OPINION ARTICLE

Statin (3-hydroxy-3-methylglutaryl-coenzyme A reductase inhibitor)-based therapy for hepatitis C virus (HCV) infection-related diseases in the era of direct-acting antiviral agents [version 1; referees: 1 approved, 1 approved with reservations]

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Abstract

Recent improvements have been made in the treatment of hepatitis C virus (HCV) infection with the introduction of direct-acting antiviral agents (DAAs). However, despite successful viral clearance, many patients continue to have HCV-related disease progression. Therefore, new treatments must be developed to achieve viral clearance and prevent the risk of HCV-related diseases. In particular, the use of pitavastatin together with DAAs may improve the antiviral efficacy as well as decrease the progression of liver fibrosis and the incidence of HCV-related hepatocellular carcinoma. To investigate the management methods for HCV-related diseases using pitavastatin and DAAs, clinical trials should be undertaken. However, concerns have been raised about potential drug interactions between statins and DAAs. Therefore, pre-clinical trials using a replicon system and human hepatocyte-like cells from human-induced pluripotent stem cells should be conducted. Based on these pre-clinical trials, an optimal direct-acting antiviral agent could be selected for combination with pitavastatin and DAAs. Following the pre-clinical trial, the combination of pitavastatin and the optimal direct-acting antiviral agent should be compared to other combinations of DAAs (e.g., sofosbuvir and velpatasvir) according to the antiviral effect on HCV infection, HCV-related diseases and cost-effectiveness.
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Introduction
Hepatitis C virus (HCV) infection is an important public health problem worldwide, as many HCV-infected patients develop liver cirrhosis and/or hepatocellular carcinoma (HCC)1.

Recent improvements in the treatment of HCV infection have focused on the use of direct-acting antiviral agents (DAAs)3. However, despite successful viral clearance, many patients with advanced fibrosis continue to have HCV-related disease progression3–5. Therefore, new treatments must be developed to achieve viral clearance and prevent the risk of HCV-related diseases.

Statins (inhibitors of 3-hydroxy-3-methylglutaryl-coenzyme A reductase) have been proposed as a new candidate for the treatment of HCV infection. According to previously conducted clinical studies using statins in HCV-infected patients3–5, the antiviral effect of statins on HCV infection depends on the type of statin used. Among various statins, pitavastatin showed the highest antiviral efficacy against HCV genotype 1b infection in vitro6. Therefore, pitavastatin represents a novel candidate for the treatment of HCV infection.

Clinical trials for pitavastatin against HCV infection
After performing a search of the PubMed database, we identified three clinical trials using pitavastatin for the treatment of HCV infection8,9,10. When Shimada et al.7 considered two reports published in 2010 investigating the antiviral efficacy of pitavastatin against HCV infection in vitro8,9, they conducted a randomized controlled trial7. The proof-of-concept studies demonstrated the antiviral efficacy and safety of pitavastatin against HCV infection using a replicon system and human hepatocyte-like cells from human induced pluripotent stem cells (hiPSCs)10, and these effects of pitavastatin were confirmed in the randomized controlled trial7,11. Indeed, this series of studies7,11 would be the first to report the clinical applications of hiPSCs12. After the clinical trial performed by Shimada et al., the antiviral efficacy and safety of pitavastatin against HCV infection were further confirmed in two clinical studies8,10. Thus, investigating the antiviral efficacy and safety of pitavastatin against HCV infection using a replicon system and human hepatocyte-like cells from hiPSCs10 constitutes a rational approach to discover new drugs and/or new therapeutic methods.

Clinical trials to investigate the management methods of HCV-related diseases in the era of DAAs
Butt et al.13 showed that statin use was associated with improved antiviral efficacy as well as decreased progression of liver fibrosis and a reduced incidence of HCC among a large cohort of HCV-positive veterans. Furthermore, the use of statins among patients with HCV and compensated cirrhosis (n=40,512) was associated with a more than 40% lower risk of cirrhosis decompensation and death14. Moreover, statin users showed a significant reduction in the incidence of HCC15. In addition, pitavastatin showed anti-cancer effects against human hematoma cell lines16–18.

DAAs in combination with statins have been shown to generate increased antiviral efficacy against HCV infection19. However, concerns have been raised about the drug interactions between various statins and DAAs20. For instance, simvastatin and lovastatin should be avoided in patients with HCV infection who are using boceprevir or telaprevir as a DAA21. Atorvastatin should be avoided in patients with HCV infection who are using telaprevir21, and pravastatin plus boceprevir may also pose risks21. Although rosuvastatin could be considered for use in combination with telaprevir and boceprevir21, the drug interactions between pitavastatin and DAAs remain unknown.

Furthermore, according to our search of the PubMed database and UMIN Clinical Trials Registry System (http://www.umin.ac.jp/icdr/index.html), no clinical trial has been conducted for the combination of statins and DAAs. Therefore, although there is likely only a minimal additive benefit for viral clearance using statins, as new combinations of DAA (sofosbuvir and velpatasvir) therapy have shown sustained virologic response (SVR) rates above 95%12, conducting a clinical trial for the combination of pitavastatin and DAAs may be meaningful to investigate management methods to prevent fibrosis and cirrhosis or the development of HCC and other HCV-related diseases in the era of DAAs. However, in pre-clinical trials, the antiviral effects of the combination of pitavastatin and DAAs should be evaluated using a replicon system10. Furthermore, hepatotoxicities should also be evaluated for the combination of pitavastatin and DAAs using human hepatocyte-like cells from hiPSCs11. Using a replicon system and human hepatocyte-like cells from hiPSCs in a pre-clinical trial11, an optimal direct-acting antiviral agent could be selected for use in the combination of pitavastatin and DAAs. After the pre-clinical trial, the combination of pitavastatin and the optimal direct-acting antiviral agent should be compared with other DAA combinations (e.g., sofosbuvir and velpatasvir) according to their antiviral efficacy against HCV infection and prevention of HCV-related diseases. Furthermore, because the cost for DAA combination treatment is very high ($83,000 to $153,000 per course of treatment)22, the new and effective hepatitis C treatments seem beyond the reach of low- and middle-income countries23. However, the cost ($0.79 to $2.59 per day) of pitavastatin (2mg) is low (http://www.pharmacychecker.com/generic/price-comparison/pitavastatin/2+mg/). Therefore, the above-mentioned comparison should also be investigated in light of cost-effectiveness.

In conclusion, a pre-clinical trial investigating the combination of pitavastatin and DAAs against HCV infection using a replicon system and human hepatocyte-like cells from hiPSCs10,11 represents a rational approach to discovering a new therapeutic method.

Author contributions
All authors (Reem Mohamed Fathy El-Shenawy, Sara Kishta and Sobhy Kishta) equally contributed to the writing of the manuscript.

Competing interests
No competing interests were disclosed.

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References


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This article would be improved with statements quantifying how common progression to disease occurs in patients treated with anti-HCV compounds in the absence of other drugs. The main point of the article, that there should be pre-clinical trials to test for efficacy and toxicity of dual therapy with Pitavastatin are valid. However, a replicon system in hepatocellular cells is not likely to duplicate the full gamut of potential side effects seen in humans receiving pitavastatin and anti-HCV drugs. Although it is an important place to start, the risk that complications will not be revealed in a preclinical trial using this system should be acknowledged.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Competing Interests: No competing interests were disclosed.

Author Response 28 Jun 2016
Sara Kishta, National Research Center, Egypt, Egypt

First of all, thank you very much for your suggestions. We think that your comments were very helpful. We agree with your suggestions. If you agree with our revisions, we are very happy.

1) According to your suggestions, we added the following sentences to our text (please see, last paragraph and conclusion).

On the other hand, a replicon system in hepatocellular cells is not likely to duplicate the full gamut of potential side effects seen in patients with HCV infection receiving pitavastatin and DAAs. The risks (potential side effects) that will not be revealed in a pre-clinical trial using this system should be acknowledged. Therefore, in order to identify the risk (potential side effects) that will not be revealed in a pre-clinical trial using this system, the evaluations using human neurons and human cardiomyocytes from hiPSCs24, 25 should also be done in pre-clinical trials.

In conclusion, a pre-clinical trial investigating the combination of pitavastatin and DAAs against HCV infection using a replicon system, human hepatocyte-like cells, human neurons and human cardiomyocytes from hiPSCs9,11,24,25 represents a rational approach for discovering a new therapeutic method.
2) We revised our abstract. Pre-clinical trials using a replicon system, human hepatocyte-like cells, human neurons and human cardiomyocytes from human-induced pluripotent stem cells should be conducted.

3) We added the new references (No. 24 and 25) in the reference section.

**Competing Interests:** No competing interests were disclosed.

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Recent improvements in treatment strategies for Hepatitis C virus infected patients have been successful at reaching sustained virological responses yet did not prevent liver fibrosis and progression to hepatocellular carcinoma.

The idea proposed in this paper that new treatments must be developed to achieve viral clearance while preventing disease progression although meritorious is not original and should be complemented with further understanding of mechanisms underlying hepatitis C disease progression in the presence of sustained virological response (i.e. absence/ or low level of viral replication).

The authors stated that pitasvatin represents a novel candidate to treat HCV infection yet the rationale for selecting this statin, as the likely candidate is not clearly justified. The authors based their selection on data from a retrospective analysis demonstrating no significant difference in sustained virological response rate per protocol analysis between patients treated with PEG-IFN/ribavirin in the presence or absence of pitavastatin. Findings from a second prospective trial demonstrates the safety of pitavastatin in combination with Peg IFN plus ribavirin although the decrease in HCV RNA was only significant at 2 (4 and 12 weeks) of 6 evaluated time points of treatment. The third trial used pitavastin in combination with eicosapentaenoic acid and identified this combination therapy as predictive of sustained virological response in multifactorial analysis only after genetic variation in IL28B was excluded from these factors. Thus the authors could have presented a better rationale for the selection of this statin.

An additional point of discussion that should have been included in this paper would have been the evidence that statin use decreased progression to liver fibrosis and hepatocellular carcinoma yet this effect was independent of having attained a sustained virological response. These findings raised the possibility that the statins-mediated delay in liver fibrosis are related to their immunomodulatory rather than their antiviral effects. This assumption is supported by findings demonstrating little or no effect of statins use on HCV replication.

**I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.**
Competing Interests: No competing interests were disclosed.