

 The addition of statin to the PEG IFN + ribavirin treatment could influence not only the SVR, but also the risk of complications, such as progressive fibrosis or HCC. The HALT-C study included non-responders to the previous interferon based treatment with advanced fibrosis (Ishak score ≥ 3). Patients that did not achieve virological response (HCV RNA negativity in week 20 were treated with PEG IFN + ribavirin for the next 3.5 years. Patients who were concomitantly treated with statin displayed a decrease of liver fibrosis (-0.34 ± 0.94 points) compared to non-statin users, where fibrosis progressed (0.42 ± 1.42), *p =* 0.006. Overall, fibrosis progression was found in 10% of statin users and 29% of non-users (adjusted HR = 0.31; 95%CI: 0.10–0.97). Statin treatment did not significantly influence the histology activity index or the plasmatic activity of ALT[[36](#_ENREF_36)]. Similar data was reported from a registry-based study (ERCHIVES Registry) of 9135 HCV infected veterans treated with interferon based therapy in the years 2001–2014. Liver cirrhosis occurred in 1649 patients and HCC in 239 patients. The risk of cirrhosis in statin users was 44% lower (adjusted HR = 0.6, 95%CI: 0.53-0.68) and the risk of HCC was lower by 49% (adjusted HR = 0.51, 95%CI: 0.36-0.72) compared to non-users. The strongest antifibrotic effect was attributed to atorvastatin and fluvastatin[[37](#_ENREF_37)].

 The addition of statin to the interferon-based therapy has the potential to decrease the degree of fibrosis and the risk of HCC; however, patients with chronic hepatitis C receive statins less frequently compared to patients without HCV infection[[38](#_ENREF_38)]. The introduction of direct acting antivirals, with SVR rates up to 100% also in cirrhotics and nonresponders to previous IFN based treatment, limits the benefit of statin treatment in HCV infected patients[[39-41](#_ENREF_39)]. However, statins may potentially play a role in other aspects of chronic HCV infection[[42](#_ENREF_42)].

 There is no relevant information about the statin influence on hepatitis B virus both in terms of hepatitis B virus (HBV) DNA dynamics or fibrogenesis. However, in an earlier study we reported that cholesterol has a significant quadratic relationship with HBV DNA. Thus, patients with cholesterol levels above and below the normal range had higher levels of HBV DNA (Figure 1)[[43](#_ENREF_43)]. It has been well documented that HBV DNA level is the strongest predictor of fibrosis progression. According to Iloeje *et al*[[44](#_ENREF_44)] “the cumulative incidence of cirrhosis is 4.5% in patients with HBV DNA < 300 copies/mL compared to 36.2% in patients with HBV DNA ≥ 106 copies/mL(*p <* 0.001)”. However, it is unclear if statin treatment in hypercholesterolemic patients would in any way influence HBV DNA levels. There is some in vitro data that simvastatin might increase the antiviral activity of nucleot(s)ide analogues[[45](#_ENREF_45)], but it is unlikely that this information will have any clinical meaning, because of the high efficacy of currently available tenofovir and entecavir.

**Statins and fibrogenesis**

Statins in general positively influence endothelial dysfunction and this effect is also present in intrahepatic sinusoids. They show an anti-inflammatory effect in the inflammatory response caused by endotoxin, angiotensin II or hypovolemia, and the diminished activation of hepatic stellatae cells (HSC)[[46](#_ENREF_46)]. Statins inhibit non-canonical hedgehog signaling and cirrhotic portal hypertension[[47](#_ENREF_47)], resulting in a protective effect in ischemic hepatitis[[48](#_ENREF_48)] and have protective effects against the thrombosis of hepatic sinusoids and portal vein[[49](#_ENREF_49)].

 Parenchymal extinction theory of fibrogenesis proposes the microthrombosis of liver sinusoids as the driving force of inflammation and fibrosis. This is supported by the frequent finding of factor V Leiden mutations, protein C deficiency and increased factor VIII expression in cirrhotics[[50](#_ENREF_50)]. Thrombin, generated as the result of coagulation cascade activation, might activate HSC through protease activated receptors 1 and 4[[51](#_ENREF_51)]. The administration of statin in these circumstances increases protein C activity[[52](#_ENREF_52)] and decreases thrombin generation in plasma[[53](#_ENREF_53)]. One of the key prothrombotic factors in liver cirrhosis is von Willebrand factor antigen (vWF:Ag). This is released by endothelial cells and megacaryocytes and promotes the endothelial adhesion of thrombocytes, the transport and binding of factor VIII and thrombus formation. The level of vWF:Ag directly correlates with the degree of liver fibrosis (Figure 2). Maieron *et al*[[54](#_ENREF_54)] developed a novel scoring system, that included vWF:Ag divided by thrombocytes (VITRO) for the prediction of liver cirrhosis with AUC 0.893 compared to Forns score AUC 0.874, *p =* ns (Figure 3). The VITRO score is also more accurate for the noninvasive diagnosis clinically significant portal hypertension than the ELF or APRI score[[55](#_ENREF_55)]. Simvastatin and pravastatin significantly decrease vWF:Ag levels [SMD: –0.54, 95%CI: –0.87-(–0.21), *p =* 0.001], in contrast to fluvastatin, atorvastatin and rosuvastatin. The greatest decrease of vWF:Ag level was observed after 12 weeks of statin administration[[56](#_ENREF_56)]. Statins in animal models upregulate Kruppel-like factor 2(KLF2) signaling pathways that leads to the decrease of circulating vWF:Ag, decreased activation of HSC, and the regression of liver fibrosis[[46](#_ENREF_46),[57](#_ENREF_57),[58](#_ENREF_58)].

 Increasing evidence from interventional studies provides support for the microthrombotic theory of fibrosis. The Italian authors evaluated 48 wk of enoxaparin treatment in cirrhotic patients (Child Pugh 7-10 points) with parent portal vein. Enoxaparin decreased not only the incidence of portal vein thrombosis, but also the incidence of decompensation and mortality[[49](#_ENREF_49)]. It is unclear if statin treatment alone could decrease the incidence of portal vein thrombosis in cirrhotics; however, in patients with diagnosed malignancy, this treatment significantly decreases the cumulative incidence of deep vein thrombosis. Six months after the start of the statin the incidence of deep vein thrombosis was only sporadic[[59](#_ENREF_59)]. This data indicates the need to further study statins in compensated and decompensated cirrhotics aimed at the prevention of portal vein thrombosis.

**Statins and the clinical course of cirrhosis**

Two retrospective observational studies evaluated the effect of statins on clinical outcomes of cirrhotic patients. The first study included 81 cirrhotics on statin treatment and 162 controls. The median follow-up was 36 months in the statin group and 30 mo in the control group. There was no difference in etiology, age, Child-Pugh, MELD, HCC prevalence, beta blockers use, esophageal varices or selected biochemical parameters at inclusion. Patients in the statin group had a significantly higher prevalence of coronary heart disease and a lower prevalence of diabetes mellitus. The mean survival time of patients in the statin group was 10.8 years compared to 6.3 years in the control group (*p =* 0.06). The mean survival time of Child Pugh A patients was 14.4 years in the statin group and 7 years in the control group (*p =* 0.01). The adjusted hazard ratio for overall mortality was 0.53, *p =* 0.01 in statin users. The authors also reported lower risk of cirrhosis decompensation in statin users[[60](#_ENREF_60)]. Another study performed by Mohanty *et al*[[61](#_ENREF_61)] was registry-based and included patients with cirrhosis caused by hepatitis C infection between 1996-2009. The study cohort included 40 512 patients, 98% of which were male with an average age of 56 years, 2802 patients were using statins. The authors compared the propensity matched cohorts of statin users and non-users and found that patients using statin had a lower risk of decompensation (HR = 0.55, 95%CI: 0.39-0.77) and death (HR = 0.56, 95%CI: 0.46-0.69). These observational studies provide a solid foundation to consider a randomized controlled trial with statin in liver cirrhosis, despite the already decreased level of cholesterol in cirrhotics that correlates with the prognosis[[62](#_ENREF_62)].

**Statins and portal hypertension**

Changes in intrahepatic microcirculation, increased intrahepatic vascular resistance and splanchnic vasodilation are the main factors leading to portal hypertension[[63](#_ENREF_63)]. Nitric oxide (NO) is the main modulator of the vascular tonus both in the liver and in the splanchnic region. The physiological production of NO is associated with anti-fibrotic, anti-inflammatory and anti-thrombotic effects. Decreased NO production in the sinusoidal endothelial cells has a proinflammatory and profibrotic effect in the liver[[48](#_ENREF_48),[64](#_ENREF_64)]. Simvastatin increases NO production in hepatosplanchnic region, decreases vascular resistance, and ameliorates the postprandial increase of portal pressure in cirrhotic patients without a substantial effect on systemic circulation[[65](#_ENREF_65)]. Abraldes *et al*[[66](#_ENREF_66)] performed a randomized controlled trial in patients with portal hypertension that evaluated the efficacy of 20 mg simvastatin, later titrated to 40 mg on the hepatic vein pressure gradient (HVPG). The decrease of HVPG was greater in the statin group (8.3% ± 12.2% *vs* 1.6%± 12.3%, *p =* 0.041). Statin treatment led to the decrease of HVPG both in patients treated (-11%, *p =* 0.033) and not treated (-5.9%, *p =* 0.013) with beta-blocker. Statin treatment did not affect systemic circulation and the incidence of adverse effects was the same in the treatment and control group. Another prospective study by Pollo-Flores *et al*[[67](#_ENREF_67)] included 34 patients with portal hypertension. Fourteen patients received 40 mg of simvastatin and 20 patients received placebo for 3 mo. Three patients in the statin group were excluded because of a contrast medium reaction and newly diagnosed HCC, while seven patients were excluded from the control group. In the per-protocol analysis the authors reported the decrease of HVPG in the statin group compared to no change in the control group (2 ± 2.2 Torr *vs* 0 ± 1.1 Torr, *p =* 0.02). Primary endpoint (the decrease HVPG of at least 20% from the baseline or under 12 mmHg) was achieved in 55% of patients in the statin group and 0% of patients in the control group (*p =* 0.036). Clinical outcomes related to portal hypertension, particularly variceal re-bleeding, have been evaluated in the BLEPS study which was a multicenter double-blind randomized controlled trial. It included 69 patients in the active group that received 20 mg of simvastatin titrated to 40 mg after 15 d and 78 patients in the control group. Patients were followed up for 24 months. The primary endpoint was re-bleeding or death. Nine percent of patients in the statin group and 22% of patients in the control group died during the study (HR = 0.39, 95%CI: 0.15-0.99, *p =* 0.030). Simvastatin treatment reduced the relative risk of death compared to the placebo by 61%. The rate of re-bleeding did not differ significantly between the two groups. Two patients from the statin group developed rhabdomyolysis during the statin treatment[[68](#_ENREF_68)]. As practically all of the studies used simvastatin it is not clear if this effect is a class effect of all statins or is limited to simvastatin.

**Statins and infections in cirrhosis**

Infections are common in cirrhotic patients and increase mortality by approximately four-fold. Thirty percent of patients die in the first month after infection diagnosis and another 30% in the following year[[69](#_ENREF_69)]. Motzkus-Feagans *et al*[70] evaluated the effect of statin treatment on the incidence of infections. The study included 19379 patients with compensated cirrhosis from United States Veterans Health Administration database, with a mean follow-up of 1194 (365-3103) d. 2468 patients were receiving statin, the most common was simvastatin. Infection was diagnosed in 12.4% of patients during follow-up, with a mean time to infection of 608 d. The most common infections were pneumonia and skin infections. Statin treatment was associated with reduced infection rate and mortality rate in the whole cohort (aHR = 0.42, 95%CI: 0.36-0.48), as well as in the propensity score matched sample that included 503 statin users and 1760 statin non-users (aHR = 0.67, 95%CI: 0.47-0.95)[[70](#_ENREF_70)]. The question remains if statins improve the outcome of patients with severe infection or sepsis. Although no data exists about this particular issue in cirrhotic patients, there are many studies about this topic in the general population. Meta-analysis showed that patients with severe infections or sepsis, who were given statin had lower mortality for sepsis [aOR = 0.40 (95%CI: 0.23-0.57)], pneumonia [aOR = 0.33 (95%CI: 0.09-0.75)] and mixed infection-related mortality [aOR = 0.50 (95%CI: 0.18-0.83)] compared to statin non-users[[71](#_ENREF_71)]. These findings, however, were not confirmed in the randomized placebo-controlled trial with 40 mg of atorvastatin. Although statin treatment reduced the conversion rate from sepsis to severe sepsis (4% *vs* 24%, *p =* 0.007), no significant differences were found in mortality, length of hospital stay, or the number of re-hospitalizations[[72](#_ENREF_72)]. Therefore, more studies are needed to evaluate the clinical benefit of statins in cirrhotics with infections or sepsis.

**Statins and HCC**

Statin treatment does not generally affect the incidence of cancer or cancer related mortality[[73](#_ENREF_73)]. Hepatocellular carcinoma (HCC) occurs mostly in cirrhotic liver, with less than 20% of HCC occurring in non-cirrhotic liver[[74](#_ENREF_74)]. Therefore, statin therapy may indirectly influence the risk of HCC with its anti-fibrotic effect. Accumulated evidence from mostly observational studies suggest, that statins could also decrease the incidence of HCC by direct chemopreventive effect. The carcinogenesis of HCC along with potential targets for prophylaxis or treatment is depicted on figure 4. Multiple target sites of statins include the inhibition of post-translational prenylation of Ras/Raf proteins, inhibition of the proteasome pathway activation, limitation of the cyclin-dependent kinase inhibitors p21 and p27 degradation, and the blocking of Myc phosphorylation and activation, suppressing tumor initiation and growth[[75](#_ENREF_75)].

 Multiple clinical studies evaluated the effect of statin treatment on the incidence of HCC. Singh *et al*[[76](#_ENREF_76)] included 10 of the studies in the meta-analysis published in 2013. Seven studies were observational (3 case-control and 4 cohort studies) and three were RCTs, six studies included Western population and four studies Asian population[[73](#_ENREF_73),[77-85](#_ENREF_77)]. A total of 1459417 patients were included. The chemopreventive effect of statin administration was reported in half of the studies. Overall, statin administration was associated with lower risk of HCC (aOR = 0.63; 95%CI: 0.52– 0.76). The risk of HCC was lower in statin users in both the Western (aOR = 0.67; 95%CI: 0.53– 0.85) and Asian population (aOR = 0.52; 95%CI: 0.42– 0.64). These findings were confirmed in an updated meta-analysis by Shi et al that included 12 studies (6 case-control studies, 5 cohort studies and 1 randomised controlled study)[[73](#_ENREF_73),[77-82](#_ENREF_77),[85-89](#_ENREF_85)]. The relative risk of HCC in statin users was 0.58, (95%CI: 0.51-0.6), Figure 5[[90](#_ENREF_90)]. A smaller meta-analysis by Zhou *et al*, that included five observational studies[[78](#_ENREF_78),[85](#_ENREF_85),[87](#_ENREF_87),[91](#_ENREF_91),[92](#_ENREF_92)], also showed a significant risk reduction of HCC in statin users. Odds ratios were 0.63, 95%CI: 0.45-0.89 for atorvastatin and OR = 0.58, 95%CI: 0.40-0.85 for fluvastatin[[93](#_ENREF_93)]. The chemopreventive effect of statins was also described in patients with chronic hepatitis C without cirrhosis. A registry-based study from Taiwan included 35023 statin users and 225841 non-statin users. The authors reported a significant dose response relationship between statin use and the risk of HCC with an aHR of 0.66, 95%CI: 0.59-0.74, aHR = 0.47, 95%CI: 0.40-0.56 and aHR = 0.33, 95%CI: 0.25-0.42) in groups with cDDD of 28-89, 90-180 and more than 180 respectively[[89](#_ENREF_89)].

 The incidence of HCC in patients with chronic hepatitis B partially depends on the viral load. Levels of HBV DNA ≥ 10000 copies/mL are a significant predictor of HCC independent of ALT levels, HBeAg or the presence of liver cirrhosis[[94](#_ENREF_94)]. Statin use significantly decreased the risk of HCC in patients with hepatitis B in a registry-based observation from Taiwan with dose-dependent relationship. Adjusted hazard ratios were 0.66, 95%CI: 0.44-0.99, 0.41, 95%CI: 0.27-0.61 and 0.34, 95%CI: 0.18-0.67 for cDDD of 28-90, 91-365 and greater than 365 respectively[[85](#_ENREF_85)]. Another study from Hong Kong also reported that statin use was associated with 32% risk reduction of HCC development [weighted sub-hazard ratio (SHR) = 0.68; 95%CI: 0.48-0.97], however statins did not reduce the risk of mortality (weighted HR = 0.92; 95%CI: 0.76-1.11). The addition of statin to the standard nucleot(s)ide analogue treatment reduced the risk of HCC by 59% (weighted SHR 0.41; 0.19-0.89) compared to patients with only nucleot(s)ide analogue treatment[[95](#_ENREF_95)]. This corresponds with the presumed synergy between nucleot(s)ide analogue and statins for HCC risk reduction[[45](#_ENREF_45)].

 It has been shown that statins do not influence the incidence of cancer in the general population[[73](#_ENREF_73)]. Interestingly, the risk reduction seems to be significant in patients with chronic hepatitis B. An observational study by Chen et al. included 71847 patients with chronic hepatitis B. Statin users from this study had significantly lower risk of not only liver cancer but also all malignancies in general. The concomitant use of statin and metformin reduced the risk of malignancies even further in patients with chronic hepatitis B (Table 1)[[96](#_ENREF_96)].

Statins have also been tried as a concomitant therapy in patients with confirmed HCC. Two randomized controlled trials evaluated the role of statins in the treatment of advanced hepatocellular carcinoma. Japanese authors randomized 83 patients with non-resectable HCC undergoing transarterial chemoembolisation into 40 mg pravastatin and control group. The mean survival rate was significantly longer in the statin group (18 mo *vs* 9 mo)[[97](#_ENREF_97)]. These results were confirmed in a similarly designed German RCT that included 131 patients. Survival in the statin group was 20.9 mo, 95%CI: 15.5–26.3 compared to 12.0 mo, 95%CI: 10.3–13.7, *p =* 0.003 in the control group[[98](#_ENREF_98)]. Similar data was reported from observational studies in Taiwan and United States. The Taiwanese authors observed 20200 patients who received palliative treatment for HCC with median follow-up of 1.66 years. Statin treatment in this group was associated with lower HCC-related deaths in all stages of HCC. The risk of HCC-related death was reduced in 50% during 18 mo´ follow-up in patients with stage II and III[[99](#_ENREF_99)]. The American authors observed 1036 with early HCC (stage I or II) undergoing standard treatment for HCC. Patients who used statin lived significantly longer (23.9 *vs* 18.9 years, *p =* 0.047). However, after adjustment for confounders and immortal time bias, statin use did not confer lower risk of death (HR = 0.98, 95%CI: 0.80-1.20)[[100](#_ENREF_100)]. The reviewed studies suggest that the addition of statin to the treatment of patients with advanced HCC could extend survival by 5-9 mo. Surprisingly, the results of two RCTs are more favorable than the results of the SHARP study, where sorafenib treatment extended the survival of patients with non-resectable HCC by only 2.8 mo[[101](#_ENREF_101)].

**Limitations of statin use in patients with liver disease**

The conclusions that can be drawn from this review are limited by the mostly observational nature of the reviewed studies. However, the risk of bias seems to be relatively low because the control groups come from the same population as treated patients[[102](#_ENREF_102)]. Moreover, the evidence from RCTs is accumulating as well. There is not enough data to conclude if the various benefits of statins are related to the class effect, or if they are limited to particular a molecule (fluvastatin in chronic hepatitis B, simvastatin in portal hypertension, atorvastatin or fluvastatin in HCC risk reduction and pravastatin in palliative treatment of HCC).

 A second concern about statins in liver disease is the potential hepatotoxicity. There are two possible reactions to the the statin treatment. The most common is an asymptomatic, dose-dependent increase of plasma transaminase activity. This is present in as much as 2.7% of all high dose statin users, and is also dependent on the particular statin molecule. For example, rosuvastatin has the least chance of causing the elevation of liver enzymes[[103](#_ENREF_103)]. The second possibility is a drug induced liver injury (DILI) that is a result of idiosyncratic reaction, dose independent, and has potentially serious consequences. The diagnostic criteria for DILI increased of ALT ≥ 5-times above the upper limit of norm, or the increase of ALP ≥ 2-times above the upper limit of norm. According to data from the Swedish Adverse Drug Reactions Advisory Committee, only 73 cases of DILI, two deaths and one liver transplant occurred over 23 years (1988-2010) of statin use in Sweden. That represents about 1.2 cases of DILI per 100000 users. The lowest rate of DILI was reported for pravastatin and highest for fluvastatin[[104](#_ENREF_104)]. This rate is at the lower end of the range reported for general DILI incidence (from 1:10 000 to 1:100 000)[[105](#_ENREF_105)]**.** Despite this information, almost 50% of academic physicians hesitate to prescribe statin if ALT is greater than 1.5 times the upper limit of norm[[106](#_ENREF_106)].

 Finally, the most common complication of statin treatment that leads to the statin treatment being stopped is drug-induced myopathy. This condition is associated with single nucleotide polymorphism in SLCO1B1[[48](#_ENREF_48)]. The risk of statin-induced myopathy can be lowered by the administration of coenzyme Q10 alone, or in combination with selenium[[107](#_ENREF_107)].

**Conclusion**

This review summarized the potential uses of statins in patients with various liver disease states. In patients with chronic hepatitis C the addition of statin improves SVR rates of PEG IFN treatment and slows down fibrogenesis,. While it is not clear if statins influence HCV RNA levels, the main benefit is in patients with advanced fibrosis or cirrhosis. Statins have the potential to decrease the rate of fibrosis possibly through the prevention of hepatic sinusoidal microthrombosis. Statins decrease HVPG in patients with portal hypertension, and improve the survival of patients after variceal bleeding. Lower rates of infections were observed in patients with cirrhosis who received statin treatment. Statins decrease the risk of HCC in patients with advanced liver disease in general but particularly in patients with chronic hepatitis B and C. The addition of statin could prolong the survival of patients with advanced HCC. Most of the presented information comes from observational studies, randomized controlled trials are warranted to confirm these effects and allow the routine clinical use of statins in new indications.

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**Figure 1 Association between total cholesterol and hepatitis B virus DNA load[**[**43**](#_ENREF_43)**].** With permission from Elsevier.



**Figure 2 Dotplots for vFW: Ag according to fibrosis stage showing mean values and IQR.** *P <* 0.001 for all fibrosis stages, F3 *vs* F4 *P <* 0.0001[[54](#_ENREF_54)]. With permission from John Wiley and sons.



**Figure 3 Receiver operating characteristics curves for vWF:Ag, VITRO score and FORNS in the diagnosis of cirrhosis (F4).** AUC vWF:Ag = 0.835, VITRO score = 0.893 and FORNS = 0.874 (*P =* NS)[[54](#_ENREF_54)]. With permission from John Wiley and sons.

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**Figure 4 Pathogenesis of hepatocellular carcinoma and targets for chemopreventive agents.** Tyrosin kinase associated receptor pathways induce MAPK and PI3K–Akt kinase pathways in > 50% of HCCs. The resulting disruption of the mTOR pathwayh is seen in 40%–50% of cases of HCC, leading to inactivation of tumour suppressors such as PTEN. Statins block post-translational prenylation of Ras/Raf proteins, inhibit the activation of the proteasome pathway, limiting the degradation of the cyclin-dependent kinase inhibitors p21 and p27, and block Myc phosphorylation. Metformin activates AMPK, which inhibits the mTOR pathway. Thiazolidinediones inhibit the ubiquitin-proteasome system and extracellular signal-regulated kinase pathway. Insulin and sulphonylureas might promote hepatocarcinogenesis by increasing IGFR1 activity, enhancing growth-factor-dependent cell proliferation. AMPK: Adenosine monophosphate-activated protein kinase; HCC: Hepatocellular carcinoma; IGFR1: Insulin-like growth factor receptor 1; IR: Insulin receptor; MAPK: Ras mitogenactivated protein kinase; mTOR: Mammalian target of rapamycin; PI3K: Phosphatidylinositol 3-kinase; PPAR-γ: Peroxisome proliferator activated receptor γ[[75](#_ENREF_75)]. With permission from Nature publishing group.



**Figure 5 Overall meta-analysis of statin use and liver cancer risk[**[**90**](#_ENREF_90)**].**

**Table 1 Risk of overall and individual cancer with statin or metformin use in hepatitis B virus patients[**[**96**](#_ENREF_96)**]**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **All group (*n =* 71824)** | **No. of patients** | **Nonuser (*n =* 53037)** | **Only-metformin (*n =* 4774)** | **Only-statin (*n =* 8861)** | **M + S (*n =* 5152)** |
| **Adjusted HR (95%CI)** | **Adjusted HR (95%CI)** | **Adjusted HR (95%CI)** | **Adjusted HR (95%CI)** |
| Total cancer | 5434 | 1.00 | 1.03 (0.94–1.14) | 0.60 (0.55–0.66)c | 0.46 (0.40–0.52)c |
| Liver cancer | 1735 | 1.00 | 1.25 (1.06–1.47) b | 0.34 (0.27–0.42) c | 0.35 (0.27–0.45)c |
| Nonliver cancer | 3699 | 1.00 | 0.94 (0.83–1.06) | 0.72 (0.65–0.80) c | 0.50 (0.44–0.58)c |
| Lung cancer | 439 | 1.00 | 0.91 (0.66–1.26) | 0.51 (0.37–0.70) c | 0.49 (0.34–0.71)c |
| Stomach cancer | 144 | 1.00 | 0.77 (0.42–1.42) | 0.59 (0.35–1.00)a | 0.31 (0.14–0.69)b |
| Colorectal cancer | 572 | 1.00 | 1.14 (0.85–1.53) | 0.84 (0.65–1.09) | 0.51 (0.35–0.75)c |
| Esophagus cancer | 93 | 1.00 | 1.19 (0.61–2.31) | 0.38 (0.17–0.86)a | 0.30 (0.11–0.87)a |
| Pancreatic cancer | 127 | 1.00 | 1.33 (0.74–2.41) | 0.73 (0.40–1.31) | 0.70 (0.34–1.43) |
| Prostate cancer | 225 | 1.00 | 0.94 (0.59–1.50) | 0.77 (0.51–1.15) | 0.63 (0.37–1.05) |
| Breast cancer | 288 | 1.00 | 0.80 (0.47–1.32) | 0.91 (0.63–1.33) | 0.56 (0.33–0.95)a |
| Cervical cancer | 105 | 1.00 | 0.70 (0.31–1.58) | 0.67 (0.35–1.25) | 0.28 (0.10–0.79)a |
| Other cancers | 1706 | 1.00 | 0.91 (0.76–1.09) | 0.51 (0.42–0.64)c | 0.75 (0.65–0.88)c |

a*P* < 0.05, b*P* < 0.01, c*P* < 0.001 *vs* control. Adjusted HR is adjusted for baseline propensity score. HBV: Hepatitis B virus; HR: Hazard ratio; M: Metformin; S: Statin.