Elimination half-life may explain the relative efficacy of boceprevir and telaprevir in the treatment of hepatitis C virus genotype 1.

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(Article begins on next page)
**Clinical Infectious Diseases**

Comment on "The Relative Efficacy of Boceprevir and Telaprevir in the Treatment of Hepatitis C Virus Genotype 1"

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Dear Editor,

Please find as online submission our manuscript entitled Comment on “The Relative Efficacy of Boceprevir and Telaprevir in the Treatment of Hepatitis C Virus Genotype 1” to be considered for publication in Clinical Infectious Diseases.

We appreciated the analysis by Kirean and colleagues trying to assess the efficacy of telaprevir and boceprevir in the absence of direct comparison trials. We suggest that the observed superiority in telaprevir recipients could be justified by the drug longer elimination half-life and therefore forgiveness. We therefore apply what we learnt from antiretroviral treatment experience and we suggest that adherence should be taken into account, measured and enhanced by clinicians and researchers.

Regarding potential conflicts of interest, no author has specific funding to disclose (a transparency declaration is at the end of the document).

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Giovanni Di Perri,
We hope that this manuscript can be of interest to the readers of CID, and look forward to receiving from you.

Yours truly.

Dr. A. Calcagno
Comment on “The Relative Efficacy of Boceprevir and Telaprevir in the Treatment of Hepatitis C Virus Genotype 1”

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Dear Editor,

in their very comprehensive meta-analysis Dr. Kieran and co-workers compared the efficacy of telaprevir and boceprevir as third agents in the treatment of chronic HCV hepatitis; they found a significantly greater efficacy rate for telaprevir in the specific setting of prior relapers as compared to standard pegylated interferon/ribavirin (PegIFN/RBV) therapy [1]. Such a meta-analysis is so far the only attempt to compare the two new anti-HCV antivirals, which have been released into the market without any preference for either agent in treatment recommendations [2]. A number of parameters have been proven to influence the outcome of anti-HCV treatment, such as genetic variation in *IL28B*, the type of PegIFN administered, baseline HCV-RNA, RBV pharmacokinetic exposure and degree of liver fibrosis. As stated by the Authors, dependency of the treatment outcome on the third drug may be thus be rather variable, and even of borderline significance when multiple favourable factors coincide in the same patient. We think that further to what suggested by the Authors in terms of possible reasons accounting for the higher efficacy of telaprevir in prior relapers, its longer elimination half-life ($t_{1/2}$, 9-11 hours) as compared to boceprevir (3.4 hours) should also be taken into consideration [3,4]. In anti-HIV therapy a longer half-life is the major determinant of what we call “forgiveness”, such as the property of maintaining effective concentrations in spite of a missed dose of the drug/regimen [5]. In the field of antiretroviral therapy, where numerous head-to-head comparisons have been made, a tendency to a better virological outcome is almost always recognizable in favour of the regimen containing the drug/s with longer half-life [6-8], with the notable exception of integrase inhibitors (that being associated with a faster viral clearance may compensate
for the drug shorter half-life) [9]. Although patients’ adherence has been far less characterized in the HCV setting than in antiretroviral therapy, we might reasonably envisage how patients under triple anti-HCV therapy taking oral drugs three times daily are at risk of suboptimal adherence. Supposing an equal degree of non-adherence, effective pharmacokinetic exposure of telaprevir persist longer than in case of boceprevir intake when a dose is missed, thus allowing a greater chance of maintaining adequate antiviral concentration despite of irregular drug intake. This pharmacokinetic property of telaprevir has been recently further testified by the successful validation of twice daily intake of the drug at equal total daily dose [10]. Since controlled head-to-head comparative trials between telaprevir and boceprevir are unlikely to be performed (and might soon lose interest with the new anti-HCV drugs being developed) the meta-analytic comparison carried out by the Authors might remain the sole to rely upon. Based on these considerations we believe that whenever patient’s adherence is perceived to be particularly at risk, the choice of telaprevir might provide an advantage in terms of pharmacokinetic coverage.

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References

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Transparency Declarations

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