APASL consensus statements and recommendation on treatment of hepatitis C

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Abstract The Asian-Pacific Association for the Study of the Liver (APASL) convened an international working party on the “APASL consensus statements and recommendation on management of hepatitis C” in March, 2015, in order to revise “APASL consensus statements and management algorithms for hepatitis C virus infection (Hepatol Int 6:409–435, 2012)”. The working party consisted of expert hepatologists from the Asian-Pacific region gathered at Istanbul Congress Center, Istanbul, Turkey on 13 March 2015. New data were presented, discussed and debated to draft a revision. Participants of the consensus meeting assessed the quality of cited studies. Finalized recommendations on treatment of hepatitis C are presented in this review.

Keywords APASL · DAAs · HCV · Interferon-free · Turkey

Introduction

The major aim of antiviral treatment for chronic hepatitis C is to prevent liver-related complications, including HCC, by achievement of sustained virologic response (SVR) [1, 2].
The combination of peginterferon plus ribavirin can lead to ~50 and ~80 % SVR in patients infected with HCV genotype (GT)-1/GT-4 or HCV GT-2/GT-3, respectively [3]. In the direct-acting antivirals (DAAs) era, DAAs with peginterferon plus ribavirin can shorten the treatment duration and lead to ~90 % SVR in HCV GT-1-infected patients with interleukin-28B (IL28B, IFN-lambda 3) favorable single nucleotide polymorphism (SNP). But these treatments have many adverse events associated with interferon use which hamper the patients in accomplishing the treatments. Recently, interferon-free treatment has played a central role in the eradication of HCV. In this article, we aim to introduce the recent advances of interferon-free therapies for the patients with chronic HCV various genotypes in Asian-Pacific countries. Grading of evidence and recommendations are shown in Supplementary Table 1.

**Interferon supersensitive group**

**Some viral and host factors are related to supersensitivity to the interferon-based therapy**

HCV GT-2 is supersensitive to interferon-based therapy. The sustained virological response (SVR) rates with 24-week peginterferon and ribavirin treatment in chronic HCV GT-2 are around 80 % in western countries [4–8]. In the chronic HCV GT-2 trial from Taiwan, the SVR rates by treating with peginterferon and ribavirin can be higher than 95 % if the patients achieve a rapid virologic response (RVR) [9].

For chronic HCV GT-1 patients with RVR, high SVR rates, near 90 %, can be obtained by the peginterferon and ribavirin combination therapy [10]. The SVR rates were as high as 95 % by treating with peginterferon and ribavirin combination therapy in chronic HCV GT-1 patients with RVR and low viral load at baseline, even with the treatment duration shortened to 24 weeks [11–14]. The study by Atlanta Medical Center also demonstrated that the 24-week peginterferon and ribavirin regimen may have equal efficacy as a 28-week lead-in then boceprevir and peginterferon plus ribavirin triple therapy in chronic HCV GT-1 patients with low viral load and RVR [15].

Several genome-wide association studies have demonstrated that host SNPs near the IL28B gene are associated with SVR to the treatment with peginterferon alfa and ribavirin in chronic hepatitis C patients [16–18]. These SNPs are also associated with spontaneous clearance of HCV in acute HCV infection. The various distributions of IL28B polymorphisms among different populations worldwide may, at least partly, explain the heterogeneity in the responses to interferon-based treatments among different ethnic groups. The IL28B SNPs are strongly associated with SVR rates in patients who are infected with HCV GT-1 or GT-4 and receive combination treatment with peginterferon alfa and ribavirin [16–21]. However, the association between IL28B variations and treatment response in patients infected with HCV GT-2 or GT-3 is still controversial [22–24].

IL28B variations are associated with very early on-treatment viral kinetics in chronic hepatitis C patients who undergo interferon alfa-based therapy, and are the strongest pretreatment predictor of treatment response in patients infected with HCV GT-1 [19, 21–24]. Regarding the retreatment with peginterferon and ribavirin in chronic HCV GT-1 and GT-2 patients, the SVR rate for prior-relapser was quite good [25, 26]. However, the response was only found in the relapsers with favorable IL28B genotypes. Therefore, for the countries in which direct acting antiviral agents are available, generic drugs are not available, and only have limited resources, peginterferon and ribavirin combination therapy may be considered in chronic HCV GT-1, GT-4, or GT-6 treatment-naive patients with low viral load and favorable IL28B genotypes and in HCV GT-2 or GT-3 naïve patients [27]. For the treatment-experienced patients, the peginterferon and ribavirin combination therapy can only be considered in relapsers with favorable IL28B genotype and non-cirrhotic patients.

#1 Consensus statements and recommendation on interferon supersensitive group

1. For the countries in which direct acting antiviral agents are not available, or only have limited resources, peginterferon and ribavirin combination therapy may be considered in chronic HCV GT-1, GT-4, or GT-6 treatment-naive patients with low viral load and favorable IL28B genotypes and in HCV GT-2 or GT-3 treatment-naive patients. (A1)

2. For the treatment-experienced patients, the peginterferon and ribavirin combination therapy can only be considered in relapsers with favorable IL28B genotype and non-cirrhotic patients. (B2)

Is interferon still needed in DAA affordable countries?

The development of direct acting antiviral agents and their inclusion in all-oral, interferon-free regimes has been the central element of the revolution in therapy for HCV infection. These new treatments are safe and very effective and there are virtually no medical reasons to withhold therapy [28]. Penetration of new interferon-free therapies into standard management plans in many countries in the Asian-Pacific region has been very slow despite
outstanding responses to therapy in virtually every sub-
group of patients treated to date [29, 30]. The delay in
making these drugs available to patients is their high cost
which is limiting uptake—irrespective of whether funding
is patient-based or by government reimbursement. Restricted access to therapy is not limited to resource-re-
stricted countries as many people living in countries with
relatively high gross domestic products (GDP) still have
limited availability. It is within this context that providers
of health care are attempting to determine if there is a place
remaining for the less expensive interferon-containing
regimes in the emerging treatment paradigms.

Interferon has been a partner in the treatment of HCV
for three decades and has played an important role in
curing many patients and reducing their subsequent risk
of cirrhosis or HCC [31]. Indeed, we have learnt much in
the decades of experience gained using interferon-contain-
ing regimes. Optimising dosing regimens [5], PEGy-
lation to improve pharmacokinetics [32], adding ribavirin
[33] and discovering host genetic and other factors as
well as viral characteristics that influence response [34]
have all been major advances that enhanced treatment response rates. However, there are a number of disad-
vantages of interferon therapy that have limited its uptake
and ensured that its role in antiviral therapy for HCV will
eventually become largely obsolete. The side-effect pro-
file and adverse effects of interferon therapy are sub-
stantial. In one report from Asia, up to 50 % of patients
were judged to be unsuitable to commence therapy. Interferon-based therapy is lengthy—up to 1 year for
genotype 1 patients—and requires substantial human and
laboratory infrastructure to ensure safety, compliance and
optimal dosing and treatment duration. The nature of the
therapy limits its use to major treatment centers with only
limited, if any, uptake in isolated areas. For these reasons,
treatment uptake rates with interferon-based regimes have
been modest, with some countries reporting treatment rates of only 5 % even when diagnosis rates were greater
than 50 % [35].

In contrast, direct acting antiviral regimes are very well
tolerated and treatment duration is shorter potentially falling to 8 weeks in some HCV populations [36]. Studies
to date have shown outstanding treatment response rates
across all genotypes, in previously treated and untreated
subjects, pre and post liver transplant patients, HIV co-
infected individuals and those with more advanced liver
disease [37–42]. As stated earlier, there are virtually no
medical reasons to withhold therapy. In this context it is
not flippant to suggest that, at present, the only indication
for interferon-based therapies is no access to direct acting
antiviral therapies.

Arguments in favor of maintaining interferon therapies
are that Asian patients respond reasonably well to these
therapies, due in part to the enrichment of the population
with the favorable IL28B allele. In HCV GT-1 Asian
subjects, SVR of 75 % have been reported [43], with even
more impressive responses in HCV GT-2 and GT-3 sub-
jects. However, the issues that limit interferon uptake still
exist as does on-treatment side effects and infrastructure
requirements. Thus with current rates of therapy and con-
tinued use of interferon-based regimes, most countries in
the Asian-Pacific region will continue on the inevitable path towards the peak of population morbidity
and mortality predicted from HCV epidemiological studies
and judged to be 10–20 years hence. Reducing the regional
impact of this disease should be a major goal of govern-
ments in the Asian-Pacific region and this aim increases
the urgency with which treatments should be made available.
Of course, to have any impact on HCV-related disease
burden, access to therapy must be increased. Without
increasing access to care, the number of treated patients
will not change. In the absence of such policies to increase
access to direct acting antivirals, we are going to com-
pound global inequalities in treatment and disease burden
and individual suffering. The efforts by Gilead to assist in
the provision of their medications to the 90 countries with
the lowest GDP are to be congratulated. Global viral
eradication was never going to be feasible with interferon-
containing regimes. In contrast, there is a distinct possi-
bility of eliminating HCV in the Asian-Pacific region with
oral antiviral agents if sufficient attention is also paid to
primary prevention.

**ALL-oral treatment for HCV GT-1 infection**

In the Asian-Pacific region, except Australia, Iran, New
Zealand, Philippines and Thailand, HCV GT-1b is the main
subgenotype in HCV GT-1 [3]. During the interferon era,
HCV GT-1 was the most intractable among all HCV GTs
when patients were treated with peginterferon plus rib-
avirin [1]. Combined peginterferon plus ribavirin treatment
for 48 weeks led to only ~50 % SVR in HCV GT-1 and
high viral load. The addition of HCV NS3/4A protease
inhibitors such as telaprevir, boceprevir or simeprevir to
peginterferon plus ribavirin can improve SVR rates and
shorten the treatment duration. These therapies can lead to
70–80 % and 90–100 % SVR in treatment-naı ¨ ve and previ-
ous-treatment relapers with HCV genotype 1b infection,
respectively, but only ~30 % SVR in previous-treatment
null responders [1, 44–48].

Furthermore, the combination of peginterferon plus
ribavirin with or without protease inhibitor therapies usu-
ally results in various adverse events, which can occa-
sionally be serious in certain patients despite those being
treated with these therapies having been selected on the
basis of SNPs of the IL28B gene and/or the inosine triphosphate pyrophosphatase (ITPA) gene [16–18, 49]. It is difficult to use peginterferon in interferon-ineligible/intolerant patients and it is hard to cure HCV-infected patients with unfavorable IL28B or with advanced liver fibrosis using interferon-including regimens [50, 51].

In 2010, interferon-free treatment for chronic HCV GT-1 infection was reported for the first time [52]. In this INFORM-1 trial, the oral combination of a nucleoside analogue polymerase inhibitor and protease inhibitor provided a proof-of-concept of “interferon-free treatment for chronic HCV GT-1” without treatment-related serious or severe adverse events [52]. New DAAs against HCV have since been developed, and these combinations without peginterferon can improve the SVR rates and shorten the treatment durations.

**Ledipasvir and sofosbuvir**

*Treatment-naive HCV GT-1 patients*

Ledipasvir is an HCV NS5A inhibitor with antiviral activity against HCV GT-1 [53]. Sofosbuvir is a nucleotide polymerase inhibitor with antiviral activity against HCV pan-genotypes [54]. After a phase 2 trial [54], an ION-1 study (n = 865) was conducted: a multicenter, randomized, open-label phase 3 trial for treatment-naive HCV GT-1 patients with 12 or 24 weeks of a fixed-dose combination of ledipasvir (90 mg) and sofosbuvir (400 mg), with or without ribavirin (Fig. 1a) [30]. A total of 67 % of the patients had HCV GT-1a infection, 70 % had the non-CC IL28B (rs12979860) genotype, and 16 % had cirrhosis. Of the 865 patients, only 3 patients had virological failure: one with HCV GT-1b had virological breakthrough, while the other two, one with HCV GT-1a and one with HCV GT-1b, had virological relapse. Concerning HCV NS5A-resistant associated variants (RAVs), the relapser with HCV GT-1a had the L31M variant, and both patients with HCV GT-1b had the Y93H variant at the time of virological failure. Two of the relapers also had HCV NS5A RAVs at baseline [30]. Treatment with 12 weeks of a fixed-dose combination of ledipasvir and sofosbuvir, with or without ribavirin, led to SVR 12 weeks after the end of treatment (SVR12) in 97 % (211/217) or 99 % (211/214) of the patients, respectively. Treatment with 24 weeks of a fixed-dose combination of ledipasvir and sofosbuvir, with or without ribavirin, led to SVR in 99 % (215/217) or 98 % (212/217) of the patients, respectively. Of the 33 patients who had a serious adverse event during treatment, 25 and 8 were in the 24- and 12-week regimens, respectively. Only 6 serious adverse events were observed, as follows: cellulitis, chest pain, gastroenteritis, hand fracture, non-cardiac chest pain and pneumonia, although fatigue, headache and nausea were the most common adverse events [54]. No patient in the 12-week group discontinued due to adverse events.

The ION-3 study (n = 647) consisted of a multicenter, randomized, open-label phase 3 trial for treatment-naive HCV GT-1 patients without cirrhosis with 8 weeks of the combination of ledipasvir (90 mg daily) and sofosbuvir (400 mg daily), with or without ribavirin, or with 12 weeks of the combination of ledipasvir and sofosbuvir without ribavirin (Fig. 1a) [36]. A total of 80 % of the patients had HCV GT-1a infection and 75 % of the total patients had non-CC IL28B (rs12979860) genotype. Treatment with 8 weeks of the combination of ledipasvir and sofosbuvir, with or without ribavirin, led to SVR12 in 93 % (201/216) or 94 % (202/215) patients, respectively. Treatment with 12 weeks of combination of ledipasvir and sofosbuvir without ribavirin led to SVR12 in 95 % (206/216). Of the 23 patients who had a relapse, 15 had HCV NS5A RAVs at the time of relapse and 8 did not. Of these 15 relapsed patients, 9 had RAVs at baseline and 6 did not. Adverse events were more associated with ribavirin [36].

A Japanese study [55] also showed that treatment with 12-week treatment with the combination of ledipasvir
(90 mg daily) and sofosbuvir (400 mg daily), with or without ribavirin, led to SVR12 in 96.4 % (80/83) or 100 % (83/83) of treatment-naive HCV GT-1 patients, respectively. At present, sofosbuvir cannot be used in patients with estimated glomerular filtration rate (eGFR) ≤30 mL/min per 1.73 m² or on hemodialysis.

**Treatment-experienced HCV GT-1 patients**

An ION-2 study (n = 440) has been carried out, consisting of a randomized, open-label phase 3 trial for HCV GT-1 patients who had not had SVR after treatment with peginterferon and ribavirin, with or without a protease inhibitor, with 12 or 24 weeks of a fixed-dose combination of ledipasvir (90 mg) and sofosbuvir (400 mg), with or without ribavirin (Fig. 1b) [29]. A total of 79 % of the patients had HCV GT-1a infection, 88 % had non-CC IL28B (rs12979860) genotype, and 20 % had cirrhosis. A total of 52 % of the patients had received prior treatment with a protease-inhibitor regimen. Of the total of 440 patients, 55.7 % (245) and 44.3 % (195) were previous treatment relapsers or patients with breakthrough and patients with no response, respectively. Treatment with 12 weeks of the fixed-dose combination of ledipasvir and sofosbuvir, with or without ribavirin, led to SVR12 in 96 % (107/111) or 94 % (102/109) patients, respectively. Treatment with 24 weeks of the fixed-dose combination of ledipasvir and sofosbuvir, with or without ribavirin, led to SVR in 99 % (110/111) or 99 % (108/109) patients, respectively. Of the 11 patients with relapse, 6 had HCV NS5A RAVs at baseline. All these 11 relapsers had HCV NS5A RAVs at the time of relapse. No patients discontinued treatment due to adverse events, of which fatigue, headache and nausea were the most common [29]. In patients with decompensated cirrhosis, ribavirin plays a role in the achievement of SVR [56].

The Japanese study [55] also showed that 12-week treatment with a combination of ledipasvir (90 mg daily) and sofosbuvir (400 mg daily), with or without ribavirin, led to SVR12 in 100 % (87/87) or 100 % (88/88) of treatment-experienced HCV GT-1 patients, respectively.

**Paritaprevir/ritonavir and ombitasvir**

A Japanese study (n = 73) comprised a randomized, open-label phase 2 trial for HCV GT-1b patients who had not had SVR after treatment with peginterferon and ribavirin with 12 or 24 weeks of the combination of NS3/4A inhibitor paritaprevir (ABT-450)/ritonavir (100/100 mg daily) and NS5A inhibitor ombitasvir (ABT-267) (25 mg daily) (Table 1) [57]. SVR 24 weeks after the end of treatment (SVR24) was high (88.9–100 %) regardless of the paritaprevir dose or treatment duration [57]. The most common adverse events were nasopharyngitis (29 %) and headache (14 %).

**Paritaprevir/ritonavir, ombitasvir and dasabuvir**

The Turquoise-II study (n = 380) consisted of a phase 3 trial for HCV GT-1 patients with Child-Pugh class A cirrhosis with 12 or 24 weeks of the combination of paritaprevir (ABT-450)/ritonavir (150/100 mg daily) and dasabuvir (250 mg, twice daily) and ribavirin (Table 1) [39]. Treatment for 12 or 24 weeks led to SVR12 in 91.8 % (191/208) or 95.9 % (165/172) patients, respectively. Drug discontinuations due to adverse events were uncommon (2.1 %).

**Table 1** Future perspectives of all-oral treatment for HCV genotype 1 in Asian-Pacific Region

| Companies (regimen) [references] | Drug targets | Ribavirin Duration of treatment (weeks) | SVR (%) |
|----------------------------------|--------------|---------------------------------------|---------|
| Gilead [29, 30, 36, 55]          | –            | Ledipasvir (90 mg daily)              | Sofosbuvir (400 mg daily) | ± | 12 | 93–100 |
| Abbvie (2D) [57]                 | Paritaprevir (150 mg daily)/ritonavir (100 mg daily) | Ombitasvir (25 mg daily) | – | ± | 12 | 89–100 |
| Abbvie (3D) [39, 58]             | Paritaprevir (150 mg daily)/ritonavir (100 mg daily) | Ombitasvir (25 mg daily) | Dasabuvir (500 mg daily) | ± | 12 | 90–100 |
| Bristol-Myers Squibb (2D) [60, 62] | Asunaprevir (200 mg daily) | Daclatasvir (60 mg daily) | – | – | 24 | 81–91 |
| Bristol-Myers Squibb (3D) [64]   | Asunaprevir (200 mg daily) | Daclatasvir (60 mg daily) | BMS-791325 (75 or 150 mg daily) | – | 12 or 24 | 89–94 |
| MSD [65]                         | Grazoprevir (MK-5172) (100 mg daily) | Elbasvir (MK-8742) (50 mg daily) | – | – | 12 | 90–100 |

Abbvie 2D regimen and Bristol-Myers 2D regimen are only available for HCV genotype-1b infection. Ritonavir is a booster.
The PEARL-III and PEARL-IV studies were conducted via a phase 3 trial for HCV GT-1a \((n = 419)\) or GT-1b \((n = 305)\) patients without cirrhosis, respectively, with a 12-week combination of paritaprevir (ABT-450)/ritonavir (150/100 mg daily), ombitasvir (ABT-267) (25 mg daily), and NS5B polymerase non-nucleoside inhibitor dasabuvir (ABT-333) (250 mg, twice daily), with or without ribavirin (Table 1) \([58]\). The 12-week treatment, with or without ribavirin, led to SVR12 in 97.0 % \((97/100)\) or 90.2 % \((185/205)\) of the HCV GT-1a patients, respectively. As for the HCV GT-1b patients, the 12-week treatment, with or without ribavirin, led to SVR12 in 99.5 % \((209/210)\) or 99.0 % \((207/209)\), respectively. Drug discontinuations due to adverse events were rare \((0.3\%)\). The most common adverse events were fatigue, headache and nausea.

SAPPHIRE-II studies were conducted by a phase 3 trial for treatment-experienced HCV GT-1 patients without cirrhosis, respectively, with 12-weeks of the combination of paritaprevir (ABT-450)/ritonavir (150/100 mg daily), ombitasvir (ABT-267) (25 mg daily), and dasabuvir (ABT-333) (250 mg, twice daily) with or without ribavirin, \((1000\ or\ 1200\ mg\ daily)\) (Table 1) \([40]\). The total SVR12 rate was 96.3 % \((286/297)\):95.3 % \((82/86)\) in those with prior relapse, 100 % \((65/65)\) with prior partial response, and 95.2 % \((139/146)\) with prior null response.

**Asunaprevir and daclatasvir**

After a phase 2 trial \([59]\), showing that the combination of asunaprevir and daclatasvir was effective only for HCV GT-1b, a Japanese \((n = 222)\) randomized, open-label phase 3 trial for HCV GT-1b patients \((135\ interferon-ineligible/intolerant\ and\ 87\ non-responder\ patients)\) with 24 weeks of the combination of NS3/4A inhibitor asunaprevir \((100\ mg,\ twice\ daily)\) and NS5A inhibitor daclatasvir \((60\ mg,\ once\ daily)\) was conducted (Table 1) \([60]\). SVR24 rates were 87.4 % \((118/135)\) or 80.5 % \((70/87)\) in interferon-ineligible/intolerant or nonresponder patients, respectively. The most common adverse events were nasopharyngitis, ALT and AST elevations, headache, diarrhea and pyrexia. It was also reported that drug-induced immunoallergic hepatitis occurred during the combination therapy with daclatasvir and asunaprevir \([61]\).

The HALLMARK-DUAL study presented a randomized, open-label phase 3 trial for HCV GT-1b patients \((205\ treatment-naïve, 205\ non-responders\ and\ 235\ ineligible\ and/or\ intolerant\ patients)\) with the 24-week combination of asunaprevir \((100\ mg,\ twice\ daily)\) and daclatasvir \((60\ mg,\ once\ daily)\) (Table 1) \([62]\). The SVR12 rate was 90 % \((182/205)\), 82 % \((168/205)\), or 82 % \((192/235)\) in treatment-naïve, non-responders, or ineligible and/or intolerant patients, respectively. They \([62]\) detected RAVs at NS5A-L31, NS5A-Y93 and NS3-D168, or a combination of two or more, in 75 of 596 patients. Of these 75 patients, only 29 \((39\%)\) achieved SVR12. Of note, these RAVs were associated with virological failure.

Asunaprevir and daclatasvir are mainly eliminated through liver. Therefore, this combination therapy for 24 weeks is not contraindicated in patients with severe renal impairment, including HCV GT-1 patients with hemodialysis and was highly effective and well tolerated \([63]\).

**Asunaprevir, daclatasvir and BMS-791325**

A joint USA/France study \((n = 64)\) performed a randomized, open-label phase 2a trial for treatment-naïve HCV GT-1 patients with 12 or 24 weeks of the combination of asunaprevir \((200\ mg,\ twice\ daily)\), daclatasvir \((60\ mg,\ once\ daily)\) and BMS-791325 \((75\ or\ 150\ mg,\ twice\ daily)\) (Table 1) \([64]\), SVR12 was achieved in 92 % \((61/64)\) of the patients.

**Grazoprevir (MK-5172) and elbasvir (MK-8742)**

Another USA/Europian countries study \((n = 253)\) consisted of a randomized, open-label phase 2 trial for HCV GT-1 patients with cirrhosis, with 12 or 18 weeks of the combination of HCV NS3/4A inhibitor grazoprevir (MK-5172) \((100\ mg\ daily)\) and NS5A inhibitor elbasvir (MK-8742) \((50\ mg\ daily)\), with or without ribavirin (Table 1) \([65]\). SVR12 was achieved in 90–97 % of the patients. Less than 1 % of grazoprevir and elbasvir are renally excreted. Therefore, this combination therapy for 12 weeks is not contraindicated in GT-1 patients with stage 4–5 chronic kidney disease and was highly effective and well tolerated \([66]\).

**Future perspectives of chronic HCV GT-1 patients in Asian countries**

There are differences between HCV GT-1a and GT-1b according to the several regimens of all-oral treatment for HCV GT-1. HCV subgenotyping and RAVs at HCV NS5A should be analyzed before the commencement of asunaprevir and daclatasvir combination treatment \([67]\). However, all-oral treatment can be available to cure almost all HCV GT-1 patients including “difficult-to-treat” patients (Fig. 2). Of course, further studies will be needed.

#2 Consensus Statements and Recommendation on all-oral treatment for HCV GT-1 infection

1. In treatment-naïve patients and previously treated-with peginterferon plus ribavirin patients with chronic HCV GT-1 infection, the following all-oral treatments apply:
1. For HCV GT-1 patients (treatment-naïve or -experienced with peginterferon plus ribavirin), treatment with daily ledipasvir (90 mg) plus sofosbuvir (400 mg) for 12 weeks is recommended. (A1)

2. For HCV GT-1b patients (treatment-naïve or -experienced with peginterferon plus ribavirin), other regimens such as paritaprevir/ritonavir, ombitasvir and dasabuvir for 12 weeks or grazoprevir (MK-5172) and elbasvir (MK-8742) for 12 weeks may also be applicable. (A1)

3. For HCV GT-1b patients (treatment-naïve or -experienced with peginterferon plus ribavirin) with renal impairment, other regimens such as grazoprevir (MK-5172) and elbasvir (MK-8742) for 12 weeks or asunaprevir and daclatasvir for 24 weeks may also be applicable. Before treatment with asunaprevir and daclatasvir, resistant associated variants (RAVs) with target regions should be examined. (A2)

2. In patients previously treated with NS3/4A protease inhibitor and peginterferon plus ribavirin with chronic HCV GT-1 infection, the following all-oral treatments apply:

1. For patients with minimal liver disease, treatment should be avoided, until data are available. (C1)

2. For patients with advanced liver diseases, RAVs with target regions should be examined before any treatments. (C1)

All-oral treatment for HCV GT-2 and GT-3 infection

HCV GT-2

In treatment-naïve patients with HCV GT-2 infection patients, the all-oral treatment regimen is the combination of sofosbuvir (400 mg per day) with weight-based RBV [1000 mg (<75 kg) to 1200 mg (≥75 kg)] for 12 weeks, which produces high SVR rates (Table 2). This regimen has been evaluated in 4 clinical trials (ELECTRON, FISSION, POSITRON, and VALENCE) [68–71]. The FISSION study randomized patients to receive daily peginterferon and RBV (800 mg) for 24 weeks or sofosbuvir plus daily weight-based RBV [1000 mg (<75 kg) to 1200 mg (≥75 kg)] for 12 weeks [70]. The SVR rate was higher (94 %) in patients who received sofosbuvir plus RBV than in those who received peginterferon and RBV (78 %; 52/67). Across all 3 trials [68, 70, 71], 201 (94 %) of the 214 patients with HCV GT-2 infection achieved SVR with sofosbuvir plus RBV. Among those who did not achieve SVR, sofosbuvir resistance-associated variants (RAVs) were not detected. Based on the results of these trials, both the FDA and EMA approved sofosbuvir and RBV for 12 weeks in all HCV GT-2 treatment-naïve

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Table 2 All-oral treatment for treatment-naïve HCV genotype 2 in the Asian-Pacific Region

| Companies (regimen) [references] | Drug targets | Ribavirin | Duration of treatment (weeks) | SVR (%) |
|----------------------------------|--------------|-----------|-------------------------------|--------|
|                                  | NS5A | NS5B |                  |          |
| Gilead Sciences [68–72]          |     |      | Sofosbuvir (400 mg daily)     | 12     | 88–100 |
| Bristol-Myers Squibb [73]         |     |      | Sofosbuvir (400 mg daily)     | 24     | 96     |
| Gilead Sciences [74]              |     |      | Sofosbuvir (400 mg daily)     | 24     | 96     |
| Gilead Sciences [75]              |     |      | Sofosbuvir (400 mg daily)     | 24     | 96     |
patients. An open-label phase III study in Japan confirmed these results [72]. A total of 153 patients with HCV GT-2 infection were enrolled and treated with sofosbuvir plus RBV for 12 weeks. Among them, 11 % had liver cirrhosis and 22 % were over 65 years. Overall, 148 patients (97 %) achieved SVR. Of the 90 treatment-naive patients, 88 (98 %) achieved SVR, including all the patients with cirrhosis. Of the 63 treatment-experienced patients, 60 (95 %) achieved SVR, including all cirrhosis patients except one (89 %). Thus, the overall SVR rate was 94 % in patients with cirrhosis and those over 65. Although no data were available to support an extension of therapy for treatment-naive patients with HCV GT-2 infection, longer treatment duration may improve SVR in treatment-experienced patients with cirrhosis (60 % after 12 weeks and 78 % after 16 weeks) (Table 3). Therefore, extending treatment from 12 to 16 weeks in HCV GT-2-infected patients with cirrhosis is recommended.

Daclatasvir (60 mg/day) with sofosbuvir (400 mg/day) for 24 weeks was associated with a high SVR rate (96 %) in treatment-naive patients with HCV GT-2 infection [73] (Table 2). Although RBV did not seem to be necessary to achieve SVR, the number of patients was too small to draw any firm conclusions. Ledipasvir (90 mg/day) with sofosbuvir (400 mg/day) for 12 weeks was also associated with high SVR12 rate (96 %) in HCV GT-2 infected patients including treatment-experienced and those with cirrhosis [74] (Table 2). Sofosbuvir in combination with GS-5816 was evaluated in treatment-naive patients with HCV GT-2 infection with high SVR rates [75] (Table 2).

**HCV GT-3**

With current anti-HCV agents, HCV GT-3 is the most difficult-to-cure genotype. The VALENCE study assessed the efficacy and safety of sofosbuvir (400 mg per day) plus weight-based RBV [1000 mg (<75 kg) to 1200 mg (≥75 kg)] for 24 weeks [68]. This trial included 250 treatment-naive (42 %) and -experienced (58 %) patients with HCV GT-3 infection. The overall SVR12 rate was 84 % and treatment-naive patients had a higher SVR than -experienced ones (93 vs. 77 %, respectively). These data suggest that higher SVR rates can be achieved with a 24-week sofosbuvir plus RBV therapy than those reported with 12- or 16-week therapy in the FISSION [70] (63 % after 12 weeks), POSITRON [71], (61 % after 12 weeks) and FUSION [71] (30 % after 12 weeks and 62 % after 16 weeks) trials (Tables 4, 5). The SVR rates were comparable between patients with and without cirrhosis (92 and 93 %, respectively). Recently, the phase III ALLY 3 study using daclatasvir (60 mg per day) plus sofosbuvir (400 mg per day) for 12 weeks [76]. The study included 101 treatment-naive patients with HCV genotype 3 infection and showed the SVR rate of 90 %. In treatment-naive patients without cirrhosis (Metavir F0-F3), 97 % achieved SVR, and in treatment-naive patients with cirrhosis (Metavir F4), 58 % achieved SVR.

**#3 Consensus statements and recommendation on all-oral treatment for HCV GT-2 and GT-3 infection**

In chronic HCV GT-2 or GT-3 infection, the following all-oral treatments apply:

1. **GT-2**
   
   1. For treatment-naive HCV GT-2 patients, daily sofosbuvir (400 mg) plus weight-based ribavirin (1000 or 1200 mg in patients <75 kg or ≥75 kg, respectively) for 12 weeks is recommended. (A1)
   2. Therapy can be prolonged to 16 or 24 weeks in cirrhotic patients previously treated with peginterferon plus ribavirin. (A1)
   3. Daily daclatasvir (60 mg) and sofosbuvir (400 mg) for 24 weeks is recommended for treatment-naive patients who cannot tolerate ribavirin. (B1)
   4. Daily ledipasvir (90 mg) and sofosbuvir (400 mg) for 12 weeks is recommended for treatment-naive patients who cannot tolerate ribavirin. (B1)
5. Daily velpatasvir (25 or 100 mg) and sofosbuvir (400 mg) for 12 weeks is recommended for treatment-naive patients who cannot tolerate ribavirin. (B1)

## Velpatasvir is not yet licensed but will soon be available.

2. GT-3

1. For treatment-naive HCV GT-3 patients, treatment with daily sofosbuvir (400 mg) plus weight-based ribavirin (1000 or 1200 mg in patients <75 kg or ≥75 kg, respectively) for 24 weeks is recommended. (A1)

Alternatively, daily daclatasvir (60 mg) and sofosbuvir (400 mg) for 12 weeks (no cirrhosis) (A1) or 24 weeks with or without weight-based RBV (cirrhosis) (B2) is recommended.

3. For patients previously treated with peginterferon plus ribavirin, daily daclatasvir (60 mg) and sofosbuvir (400 mg) for 12 weeks in those without cirrhosis (A1), and for 24 weeks with weight-based ribavirin for 24 weeks in those with cirrhosis (B2) are recommended.

### PAN-oral therapy for HCV GT-4, GT-5 and GT-6 infection

**Current clinical data with registered agents**

For HCV GT-4, GT-5 and GT-6, clinical trials related to the use of interferon-free pan-oral direct acting antiviral agents have been sponsored by AbbVie, Gilead Sciences and Merck (Table 6). Among Egyptians with HCV GT-4 residing in USA, SVR12 was achieved in 79 % (11/14) and 100 % (14/14) of treatment-naive patients treated for 12 weeks and 24 weeks with sofosbuvir plus weight-based RBV [1000 mg (<75 kg) to 1200 mg (≥75 kg)], respectively [77]. Similar results were achieved in a phase II study conducted in Egypt, with SVR12 rates of 84 % (21/25) and 92 % (22/24) in treatment-naive patients treated for 12 weeks and 24 weeks, respectively [78]. In an open-label study of HIV/HCV-co-infected patients (PHOTON-2), 31 treatment-naive patients with HCV GT-4 infection were treated with daily sofosbuvir plus weight-based RBV [1000 mg (<75 kg) to 1200 mg (≥75 kg)] for 24 weeks. In this study, 84 % of patients (26/31) achieved SVR12 [79]. Following the success of
ION studies, the SYNERGY trial, an open-label study evaluated a 12-week course of Harvoni (ledipasvir/sofosbuvir) in 21 HCV GT-4-infected patients, of whom 60% were treatment-naive and 43% had advanced fibrosis (Metavir F3 or F4). One patient took the first dose and then withdrew consent and SVR12 rate was 95% by intention-to-treat analysis and 100% by per-protocol analysis [80]. In PEARL-I, sponsored by AbbVie, treatment-naive patients with HCV GT-4 infection with or without cirrhosis received 12 weeks of the daily fixed-dose combination of Technivie (ombitasvir, paritaprevir and ritonavir) with or without weight-based RBV. SVR12 rates of 100% (42/42) was achieved in the group receiving RBV and 90.9% (40/44) in the group not receiving RBV. Adverse effects were generally mild, with headache, asthenia, fatigue, and nausea most commonly reported [81]. In vitro, both ledipasvir and daclactasvir are active against both HCV GT-5 and GT-6. However, clinical data related to the use of these NS5A DAAs are limited. The combination of sofosbuvir and ledipasvir, administered 12 weeks without ribavirin in treatment-naive and treatment-experienced patients infected with HCV GT-6 yielded SVR rate of 96% (24/25) [82].

**Emerging clinical study**

Future drug combinations will likely exist of two or more DAAs with the aim to achieve (1) pan-genotypic HCV activity, (2) little or no risk for resistance; (3) short duration (≤12 weeks) of treatment, and (4) SVR and definite cure of the disease.

To date, there are limited clinical data to support the use of the combination of 12 weeks of daily sofosbuvir (400 mg) plus simeprevir (150 mg) with or without weight-

### Table 6: HCV genotype (GT)-4, GT-5 and GT-6 SVR12 from different regimens

| Genotype | Company | Regimen | Non-cirrhotic (LSM < 14.7 kPa) SVR12 (%)/number of treated patients | Cirrhotic (LSM ≥ 14.7 kPa) |
|----------|---------|---------|---------------------------------------------------------------|-----------------------------|
| GT-4     | AbbVie  | Ombitasvir plus paritaprevir, 12 weeks | 91% (41/44, treatment-naive) [81] | No data available |
|          |         | Ombitasvir plus paritaprevir plus RBV, 12 weeks | 100% (42/42, treatment-naive or experienced) [81] | No data yet (study is on-going) |
|          | Gilead  | Ledipasvir and sofosbuvir (Harvoni™), 12 Weeks | 95% [19/20, treatment-naive (62%) or experienced (38%)] [80] | No data available |
|          |         | Sofosbuvir plus velpatasvir | 100% (116/116, Sofosbuvir plus velpatasvir 12 weeks) [ASTRAL-1 study] [86] | 83% (75/90, GT1-6, Sofosbuvir plus velpatasvir 12 weeks) [ASTRAL-4 study] [86] |
|          |         | | | 94% (82/87, GT1-6, Sofosbuvir plus velpatasvir plus ribavirin (RBV) 12 weeks) [ASTRAL-4 study] [86] |
|          |         | | | 86% (77/90, GT1-6, Sofosbuvir plus velpatasvir 24 weeks) [ASTRAL-4 study] [86] |
|          | MSD     | Grazoprevir plus elbasvir | 100% (18/18) [253] | 83% (75/90, GT1-6, Sofosbuvir plus Velpatasvir 12 weeks) [ASTRAL-4 study] [86] |
| GT-5     | Gilead  | Sofosbuvir plus velpatasvir | 97% (34/35, Sofosbuvir plus velpatasvir 12 weeks) [ASTRAL-1 study] [86] | 83% (75/90, GT1-6, Sofosbuvir plus Velpatasvir 12 weeks) [ASTRAL-4 study] [86] |
|          |         | | | 94% (82/87, GT1-6, Sofosbuvir plus Velpatasvir plus RBV 12 weeks) [ASTRAL-4 study] [86] |
|          |         | | | 86% (77/90, GT1-6, Sofosbuvir plus Velpatasvir 24 weeks) [ASTRAL-4 study] [86] |
| GT-6     | Gilead  | Ledipasvir and sofosbuvir (Harvoni™) | 96% [24/25, treatment-naive: 23 (92%); experienced:2 (8%); Cirrhosis: (8%)] [80] | 83% (75/90, GT1-6, Sofosbuvir plus velpatasvir 12 weeks) [ASTRAL-4 study] [86] |
|          |         | Sofosbuvir plus velpatasvir | 100% (41/41, Sofosbuvir plus velpatasvir 12 weeks) [ASTRAL-1 study] [86] | 94% (82/87, GT1-6, Sofosbuvir plus velpatasvir + RBV 12 weeks) [ASTRAL-4 study] [86] |
|          |         | | | 86% (77/90, GT1-6, Sofosbuvir plus velpatasvir 24 weeks) [ASTRAL-4 study] [86] |
Based RBV [1000 mg (<75 kg) to 1200 mg (≥75 kg)] in HCV GT-4-infected patients. In the open-label phase III RESTORE trial, 107 patients with HCV GT-4 infection, including 35 treatment-naive patients, were treated with simeprevir in combination with peginterferon and ribavirin. In treatment-naive patients, daily simeprevir (150 mg) for 12 weeks in combination with peginterferon and ribavirin for 24–48 weeks (by response-guided therapy) produced SVR in 83% (29 of 35) [83]. These results are comparable for 24–48 weeks (by response-guided therapy) produced SVR in 83% (29 of 35) [83]. These results are comparable to SVR rates observed with similar regimens in patients with HCV GT-1 infection, suggesting that efficacy of sofosbuvir plus simeprevir for HCV GT-4 infection may be roughly in line with the SVR rates of patients with HCV GT-1 infection shown in the COSMOS trial.

Recently, Merck has developed the combination of grazoprevir, NS3/4A protease inhibitor that has high potency in vitro against HCV GT-1, GT-2, GT-4, GT-5, and GT-6 and elbasvir, NS5A inhibitor active against GT-1, GT-2a, GT-3, GT-4, GT-5, and GT-6, even in the presence of RAVs associated with failure of other NS5A inhibitors, such as daclatasvir and ledipasvir [57, 73]. In the C-EDGE study with oral, once-daily, fixed-dose grazoprevir 100 mg/elbasvir 50 mg for 12 weeks being administered, 18 of 18 (100%) with HCV GT-4 and 8 (80%) of 10 with HCV GT-6, achieved SVR [84]. At the time of failure, both patients with HCV GT-6 with virological failure had NS3 resistant associated variants (RAVs), and 1 also had an NS5A RAV [84]. In the pipeline, Gilead Sciences is evaluating the safety and efficacy of an investigational all-oral pan-genotypic regimen containing the nucleotide analog polymerase inhibitor sofosbuvir (SOF) and the investigational NS5A inhibitor velpatasvir (GS-5816) for the treatment of hepatitis C infection across all genotypes. In a phase 2 study, among the 154 previously untreated hepatitis C patients without liver cirrhosis, 9% had HCV GT-4, a single individual had HCV GT-5 and 6% had HCV GT-6. About one-third had the favourable IL28B CC gene variant associated with interferon responsiveness. Participants in this open-label study were randomly assigned to receive 400 mg once-daily sofosbuvir plus either 25 mg or 100 mg once-daily GS-5816 for 12 weeks. SVR12 rates for GT-4 SVR12 rates were 100 and 86%, respectively, in the 25 mg and 100 mg dose groups. The single HCV GT-5 patient and all HCV GT-6 patients in both dose arms were cured [85]. So far, sofosbuvir plus velpatasvir (GS-5816) was well tolerated in over 800 patients with HCV infection evaluated. There was a low incidence of serious adverse effects and few discontinuations due to adverse events. The most frequently reported adverse events (>10%) were fatigue, headache, nausea and insomnia. The most frequently observed hematologic abnormality was hemoglobin decrease in the RBV-containing treatment groups. Further phase 3 studies, ASTRAL-1, -2, -3 and -4, have recently been completed, with all genotypes evaluated with sofosbuvir 400 mg/velpatasvir (VEL, GS-5816) 100 mg FDC tablet administered orally once daily, with and without RBV. Among the 1035 subjects in the ASTRAL-1, -2, and -3 studies, 21% had compensated cirrhosis and 28% had failed prior treatments. In ASTRAL-1, SOF/VEL for 12 weeks had similar adverse events compared with placebo and SVR12 was noted in 100% (116/116) GT-4, 97% (34/35) GT-5 and 100% (41/41) GT-6 [86].

Current availability of interferon-free and ribavirin-free regimens have been shown to be remarkably efficacious and well tolerated. However, the long duration of therapy (12–24 weeks) has raised concerns about adherence and cost [87]. Mirroring from the studies in HCV GT-1, one will anticipate the addition of a third potent direct-acting antiviral drug to the current dual therapy can further reduce the duration of treatment required to achieve sustained viral response in patients with chronic HCV GT-1 infection without cirrhosis. In the proof-of-concept study by Kohli et al., the use of three direct-acting drugs with different mechanisms of action in two therapeutic groups from a mono-infected urban population allowed for a shorter duration of treatment (6 weeks) and resulted in high cure rates and excellent tolerability [88]. Furthermore, viral kinetic modeling suggests that the three-drug regimen of sofosbuvir, ledipasvir, and GS-9451—targeting three different stages of the HCV lifecycle—resulted in enhanced HCV clearance compared with other regimens that target only two stages. Further larger studies will be warranted to confirm these findings and to understand the related specific host or viral factors to better refine our clinical management of these patients.

#4 Consensus statements and recommendation on all-oral treatment for HCV GT-4 infection

1. Patients without cirrhosis
   1. 12 weeks fixed-dose combination of sofosbuvir (400 mg) and ledipasvir (90 mg) in a single tablet administered once daily without ribavirin. (A1)
   2. 12 weeks fixed-dose combination of sofosbuvir (400 mg) and velpatasvir (VEL, GS-5816) 100 mg FDC tablet administered orally once daily. (A1)
   3. 12 weeks fixed-dose combination of omibitasvir (75 mg), paritaprevir (12.5 mg) and ritonavir (50 mg) in one single tablet (two tablets once daily with food) with daily weight-based ribavirin (1000 or 1200 mg in patients <75 kg or ≥75 kg, respectively. (B1)

2. Patients with compensated cirrhosis
   1. 12 weeks fixed-dose combination of sofosbuvir (400 mg) and velpatasvir (VEL, GS-5816) 100 mg FDC tablet administered orally once daily. (B1)
2. 24 weeks fixed-dose combination of sofosbuvir (400 mg) and ledipasvir (90 mg) in a single tablet administered once daily without ribavirin. (A1)
3. 24 weeks fixed-dose combination of ombitasvir (75 mg), paritaprevir (12.5 mg) and ritonavir (50 mg) in one single tablet (two tablets once daily with food) with daily weight-based ribavirin (1000 or 1200 mg in patients <75 kg or ≥75 kg, respectively). (B1)
4. 12 weeks combination of daily sofosbuvir (400 mg) and daily daclatasvir (60 mg), adding daily weight-based ribavirin (1000 or 1200 mg in patients <75 kg or ≥75 kg, respectively) and extending duration of treatment to 24 weeks without ribavirin. (B2)

## Velpatasvir is not licensed yet but will be available soon.

### Consensus statements and recommendation on all-oral treatment for HCV GT-5 and GT-6 infection

1. Patients without cirrhosis
   1. 12 weeks fixed-dose combination of sofosbuvir (400 mg) and velpatasvir (VEL, GS-5816) 100 mg FDC tablet administered orally once daily. (A1)
   2. 12 weeks fixed-dose combination of sofosbuvir (400 mg) and ledipasvir (90 mg) in a single tablet administered once daily without ribavirin. (B1)
   3. 12 weeks combination of daily sofosbuvir (400 mg) and daily daclatasvir (60 mg). (B1)
2. Patients with compensated cirrhosis
   1. 12 weeks fixed-dose combination of sofosbuvir (400 mg) and velpatasvir (VEL, GS-5816) 100 mg FDC tablet administered orally once daily. (B1)
   2. 12 weeks fixed-dose combination of sofosbuvir (400 mg) and ledipasvir (90 mg) in a single tablet administered once daily with weight-based ribavirin (1000 or 1200 mg in patients <75 kg or ≥75 kg, respectively). (B1)
   3. 24 weeks fixed-dose combination of sofosbuvir (400 mg) and ledipasvir (90 mg) in a single tablet administered once daily without ribavirin. (B1)
   4. 12 weeks combination of daily sofosbuvir (400 mg) and daily daclatasvir (60 mg), adding daily weight-based ribavirin (1000 or 1200 mg in patients <75 or ≥75 kg, respectively). (B1)
   5. 24 weeks combination of daily sofosbuvir (400 mg) and daily daclatasvir (60 mg) without ribavirin. (B1)

### Generic licensing of all-ors in Pakistan

Pakistan has the second highest burden of chronic HCV infection in the world after China. Prevalence of HCV infection is 5 %, with an estimated 8 million people infected with the virus [89]. There are, however, pockets of very high prevalence of up to 24 % of the population [90]. The major HCV genotype in Pakistan is GT-3 (90 % cases), followed by GT-1 (10 % cases). The IL28B genotype is favorable in most cases.

Both standard interferon and peginterferon have been used for treatment of HCV, in combination with ribavirin, with overall SVR rates of 65 % for standard interferon and around 75 % for peginterferon [91]. However, due to the large disease burden, Pakistan was the second country in the world after Egypt to receive sovaldi through the Gilead global access program at the heavily discounted price of 300 USD per month. The drug was formally registered in March 2015 and till now nearly 20,000 patients have been started on treatment. sovaldi is used either in combination with ribavirin for 24 weeks or with ribavirin and peginterferon for 12 weeks. Initial results suggest SVR rates that are similar to those reported for HCV GT-3, of around 85 %.

A sizeable population of difficult to treat HCV GT-3 patients with prior treatment exposure as well as cirrhosis is emerging, particularly in the tertiary care centers. For these patients, additional DAAs are urgently needed.

### New DAA in Egypt: the licensing scenario

It seems that, finally, the nightmare of HCV in Egypt will end, as by April 2015, 2 DAA brands, Sofosbuvir, and Semiprevir, as well as 2 generics for Sofosbuvir have been registered and are available in Egypt. Early in 2014, the Egyptian government represented by its Ministry of Health (MOH) and the National Committee for Control of Viral Hepatitis (NCCVH), successfully signed a memorandum of understanding (MoU) with Gilead according to which Gilead accepted to supply its DAA brand, Sofosbuvir, at only 1 % of its price in the USA. This price was close to 300 USD/28 cap box. By March 2015, the Ministry of Health signed a similar MoU with Janssen to supply its brand Simeprevir at a price of 250 USD/box. Simeprevir has recently been added to the national treatment program.

- Other approved treatment options for HCV GT-4 as, LEDIPASVIR/SOFOSBUVIR “Harvoni”, Daklatasvir and Viekirax [92] will soon be available.
- The National Committee for Control of Viral Hepatitis (NCCVH), finalized the regulations for using these new brands. This current phase of the national program to
treat HCV in Egypt was preceded by a state announcement that has been spread in all Egyptian media, starting from September 2014, requesting all Egyptian patients known to have HCV with advanced liver disease to register on the website, www.nccvh.org.e.g.

- Electronic replies to patients were used to fix individual appointment dates and referred to a center where each patient has to be assessed for eligibility to use the new treatment protocols.
- By September 18 2014, 180,000 Egyptians have registered on the website, with 10 % of those registered living outside Egypt. By March 2015, 850,000 patients in total were already registered and processed through the website.
- Thirty-four governmental centers are currently participating in the project. The number is increasing monthly to cover all governorates.
- The first doses were distributed starting October 20 2014 and 25,000 patients had been treated by March 2015. It is planned that more than 250,000 patients yearly would receive the new treatment protocols. The same rules will be applied for all patients regardless of the source of payment, and there will be no place for patients’ preferences in deciding the treatment regimen.
- Treatment in its first phase included only cases with F2, F3, F4, or compensated cirrhosis, excluding cases with decompensation or with HCC patients except after successful radical curative intervention (4 months after resection or successful local ablation) evident by triphasic CT.
- The presence of large risky esophageal varices, required prophylactic management before being treatment-eligible.
- Age limits for treatment eligibility fixed to be above 18 years and below 70 years for all patients while body mass index (BMI) will be accepted up to 35.
- For special population groups; priority for treatment will be offered for post-liver transplantation, post-kidney transplantation patients and combined HCV/ HBV infection regardless of the fibrosis stage. Other groups such as pediatric age groups and kidney disease patients will be reviewed following the availability of sufficient data. Patients with documented extra-hepatic manifestations will be prioritized for treatment according to the same guidelines.
- Treatment-experienced patients should only start 6 months after cessation of the previous therapy.
- No differentiation in treatment priority will be established based on the previous treatment experience.
- Sofosbuvir (nucleotide polymerase inhibitor) was introduced in the national treatment program through 2 treatment options:
  1. Triple Regimen including peginterferon (α2A and α2B): Sofosbuvir plus peginterferon and ribavirin for 12 weeks
  2. Dual Regimen: Sofosbuvir plus RBV for 24 weeks

Treatment was designed for those with Fibrosis score F2, F3, F4 and compensated cirrhotics. Recent published reports support this recommended protocol [93].

- The increasing number of reports that treatment with the new DAAs are well tolerated and efficacious in patients with decompensated cirrhosis and, more importantly, markers of hepatic and synthetic function improved during the short-term follow-up [94, 95], will be reviewed. The inclusion of these cases in the national program will be considered during the next phases of the national treatment program.
- In the new era of the DAAs, in addition to increased treatment efficacy with SVR more than 90 %, by increasing cases detection and reducing new infections, according to a mathematical modeling, the government strategy is to achieve <2 % prevalence by 2025 and >90 % drop in prevalence by 2030 (near-total HCV elimination) [96].

Results of the national program The initial report of the national program was published in June 2015. The report was an interim analysis of efficacy in both treatment groups (dual and triple therapy). The study included 458 patients received dual regimen and 349 received triple regimen. Both groups were evaluated for treatment results at week 12 which is considered the end of treatment for the triple regimen and midpoint of treatment in the dual regimen.

Results

Detailed results of the analysis are shown in Table 7. The overall treatment response at week 12 in both groups was 98.8 % (798/807), ETR in the triple regimen group was 99.1 % (346/349) and in the dual regimen group was 98.3 % (452/458) [97].

#6 Consensus statements and recommendation on generic licensing of all-orals in Egypt

1. In the new era of the DAAs, with the high clearance rate of the HCV and minimal side effects, the dream of global eradication of the virus seems feasible. (A1)
2. The major global obstacle in using these new DDAs is the outstanding very high prices of these drugs exceeding the capabilities of health care systems even in rich developed countries, and consequently the scenario will be harder both in poor as well as in
developing countries with low national incomes and financial resources. (B1)

3. In developing countries like Egypt having a high prevalence of HCV infection, in parallel with relatively low health care budgets and resources, the only feasible strategy to eradicate the virus is to establish deals with these new DAAs promising drug-producing companies to supply the drugs at a cheap affordable price for both the government and individuals. (B1)

### Generic licensing of all-oral in Indonesia

In Indonesia, chronic hepatitis C infection is dominated by GT-1b. Therefore, the standard of care (SOC) is the combination of peginterferon-alpha 2a or 2b with ribavirin [98]. Based on a multicenter prospective open-label trial in Indonesia using conventional interferon therapy with ribavirin, it was shown that SVR was 78.3% in patients with HCV GT-1 and GT-4, and 100% in non-GT-1 and GT-4 [99], whereas in a recent publication based on IL-28B distribution, the 73.5% SVR was achieved in chronic HCVGT-1 patients with 75.8% SVR in CC genotype using peginterferon-alpha 2a and ribavirin therapy [100].

### DAA in Indonesia

The only available DAA in Indonesia at this moment is boceprevir which should be used together with peginterferon and ribavirin. Sofosbuvir will soon be available in Indonesia with generic licensing. Since most Indonesian patients are dominated by HCV GT-1, the use of peginterferon will still be the cornerstone of standard therapy. However, the potential for an upcoming new combination of DAA agents with the possibility of generic licensing will give a new standard of care in Indonesia.

| Table 7 Patients characteristics according to treatment response | Treatment responders (n = 798) | Treatment non-responders (n = 9) | p value |
| --- | --- | --- | --- |
| Gender | | | |
| Male | 548 | 5 | 0.47 |
| Female | 250 | 4 | |
| Age (mean ± SD) | 53 ± 8 | 57 ± 6 | 0.16 |
| Treatment status | | | |
| Naïve | 540 | 3 | 0.07* |
| Experienced | 258 | 6 | |
| Baseline laboratory values (mean ± SD) | | | |
| Hemoglobin | 13.4 ± 1.7 | 13.3 ± 1.6 | 0.82 |
| WBCs | 5.8 ± 4 | 5.1 ± 2 | 0.66 |
| Platelets | 129 ± 66 | 133 ± 92 | 0.88 |
| Albumin | 3.8 ± 0.7 | 3.8 ± 0.8 | 0.87 |
| Viremia (log 10) | 5.5 ± 1 | 5.5 ± 0.7 | 0.98 |
| Fibrosis assessment | | | |
| FIB-4 median (IQR) | 4.34 (4.2) | 6.6 (4.5) | 0.23 |
| Fibrosis grading (n = 257) | 2/171/84 | 1/2/0 | <0.01* |
| Child score | | | |
| Non cirrhotic/A/B | 535/226/37 | 3/4/2 | 0.08/0.46 |
| Regimen type | | | |
| Dual | 452 | 6 | 0.79 |
| Triple (Roch/MSD) | 346 (198/148) | 3/2/1 | 0.36 |
| Abdominal U/S | | | |
| Liver cirrhosis | 482 | 5 | 0.96 |
| Ascites | 17 | 0 | 0.47 |
| Sponsorship | | | |
| Governmental | 715 | 5 | |

* Significant p < 0.05
#7 Consensus statements and recommendation on generic licensing of all-oral in Indonesia

1. In Indonesia, chronic hepatitis C infection can be treated with the standard therapy which consists of peginterferon and ribavirin. (B1)

2. Boceprevir might be used in addition to standard therapy if only there is no or partial response to the standard therapy. (B2)

3. Sofosbuvir might be used in combination with ribavirin in HCV GT-2 and GT-3 patients. (B1)

4. An upcoming new combination of DAA agents might be used or replace the use of peginterferon as a therapy for chronic HCV infection in all genotypes. (A1)

HCV infection in patients with hepatic decompensation, liver transplant candidates and recipients

HCV reinfection after liver transplant is universal and inevitable, and can be the key factor associated with premature graft loss. Treatment of HCV infection is crucial, as successful treatment has the potential to alter the outcomes of decompensated liver disease, liver transplant candidates and in patients with HCV recurrence after transplantation. Previously, interferon-based therapies have not been used in decompensated liver disease due to the high risk of infection and further hepatic decompensation [101]. The outcomes of interferon-based therapy after liver transplantation have been relatively poor compared to interferon-based therapies in the non-transplant situation [102]. The advent of interferon-free therapies enables successful treatment in liver transplant candidates, patients with hepatic decompensation (proceeding or not to liver transplantation) and patients in the post-transplant situation.

a. The liver transplant candidate with HCC and portal hypertension but no liver decompensation. There has only been one published study of interferon-free antiviral therapy in this situation. Curry et al. reported on using sofosbuvir and ribavirin in 61 patients awaiting liver transplantation for HCC as the primary indication [38]. The majority of patients had HCV GT-1. Patients had CPS < 7. Nine patients relapsed, did not respond or had viral breakthrough whilst 47 underwent liver transplantation. Of these, 3 had HCV detected at transplant and all 3 relapsed post-transplant. Forty-three patients were PCR negative at transplant and 10 relapsed post-transplant. The authors claimed that 24/25 patients who had the virus not detected for >4 weeks and underwent transplant had no recurrence. Thus, it seems that 30/61 patients (49 %) on an intention-to-treat basis had no HCV recurrence in the allograft. This paper has established the proof of principle that prolonged viral suppression for >4 weeks pre-transplant with sofosbuvir-based therapy can prevent reinfection of the liver allograft in at least 50 % of patients.

b. Decompensated Cirrhosis. There are now several reports (all still in abstract form) using sofosbuvir-based interferon-free antiviral therapy (AVT) in this situation [103–107]. These studies mainly on HCV GT-1 report SVR rates for sofosbuvir and ledipasvir of 80–90 % in Child-Pugh Score C patients but this may be as low as 70 % [103–107]. In these studies, an improvement of >2 in MELD scores was seen in 40–50 % of patients but worsened in about 5–10 %. In an open label study of simeprevir and sofosbuvir in HCV GT-1 patients, a similar SVR (approximately 80 %) was seen. However, simeprevir drug levels may have a significantly increased area under the curve in decompensated diseases and is generally not recommended. In one study of HCV GT-3 patients, sofosbuvir plus daclatasvir resulted in a SVR of 70 versus 60 % versus sofosbuvir plus ledipasvir [105]. It remains unclear whether ribavirin results in an increased SVR in these patients using such regimes. It seems that HCV GT-2 patients may have similar response rates using sofosbuvir and ribavirin.

c. Treating post-liver transplant HCV recurrence. A study of treatment of mild hepatitis C allograft infection with an AbbVie 3D regimen (ritonavir boosted paritaprevir, ombitasvir and dabuvir) in HCV GT-1 infection led to a 90 % SVR rate [37]. Similar SVR rates have been seen in sofosbuvir and ledipasvir therapy including some patients with more advanced disease [108–112]. Studies with sofosbuvir-based therapies in the post-transplant population in very severe liver disease such as cholestatic hepatitis C have led to 60–70 % SVR rates, but in such studies there was a mortality of approximately 13 % due to advanced disease [113, 114]. Ribavirin should be added on the sofosbuvir-based therapies and the duration of the regimens should be 24 weeks in patients with decompensated liver disease or in HCV GT-3 patients of post-transplant patients [56].

In conclusion there seem to be limitations to the outcomes from antiviral therapy particularly in decompensated liver disease. Current data suggest that interferon-free AVT will prevent HCV allograft reinfection in a significant number of patients. However, many patients with severe portal hypertension and ascites may continue to have hepatic decompensation and severe portal hypertension despite achieving SVR, and thus will still require liver transplantation. It is not
clear what parameters exist that are determining successful reversible portal hypertension or not. Furthermore, sofosbuvir-based therapies have not been studied extensively in patients with renal dysfunction which is not an uncommon problem in advanced liver disease. (Sofosbuvir is currently contraindicated in patients with creatinine clearance rates of less than 30 mL/min.) Thus, sofosbuvir-based regimes may not be a viable option in patients with very high MELD scores who have overt or incipient renal failure.

#8 Consensus statements and recommendation on decompensated liver disease

1. All the patients with decompensated liver disease can be treated with interferon-free-based therapies. Emerging data suggest that sofosbuvir-based treatment plus or minus ledipasvir (HCV GT-1) or daclatasvir (HCV GT-3) may be effective in this setting. Patients should be treated for a period of 12–24 weeks. Anticipated SVR rates at between 70 and 90 %. (B1)

#9 Consensus statements and recommendation on awaiting liver transplant

1. All patients awaiting liver transplantation can be treated as above with interferon-free antiviral therapy (sofosbuvir-based treatment plus or minus ledipasvir (HCV GT-1) or daclatasvir (HCV GT-3) for a minimum of 3 months before liver transplantation to prevent allograft recurrence. If viral clearance persists for >4 weeks even on-treatment, allograft re-infection is very likely to be prevented.(A1). It remains unclear how many patients improve MELD and CTP scores that allow removal from waiting lists. Thus, interferon-free therapy may best be introduced post-transplant rather than pre-transplant in patients with very high MELD scores. (C1)

#10 Consensus statements and recommendation on following liver transplant

1. Following liver transplantation recurrent HCV infection should be treated with interferon-free based therapies for 3 months. It is reasonable to commence therapy between 1 and 3 months post-transplant. These regimens may include sofosbuvir and ledipasvir or daclatasvir and the Abbvie 3D regime (ritonavir-boosted paritaprevir, ombitasvir and dasabuvir) for mild disease. If the latter is used, dose adjustments (significant reductions) of cyclosporin and tacrolimus are required. For more severe disease, evidence exists for sofosbuvir-based therapy in the combination of ledipasvir (GT-1) or daclatasvir (GT-3). (B1)

2. With the introduction of universal antiviral treatment post-transplant, the issues of immunosuppression should not be different in this population than in other liver transplant populations, although immunosuppressive drug doses may need adjustment depending on the regime used. (C2)

SVR for renal failure and co-infection with HBV/HIV

SVR for HCV infection in patients with renal failure

Comorbidities of HCV infection and chronic kidney disease (CKD) might present in two ways: HCV infection during maintenance dialysis and HCV-associated kidney disease. These disorders can occur both in native kidneys and in renal allografts. Therefore, all patients with kidney diseases should be evaluated for possible underlying HCV infection. It is suggested that HCV-infected patients be tested at least annually for proteinuria and hematuria.

The prevalence of HCV increases with the time that CKD patients are on dialysis which suggests that nosocomial transmission may be the route of infection in previously uninfected dialysis patients [115]. Sensitive quantitative RT-PCR tests for HCV should be administered to hemodialysis patients with unexplained abnormal aminotransferase(s) levels. Patients with end-stage renal disease (ESRD) have lower serum ALT levels than the general population; thus, some studies suggest that the optimized cut-off ALT level be approximately 0.4–0.45 times the upper limit of normal for HCV-infected patients [1, 2, 116, 117]. Cirrhosis, Asian race and history of alcohol abuse are associated with the highest risks for the development of HCC among dialysis patients with HCV infection [118]. Even after adjusting for concurrent co-morbidities, HCV infection is associated with higher risk of liver-related mortality in these subgroups of HCV-infected patients. Furthermore, HCV infection decreases the health-related quality of life in dialysis patients.

Currently, DAA-based therapies offer the best outcomes in patients with chronic kidney disease (CKD) who have mild to moderate renal impairment, i.e., creatinine clearance (CCr) from 30 mL/min to 80 mL/min. The standard dose of sofosbuvir, fixed-dose combination of sofosbuvir/ledipasvir, simeprevir, fixed-dose combination of paritaprevir/ritonavir/ombitasvir plus dasabuvir have demonstrated good efficacy in CKD. Simeprevir was also used safely at standard dose for patients with severe renal impairment, i.e., CCr < 30 mL/min.
In many countries in Asia, the access to new DAAs is limited and variable, and thus the combination peginterferon-alpha and ribavirin may still be the standard of care available for HCV infection in many CKD patients with normal, mild, moderate, or severe decrease in the glomerular filtration rate (GFR), and even in those with kidney failure. For interferon-based treatment regimens, please refer to the 2012 edition of this clinical practice guidelines for further verification [1].

#11 Consensus statements and recommendation on HCV and CKD

1. HCV-infected patients should be screened for proteinuria and hematuria at least annually to detect early HCV-associated kidney disease. (B2)
2. Maintenance hemodialysis (CKD stage 5D) confers a significant risk of nosocomial infection. Therefore, standard precautions for prevention of nosocomial infections must be rigorously observed. (A1)
3. Patients on hemodialysis should be screened with serological tests and RT-PCR at first hemodialysis or when transferring from another hemodialysis unit. (A1)
4. Maintenance hemodialysis patients and kidney transplant candidates should be tested for anti-HCV antibodies every 6–12 months, and RT-PCR should be performed for patients with unexplained elevated aminotransferase(s). (B2)
5. In dialysis patients with chronic HCV infection, liver biopsy is not mandatory, but may be recommended when the results would influence treatment decisions and when progression of the liver disease needs to be assessed. (B2)
6. Peginterferon with or without ribavirin is recommended for HCV-infected CKD patients with normal or mild decrease in GFR [creatinine clearance (CCr) ≥ 60 mL/min]. (B2)
7. The standard dose of sofosbuvir, fixed-dose combination of sofosbuvir/ledipasvir, simeprevir, fixed-dose combination of paritaprevir/ritonavir/ombitasvir plus dasabuvir may be used in the treatment or re-treatment of patients with mild to moderate renal impairment, i.e., CCr 30–80 mL/min. The use of these agents for patients with severe renal impairment, i.e., CCr < 30 mL/min or with ESRD has not been tested. (B2)
8. Regular serological screening of dialysis staff is highly recommended. (B2)

SVR for HCV co-infection with HIV

Hepatitis C has a limited impact on HIV disease progression. Conversely, HIV alters the natural history of hepatitis C in several important areas [2]. A rapid progression of liver fibrosis and increased mortality after decompensation has been observed in HCV/HIV co-infected patients [119–121]. HCV/HIV co-infected individuals should be offered treatment or re-treatment like any other individual without HIV infection, regardless of their stage of fibrosis at diagnosis [122–125].

Second generation DAA-based therapies have demonstrated high efficacy and safety in treatment-naïve, treatment experienced and cirrhotic HCV patients co-infected with HIV. Please refer to the “PAN-oral therapy for HCV GT-4, GT-5 and GT-6 infection” section for this guideline’s recommended DAA-based regimens for HCV infection. However, caution should be exercised when using such agents due to known drug–drug interactions with antiretroviral agents. A close collaboration with HIV specialist is recommended when treating HCV/HIV co-infected individuals.

While current western guidelines do not favor peginterferon/ribavirin-based therapies any more, in Asia, many of the second-generation DDAs are not yet available and thus combination peginterferon and ribavirin is still the standard of care. Data from previous studies have indicated that SVR achieved with this regimen reduces liver-related complications and mortality in HCV/HIV co-infected patients [126, 127]. Predictors of treatment response with peginterferon/ribavirin therapy are factors largely related to HCV: rapid virologic response (RVR), HCV genotype, HCV viral load, IL28B gene variation, and liver disease stage, however, the SVR rates in HIV/HCV co-infected patients are 15–20 % lower than those in patients with HCV mono-infection. Likewise, rates of hepatic decompensation during peginterferon/ribavirin treatment are considerably higher in co-infected patients than in HCV mono-infected patients, especially among cirrhotics [2, 128–133]. For interferon-based treatment regimens, please refer to the previous report for further verification [1].

#12 Consensus statements and recommendation on HCV and HIV co-infection

1. Routine screening for HIV is recommended in patients with hepatitis C following exposure risk assessment and pretest counselling. (A1)
2. HIV/HCV co-infected patients with advanced HIV disease (CD4 count < 100/µL) should receive highly active anti-retroviral therapy (HAART). HCV treatment should be delayed until immune function is improved, preferably until a CD4 count >200/µL is achieved. (A1)
3. Antiretroviral therapy-naïve HIV/HCV co-infected patients with a CD4 count of 100–350/µL should commence HAART prior to HCV treatment. (A1)
4. HIV/HCV co-infected patients with a CD4 count >350/µL should be considered for HCV treatment and do not require HAART. (A1)

5. HCV/HIV co-infected individuals should be offered HCV treatment or re-treatment like any other individual without HIV infection, regardless of the stage of fibrosis at diagnosis. (B1)

6. In regions in Asia where DAAs are not yet available/accessible, peginterferon and ribavirin combination therapy for 48 weeks is still the recommended HCV treatment; weight-based ribavirin dosing should be considered for HCV genotype 1 patients. (A1)

7. Drug–drug interactions of ribavirin- and DAA-based HCV medications with antiretroviral agents may induce adverse reactions and unwanted toxicities. (A1)

8. Ledipasvir should not be used in patients with CCr < 60 mL/min when co-administered with tenofovir-containing regimens. (B2)

9. Baseline and regular on-treatment evaluation of renal function is recommended when medications known to increase tenofovir levels are used. (B2)

10. Tipranavir, cobicistat and elvitegravir (pending more current data) should not be used with fixed-dose sofosbuvir/ledipasvir. (B2) Tipranavir should not be used with sofosbuvir. (B2)

11. Fixed-dose combination paritaprevir/ritonavir/ombitasvir plus dasabuvir should not be co-administered with ritonavir-boosted lopinavir, efavirenz, rilpivirine, and darunavir. (B2)

12. When used for boosting HIV protease inhibitor (PI), adjustments in the dose of ritonavir may be necessary if used with paritaprevir/ritonavir/ombitasvir plus dasabuvir. (B2)

13. Simeprevir should not be used with efavirenz, etravirine, nevirapine, cobicistat, or any HIV protease inhibitors. (B2)

14. Regimens containing telaprevir or boceprevir and monotherapy with peginterferon, ribavirin or a DAA are not recommended. (A1)

SVR for HCV co-infection with HBV

If HCV is determined to be replicating and is the dominant driver of liver inflammation in HCV/HBV co-infection, co-infected patients should be treated with similar regimens like those with HCV mono-infection. When serum HBV DNA levels are elevated at any time before, during and after HCV treatment, nucleos(t)ide analogues may be added to current HCV therapy. Peginterferon may be an option. Please refer to the recommendations in the 2012 guidelines [1]. Similar SVR rates are achieved with HCV genotype-guided peginterferon/ribavirin therapy in HCV mono-infected versus HCV/HBV co-infected patients [134]. HBsAg seroclearance is also observed in co-infected patients treated with peginterferon/ribavirin [135–137]. Some of the newly-approved DAAs for HCV treatment have demonstrable drug–drug interactions with nucleos(t)ide analogues which may limit or preclude their combined use in the HCV/HBV co-infected patients. In patients who achieved SVR, long-term follow-up and monitoring for relapse of HBV infection is recommended [124].

#13 Consensus statements and recommendation on HCV and HBV co-infection

1. Routine screening for HBsAg is recommended in patients with chronic HCV infection, especially in IVDUs or other high-risk populations. (A1)

2. Routine testing for serum HBV DNA is not recommended in HBsAg-negative patients with chronic HCV infection. (B1)

3. HCC screening tests, including liver ultrasonography and tests for AFP levels, are also required for co-infected patients. (B1)

4. HBV and HCV co-infected patients may be selected for antiviral treatment by the same criteria as those used for patients with mono-infection. (B2)

5. Determine which virus is dominant in patients with dual infection before commencing treatment. When HCV is the dominant replicating virus, treatment regimens are similar to those with HCV mono-infection. (B1)

6. At any time the serum HBV DNA is elevated, appropriate anti-HBV treatment must be started. (A1)

7. Baseline and regular on-treatment evaluation of renal function is recommended when tenofovir is used concurrently with DAAs and other anti-HCV medications known to increase tenofovir levels. (A1)

8. Co-administration of ledipasvir with tenofovir is not recommended in patients with CCr < 60 mL/min. (B2)

9. Simeprevir and fixed-dose paritaprevir/ritonavir/ombitasvir plus dasabuvir do not have clinically significant interactions with lamivudine and tenofovir. (B2)

10. HBV vaccination should be offered for hepatitis C patients who are HBsAg negative. (A1)

In conclusion, we have reviewed the recent advances of interferon-free therapies for the patients with chronic HCV infection in Asian-Pacific countries. A summary is shown in Table 8 and Supplementary Table 2. In the near future, HCV treatment will be further evolved to achieve much higher SVR rates and much shorter treatment duration.
| Grading | TN/TE-PR | TE-NS3/4A inhibitor | TE-NS5A inhibitor |
|---------|----------|---------------------|------------------|
|         | Non-cirrhosis/cirrhosis | Non-cirrhosis/cirrhosis | Non-cirrhosis/cirrhosis |

**Table 8** Interferon-free treatment for patients infected with HCV various genotypes (GTs)

### (A) GT-1

| TN/TE-PR | Non-cirrhosis/cirrhosis | Non-cirrhosis/cirrhosis | Non-cirrhosis/cirrhosis |
|----------|--------------------------|-------------------------|-------------------------|
| A1       | SOF/LDV for 12 weeks     | SOF/LDV for 12 weeks    |                        |
|          | PrOD for 12 weeks        |                         |                        |
| A2       | GZR/EBR for 12 weeks     |                         |                        |

### (B) GT-2

| TN/TE-PR | Non-cirrhosis/cirrhosis | Non-cirrhosis/cirrhosis | Non-cirrhosis/cirrhosis |
|----------|--------------------------|-------------------------|-------------------------|
| A1       | SOF/RBV for 12 weeks     | SOF/RBV for 12 weeks    | SOF/RBV for 12 weeks    |
|          | SOF/DCV for 24 weeks     | SOF/DCV for 24 weeks    | SOF/DCV for 24 weeks    |
| B1 (for RBV-intolerant) | SOF/LDV for 12 weeks | SOF/LDV for 12 weeks | SOF/LDV for 12 weeks |
|          | SOF/VEL for 12 weeks     | SOF/VEL for 12 weeks    | SOF/VEL for 12 weeks    |

### (C) GT-3

| TN/TE-PR | Non-cirrhosis/cirrhosis | Non-cirrhosis/cirrhosis | Non-cirrhosis/cirrhosis |
|----------|--------------------------|-------------------------|-------------------------|
| A1       | SOF/RBV for 24 weeks     | SOF/RBV for 24 weeks    | SOF/RBV for 24 weeks    |
| A2       | SOF/DCV for 12 weeks     | SOF/DCV ± RBV for 24 weeks | SOF/DCV for 12 weeks |
| B2       | SOF/RBV for 16 weeks     | SOF/VEL/RBV for 24 weeks | SOF/VEL/RBV for 24 weeks |

### (D) GT-4

| TN/TE-PR | Non-cirrhosis/cirrhosis | Non-cirrhosis/cirrhosis | Non-cirrhosis/cirrhosis |
|----------|--------------------------|-------------------------|-------------------------|
| A1       | SOF/LDV for 12 weeks     |                        | SOF/LDV for 24 weeks    |
| B1       | PrO/RBV for 12 weeks     | PrO/RBV for 24 weeks    | SOF/VEL for 24 weeks    |
| B2       | SOF/DCV/RBV for 12 weeks | SOF/DCV for 24 weeks    |                         |

### (E) GT-5/GT-6

| TN/TE-PR | Non-cirrhosis/cirrhosis | Non-cirrhosis/cirrhosis | Non-cirrhosis/cirrhosis |
|----------|--------------------------|-------------------------|-------------------------|
| A1       | SOF/VEL for 12 weeks     |                        | SOF/VEL for 12 weeks    |
| B1       | SOF/LDV for 12 weeks     | SOF/DCV for 12 or 24 weeks | SOF/VEL for 12 weeks |

**TN** treatment-naïve; **TE** treatment-experienced; **PR** peginterferon plus ribavirin; **SOF/LDV** sofosbuvir/ledipasvir; **PrOD** paritaprevir/ritonavir/ombitasvir/dasabuvir; **GZR/EBR** grazoprevir/elbasvir; **ASN/DCV** asunaprevir/daclatasvir; **RAV** resistant associated variant

*Vel, velpatasvir (will soon be available)*
Compliance with ethical statements

Conflict of interest All procedures followed were in accordance with the ethical standards of the responsible committee on human experimentation (institutional and national) and with the Declaration of Helsinki 1975, as revised in 2008 (5). Informed consent was obtained from all patients for being included in the study. This article does not contain any studies with animal subjects. Alaeeddin Ibrahim, Cosmas Rinaldi Adithya Lesmana, Mamun Al-Mahtab, Alaeeddin Ibrahim, George K. K.Lau, Barjesh C. Sharma, Jose Sollano, Manoj Kumar, Ankur Jindal, A. Kadir Dokmei, Geoffrey W. McCaughan, Darrell HG Crawford, Jafri Wasim, and Shiv Kumar Sarin declare that they have no conflict of interest. Masao Omata received fees for being a speaker, consultant, and advisory board member for Bayer Co, Boehringer Ingelheim, Bristol-Myers Squibb, Otsuka, Astellas, Gilead Sciences, Chugai, Mitsubishi Tanabe, Kyorin, Merck Sharp & Dohme, Dainippon Sumitomo, Vertex Pharmaceuticals, Takeda, and Zeria. Tatsuo Kanda received lecture fees from Chugai Pharmaceutical, MSD, Tanabe-Mitsubishi, Daiichi-Sankyo, Bristol-Myers Squibb, Gilead Sciences and Abbvie, and a research grant from Chugai and MSD. Lai Wei has research grants from BMS and Roche, and consulting fee from Abbott, Abbvie, BMS, Gilead and Novartis. Ming-Lung Yu is the consultant, advise board member and have grant support from Abbvie, BMS, Gilead, ROCHE, MSD and Abbott. Wang-Long Chuang has speaker fees from Gilead, BMS, MSD, Roche and Novartis, and is a member of Advisory Board: Gilead, Abbvie and Roche. Saeed S. Hamid has conference travel support from Gilead. Jia-Horning Kao has served as a consultant for AbbVie, Bristol-Myers Squibb, Gilead Sciences and Roche, and has also served on speaker’s bureaus for Roche, Bristol-Myers Squibb, Gilead Sciences, and Novartis. Osamu Yokosuka has research grants from Chugai Pharmaceutical, Bayer, MSD, Daiichi-Sankyo, Tanabe-Mitsubishi, and Bristol-Myers Squibb, and received speakers fees from Merck Sharp & Dohme, Kowa Souku, Sysmex, Chugai Pharmaceutical Co, GlaxoSmithKline, Bristol-Myers Squibb, Ajinomoto-Seiyaku, Bayer, Abbott, Given Imaging, Mitsubishi Tanabe Pharm, Taiho Yakuhin, Dainippon Sumitomo Pharm, and Igaku-Seibutsugakuinstitute.

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