Deep-sequencing study of HCV G4a resistance-associated substitutions in Egyptian patients failing DAA treatment

**Purpose:** To study resistance-associated substitutions using next-generation sequencing in Egyptian hepatitis C virus-infected patients failing direct-acting antiviral treatment.

**Methods:** The current study describes three cases of treatment failure in patients referred to Zagazig Viral Hepatitis Treatment Center (ZVHTC), Sharkia Governorate, Egypt. RAS were identified and characterized using deep sequencing. The first patient had breakthrough while receiving a daclatasvir (DCV)+sofosbuvir (SOF) regimen, patient 2 relapsed after treatment with DCV+SOF+ribavirin (RBV), and patient 3 relapsed after DCV+SOF therapy. A serum sample was collected from each patient at failure and sent to Vall d’Hebron Research Institute at Hospital Universitari Vall d’Hebron in Barcelona (Spain) for deep-sequencing to identify and characterize the RAS present in the samples.


**Conclusion:** To the best of our knowledge, this is the first report from Egypt of patients failing DAA-based therapy, describing the associated RAS. This information will be of help to understand the natural history of HCV in Egyptian patients and guide the proper choice of retreatment protocols.

**Keywords:** resistance-associated substitutions, RAS, subtype 4a, treatment failure, Egypt, direct acting antivirals, DAA

**Introduction**

Egypt has one of the highest prevalence rates of hepatitis C virus (HCV) infection worldwide. In 2008, the Egypt Demographic and Health Survey (EDHS) reported that nearly 10 million Egyptians were infected with this virus, whereas the 2015 EDHS identified a clear decline, reporting chronic HCV infection in only 6 million individuals. Nonetheless, this is a large number of cases, and it indicates that HCV remains a serious national health problem. A systematic review of HCV genotypes has reported limited diversity in Egypt, with dominance of genotype (GT) 4, accounting for 92.5% of cases (12.1% subtype 4a and 82.3% unknown subtypes), followed by 3.6% of GT1 cases, and 3.2% of mixed infections.

It has become clear that the root causes for this high prevalence (4.5–6.7%) are strongly associated with poor compliance with infection prevention and control in both the hospital and community settings. In 2006, the Egyptian National Committee for Control of Viral Hepatitis (NCCVH) was established to set up and
implement a national HCV control program. One of the Committee’s strategies was to provide affordable and accessible treatment, based at that time on pegylated interferon/RBV. In 2014, after successful negotiation between Gilead and the Egyptian Government represented by the NCCVH, the first direct-acting antiviral (DAA) drug, sofosbuvir (SOF) was introduced. Treatment regimens using this agent led to a sustained virological response (SVR) rate of 90%. In 2015 to 2016, new combinations were approved in Egypt to improve antiviral treatment and cover all patients chronically infected with HCV.\(^5\) The HCV program is continuously updated to open the prospect of future elimination of the disease in Egypt.

Real-life studies using DAA-based regimens in the Egyptian population have reported very few cases of the breakthrough and relapse categories of treatment failure. In this study, we report on 3 HCV subtype 4a-infected Egyptian patients who failed to respond to regimens of daclatasvir (DCV) + SOF with/without RBV. A RAS study was performed using deep-sequencing to investigate the patients’ RAS profile in targeted and non-targeted regions of the HCV proteins, NS3, NS5A, and NS5B.

Materials and methods

Patient samples

The original three serum samples were from HCV-infected patients who had failed DAA-based antiviral treatments in the Zagazig Viral Hepatitis Treatment Center (ZVHTC), Sharkia Governorate, Egypt. The study was approved by ZVHTC and the three patients signed an informed consent for participation. The RAS analysis was approved by the clinical research ethic committee of Hospital Universitari Vall d’Hebron. To perform the study of resistance-associated substitutions (RAS), one sample from each patient taken during 2018 at the time of failure was delivered on dry ice to Vall d’Hebron Research Institute at Hospital Universitari Vall d’Hebron (VHIR-HUVH) in Barcelona, for characterization using a next-generation sequencing (NGS) technique adapted to the MiSeq platform.

Definitions

For the present study, HCV viral breakthrough was defined as an increase in viral load at the end of antiviral treatment, even though HCV RNA had been undetectable during the treatment period. Viral relapse was defined as confirmed detectable HCV RNA levels during the post-treatment follow-up period in patients who had an undetectable HCV RNA level at the end of treatment.

Patient 1

Patient 1 was a 65-year-old diabetic woman with hypertension. She was receiving gliclazide and metformin to manage diabetes mellitus, and amlodipine and torsemide for hypertension. On ultrasound study, her gall bladder showed a thickened wall and internal calculi. There was no ascites, but splenomegaly was visualized (16.7 cm, superior to inferior axis). The superior mesenteric vein and splenic vein were enlarged (>10 mm), and the diameter of the portal vein was 13 mm. The portal venous flow was slow (<15 cm/sec), and showed reversal or to-and-fro flow. The patient was diagnosed with compensated liver cirrhosis and splenomegaly. Her condition was classified as Child Class A, with a Child Pugh (CP) score of 5 and MELD score of 9. The signs of liver cirrhosis included the following: surface nodularity, a generally coarse echotexture, segmental atrophy, caudate width:right lobe width ratio >0.65, and a reduction (<30 mm) in the transverse diameter of the medial segment (segment IV) of the left lobe. The body mass index (BMI) was 26.4, hemoglobin measured 13.3 g/dL, platelet count was 120×10\(^9\)/L, and total leucocyte count was 4.8×10\(^9\)/L. The international normalized ratio (INR) was 1.23. Serum albumin, total bilirubin, alanine transaminase (ALT), and aspartate transaminase (AST) levels were 3.9 g/dL, 0.91 mg/dL, 95 U/L, and 165 U/L, respectively. Alpha fetoprotein (AFP) concentration was 16 ng/mL, antinuclear antibodies (ANA) tested negative, and serum creatinine was 0.6 mg/dL.

On virological testing, HBsAg was negative and the virus was identified as HCV genotype 4a (G4a). HCV viral load in serum was 1,959,000 IU/mL, as determined by the TaqMan Real Time PCR assay (Roche Applied Science, Basel, Switzerland), which had a lower limit of detection of 15 IU/mL.

The patient had never received antiviral treatment (naïve) and was cirrhotic. She was referred to ZVHTC for treatment and was prescribed SOF (400 mg daily) plus DCV (60 mg daily) for 12 weeks (12 W), reporting good treatment adherence.

On HCV RNA monitoring at week 4 (4 W) of treatment, the virus was undetectable. At 12 W, the end of treatment (EOT), HCV RNA measured 393 IU/mL and at 12 weeks follow-up after stopping therapy (12 WFU), HCV RNA level was 707,999 IU/mL. The patient was...
retreated with SOF/DCV/RBV for 24 weeks and attained SVR.

**Patient 2**

Patient 2 was a 48-year-old man with a diagnosis of HCV cirrhosis and a history of liver encephalopathy attacks. He was referred to ZVHTC for treatment. The patient was treatment-experienced and had failed a 6-month course of SOF+RBV.

Ultrasonography showed an enlarged spleen (18 cm, superior to inferior axis), portal vein diameter of 11 mm, and signs of hepatic cirrhosis (surface nodularity, overall coarse and heterogeneous echotexture, segmental hypertrophy/atrophy, caudate width:right lobe width ratio >0.65, and a reduction in the transverse diameter (<30 mm) of segment IV of the left lobe. The BMI was 36.3. Blood tests showed a hemoglobin level of 13.6 g/dL, platelet count $88 \times 10^9$/L, and total leucocyte count $2.7 \times 10^9$/L. The INR was 1.28. Liver function tests yielded the following results: serum albumin 3.7 g/dL, total bilirubin 0.84 mg/dL, ALT 174 U/L, and AST 168 U/L. AFP measured 22 ng/mL, ANA tested negative, and serum creatinine was 0.93 mg/dL. The patient received a diagnosis of portal hypertension with splenomegaly and was classified as Child Class A, with a CP score of 6 and MELD Score of 9.

On virological testing, HBsAg was negative and the virus was identified as HCV G4a. Prior to designing a second therapy, serum HCV viral load, measured with the same real-time PCR method described above, was found to be 9,285,197 IU/mL.

The patient had never received antiviral treatment (naïve). At ZVHTC he was prescribed SOF (400 mg daily) plus DCV (60 mg daily) plus RBV (600–1000 mg, with dose adjustments according to tolerability) for 12 weeks. He reported good treatment adherence.

HCV-RNA was monitored at 4 W, 12 W EOT, and 12 and 24 W follow-up after stopping therapy (12 and 24 WFU). On RT-PCR monitoring, HCV RNA was undetectable at 4 W, 12 W EOT, and 12 WFU. At 24 WFU, viral load was 402,020 IU/mL. He was diagnosed as having HCV relapse.

The patient’s clinical condition worsened, with recurrent attacks of hepatic encephalopathy, increasing ascites, and development of hepatic hydrothorax. The patient could not afford the cost of a liver transplantation. He has not been retreated and is currently under conservative treatment for liver failure.

**Methods**

A serum sample was collected from each patient at treatment failure and sent to Vall d’Hebron Research Institute at Hospital Universitari Vall d’Hebron (VHIR-HUVH, Barcelona) for deep-sequencing study. HCV RNA was obtained from 140 µL of serum by manual extraction using the QIAmp Viral RNA Mini Kit (Qiagen, Hilden, Germany) following the manufacturers’ instructions and avoiding contamination. A high-resolution HCV subtyping method was used to identify HCV subtype and mixed infections (infection with more than one subtype at the same time). The HCV NS3, NS5A and NS5B regions, which are targeted
by DAAs, were amplified by RT-PCR-Nested using G4a-specific primers (Table 1). Reverse transcription and the first round of PCR amplification were performed with the Transcriptor One-Step RT-PCR Kit (Roche Diagnostics, Mannheim, Germany), using an annealing temperature of 55 °C and 40 seconds of elongation during 35 cycles. The KAPA HiFi HotStart PCR Kit (KAPA Biosystems, Wilmington, Massachusetts, USA) was used for nested amplification, applying an annealing temperature of 55 °C and elongation time of 15 seconds for a total of 30 cycles. The KAPA Library Preparation Kit for Illumina Platforms (KAPA Biosystems, Wilmington, Massachusetts, USA) was used to prepare the DNA library, and the SeqCap Adapter Kit A/B (Nimblegen, Roche, Mannheim, Germany) to label each patient’s amplified product. KAPA Pure Beads (KAPA Biosystems, Wilmington, Massachusetts, USA) were used at a 0.8:1 ratio for the purification steps. Deep sequencing on the MiSeq (Illumina) platform was performed using the MiSeq Reagent Kit v3 for 600 cycles (Illumina, San Diego, California, USA).

An in-house R script was used to obtain RAS reports from raw data after running the MiSeq in 2×300 mode. Additional details related to the RT-PCR-Nested amplification and deep-sequencing methodology can be found in a previous study.

A bioinformatics haplotype-centric procedure was developed to filter and exclude reads with lower quality than a minimum requirement. The FLASH software program was used to overlap paired-end reads, establishing a minimum overlap between R1 and R2 of 20 bp, with a maximum of 10% differences. A detailed description of these steps can be found in the study by Soria et al.

Because of the considerable intra-subtype diversity of HCV, 1068 sequences were retrieved from the Los Alamos databank to generate a consensus sequence (defined by the most frequent amino acid at each position) for subtypes G1a, G1b, G2c, G2i, G2j, G3a, G4a, and G4d to designate amino acid changes. All the reported amino acid changes were included in a final report, highlighting those changes that have been described as resistance associated substitutions compared to the G4a consensus sequence, but also including all other changes found in this study (Figure 1 and Table 2).

### Results

The NS3, NS5A and NS5B sequences from the three patients have been deposited in the GenBank database with accession number “PRJNA554981”.

<table>
<thead>
<tr>
<th>NS3</th>
<th>RT-PCR</th>
<th>NESTED</th>
<th>NESTED</th>
</tr>
</thead>
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<tr>
<td>FW</td>
<td>5Bu4a7952</td>
<td>13u4a7952</td>
<td>13d4a8389</td>
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<tr>
<td>RV</td>
<td>13u4a7952</td>
<td>13d4a8389</td>
<td>13d4a8584</td>
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</table>

Table 1 Specific primers for G4a RT-nested-PCR amplifications of DAA-targeted regions NS3, NS5A and NS5B.
At 4 weeks of DAA-based treatment, patients 1 and 3 showed an early positive antiviral response, whereas patient 2 had a viral load of 132 IU/mL and continued treatment for 24 weeks.

Patient 1 tested HCV RNA-positive at EOT (12 W), with a value of 393 IU/mL, suggesting viral breakthrough. At 12 WFU, viral load had increased to 707,999 IU/mL. In the case of patient 2, HCV-RNA was undetectable at 12 W and 16 W during treatment, and at 24 W EOT. At 12 WFU, viral load was 865,000 IU/mL, reflecting viral relapse. Patient 3 had undetectable HCV-RNA at 12 W, and 16 W during treatment, and at 24 W EOT. At 12 WFU, viral load was 865,000 IU/mL, re

Interestingly, deep-sequencing study showed two RAS (R155C and D168E) at very low frequencies (1.3% and 1.5%, respectively) in the non-targeted NS3 region in patient 3. R155C causes resistance to glecaprevir and paritaprevir, whereas D168E is associated with resistance to voxilaprevir, simeprevir, and paritaprevir (Table 2C). We also observed the minority substitution K24R (1.9%), which confers ledipasvir resistance in HCV G1a,10,11 but has an unknown RAS effect in G4a.

No RAS were observed in the NS5B region in any patient. However, alignment of the patients’ sequences...
with the consensus G4a sequence, composed of isolates downloaded from the Los Alamos database, identified two NS5B amino acid substitutions (A235V, R304K) with an unknown RAS effect. Specifically, all three patients were carrying the A235V substitution in almost all reads (99.3–100%), whereas R304K, reported at baseline in Portuguese G4a patients, compared to a subtype 4a consensus sequence retrieved from the Los Alamos databank are shown in Figure 1. For each patient and region we report (in columns) the region, the amino acid change, the number of reads (sequences) obtained, and the frequency of each substitution among the

### Table 2A. Patient 1. Mean fold-change in resistance to DAAs of each RAS identified

<table>
<thead>
<tr>
<th>Region (Total Reads)</th>
<th>RAS (Frequency %)</th>
<th>ASV</th>
<th>GLE</th>
<th>GZR</th>
<th>PTV</th>
<th>SMV</th>
<th>MK-7009</th>
<th>VOXI</th>
</tr>
</thead>
<tbody>
<tr>
<td>NS3 (58,871)</td>
<td>NO RAS IDENTIFIED IN THIS REGION</td>
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<td></td>
<td></td>
<td></td>
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<tr>
<td>NS5A (59,492)</td>
<td>L28M (100.0)</td>
<td>10</td>
<td>n.d.</td>
<td>100</td>
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<tr>
<td></td>
<td>L30S (99.6)</td>
<td>150–157</td>
<td>4</td>
<td>100</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>L28M+L30S (99.6)</td>
<td>5000</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NS5B (63,756)</td>
<td>NO RAS IDENTIFIED IN THIS REGION</td>
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### Table 2B. Patient 2. Mean fold-change in resistance to DAAs of each RAS identified

<table>
<thead>
<tr>
<th>Region (Total Reads)</th>
<th>RAS (Frequency %)</th>
<th>ASV</th>
<th>GLE</th>
<th>GZR</th>
<th>PTV</th>
<th>SMV</th>
<th>MK-7009</th>
<th>VOXI</th>
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<tbody>
<tr>
<td>NS3 (66,949)</td>
<td>NO RAS IDENTIFIED IN THIS REGION</td>
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<td></td>
<td></td>
<td></td>
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<tr>
<td>NS5A (26,160)</td>
<td>L30R (100.0)</td>
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<td>2</td>
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<tr>
<td>NS5B (32,337)</td>
<td>NO RAS IDENTIFIED IN THIS REGION</td>
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</table>

### Table 2C. Patient 3. Mean fold-change in resistance to DAAs of each RAS identified

<table>
<thead>
<tr>
<th>Region (Total Reads)</th>
<th>RAS (Frequency %)</th>
<th>ASV</th>
<th>GLE</th>
<th>GZR</th>
<th>PTV</th>
<th>SMV</th>
<th>MK-7009</th>
<th>VOXI</th>
</tr>
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<tbody>
<tr>
<td>NS3 (56,665)</td>
<td>R155C (1.3)</td>
<td>2.6</td>
<td>59</td>
<td>n.d.</td>
<td>n.d.</td>
<td>2.5–20</td>
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<tr>
<td>NS5A (58,493)</td>
<td>L28M (100.0)</td>
<td>10</td>
<td>n.d.</td>
<td>100</td>
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<td></td>
<td>L30H (12.1)</td>
<td>155–1215</td>
<td>4</td>
<td>100</td>
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<tr>
<td></td>
<td>L30S (87.9)</td>
<td>150–157</td>
<td>64</td>
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<td></td>
<td>L28M+L30H (12.1)</td>
<td>100</td>
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<tr>
<td></td>
<td>L28M+L30S (87.9)</td>
<td>100</td>
<td></td>
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<tr>
<td>NS5B (36,253)</td>
<td>NO RAS IDENTIFIED IN THIS REGION</td>
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**Notes:** n.d. = described as resistant but with no data; empty box indicates no resistance.

**Abbreviations:** ASV, asunaprevir; DAA, direct-acting antiviral; DCV, daclatasvir; DSV, dasabuvir; EBR, elbasvir; GLE, glecaprevir; GZR, grazoprevir; LDV, ledipasvir; MK-7009, vaniprevir; OMV, ombitasvir; PIB, pibrentasvir; PTV, paritaprevir; RAS, resistance-associated substitutions; SMV, simeprevir; SOF, sofosbuvir, VEL, velpatasvir; VOXI, voxilaprevir.

Amino acid substitutions in patients 1, 2, and 3 in the three HCV antiviral target regions (NS3, NS5A, NS5B) compared to a subtype 4a consensus sequence retrieved from the Los Alamos database are shown in Figure 1. For each patient and region we report (in columns) the region, the amino acid change, the number of reads (sequences) obtained, and the frequency of each substitution among the
The emergence of HCV resistance variants (ie, Amer et al) in this study, we report in NS5A, and N130D, P135V, A190S, N105S, F108R/C, V124A, D126E, V228I, E237A, E248D, and T254A in NS5B. R155C causes resistance to glecaprevir and paritaprevir, whereas D168E confers resistance to voxilaprevir, simeprevir, and paritaprevir. 10,11 R155C causes resistance to glecaprevir and paritaprevir, whereas D168E confers resistance to voxilaprevir, simeprevir, and paritaprevir. 10,11

The level of resistance to an inhibitor depends on several conditions: the genetic barrier to resistance of the HCV region, the drug used, the type of RAS, the frequency at which RAS are present in the viral isolate, and the presence of RAS combinations in the same viral particle, all of which can significantly increase resistance to some DAA treatments. RAS combinations in the same genome have been shown to cause a dramatic increase in resistance to the inhibitor used in the failed treatment and to other inhibitors of the same family. 17 For example, L28M+L30H causes a mean-fold resistance of 10,000 to ledipasvir (Table 2C).

Our results show that L30R appeared alone in a patient who had received RBV, whereas other L30 substitutions seem to require combination with L28 mutations. 16 This observation should be confirmed in additional patients to know whether it is an occasional finding or the rule. A deep-sequencing study in a large cohort of patients treated with and without RBV has just been completed, and this molecule did not have an influence on the type or number of RAS at HCV treatment failure (data not shown, paper in preparation).

Coinciding with the results of a previous study 16 the L28M and L30 substitutions were the main RAS in G4a. However, in contrast to that same study 18 in which L30R was the majority substitution, L30S was the most prevalent in our patients. The RAS pattern in G4a was completely different from that of G4d, the most prevalent G4 subtype in Spain, in which the most common RAS was T58P (data not shown, paper in preparation). Moreover, NS5A polymorphisms at Y93 (Y93H/S/T) were not identified in the 3 patients failing therapy, in agreement with previous studies. 11,18 Additional studies are needed to demonstrate whether or not A235V/T and R304K in the NS5B protein and K24R in NS5A are resistant to the inhibitors. Other highly prevalent substitutions observed in our study, such as A150V in NS3, 19 V52I, T75I, V99I, 1101V, 20 N105S, F108R/C, V124A, D126E, 21 and V130I, 20 in NS5A, and N130D, P135V, A190S, E202D, 20 V228I, E237A, E248D, and T254A in NS5B should still be considered simple polymorphic substitutions without resistance implications, requiring further study. As there are few studies on RAS emerging in patients failing DAA treatment, more work is needed to determine whether any newly described substitutions can be classified as RAS, as adaptive mutations, or as random occurrences.

To our knowledge, this is the first study investigating resistance-associated substitutions using next-generation sequencing in Egyptian HCV-infected patients failing DAA treatment. Deep sequencing-based studies of the three antiviral targeted regions facilitate identification of major and minor substitutions, the frequency at which they are present in the viral isolate, and whether or not they are combined with others in the same viral genome. This information is useful for designing successful retreatment strategies, and is of particular importance in the absence of next-generation HCV treatments incorporating pangenotypic DAAs that will be effective in Egypt and other parts of the world having a high GT4 burden. Until future pangenotypic DAAs become affordable, accessible, and easily provided, a scenario that may not materialize for several years, precise identification of RAS leading to treatment failure is of value for better patient management, offering the possibility of improved quality of life.

Discussion
The recent introduction of DAA therapy has revolutionized the treatment of chronic HCV infection in several countries worldwide, including Egypt, with very high SVR rates in clinical trials and slightly lower rates in real-life cohorts. 13 The emergence of HCV resistance variants (ie, RAS), is the main reason for this difference, but little information on resistance in G4a patients has been reported, as most studies include only a small number of patients with this subtype, and very few have been done using deep-sequencing. 10,11,14,15 In this study, we report the RAS profile determined by this technique in Egyptian patients failing DAA-based treatments.

All samples were analyzed for RAS in the NS3, NS5A and NS5B proteins by deep-sequencing using the MiSeq (Illumina) platform. The rationale to study all three DAA-targeted regions was that extra-target mutations and hitchhiking mutations can be selected at failure and negatively impact salvage treatments. In this sense, two minority RAS (R155C and D168E) in the non-targeted NS3 region were observed in patient 3 at very low frequencies (1.3% and 1.5%, respectively) (Table 2C). Neither of these minority RAS would have been detected using population (Sanger) sequencing, and both of them could have reduced retreatment options. 14,16 R155C causes resistance to glecaprevir and paritaprevir, whereas D168E confers resistance to voxilaprevir, simeprevir, and paritaprevir. 10,11

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In view of lack of facilities or very high cost of RAS testing in Egyptian laboratories together with the huge number of patients treated daily, the NCCCVH recommends empirical retreatment of patients who fail NS5A inhibitor-based therapy with a combination of HCV multtargeted drugs. Quadruple therapy with sofosbuvir, daclatasvir, and simeprevir plus ribavirin (SOF/DCV/SMV/RBV) or sofosbuvir, ombitasvir, paritaprevir, and ritonavir plus ribavirin (SOF/OBV/PTV/r/RBV) for 12 weeks is recommended for these patients, provided they have neither decompensated liver function nor an estimated glomerular filtration rate (eGFR) below 30 mL/min/1.73 m² body surface area. The efficacy and safety of SOF/OBV/PTV/r/RBV quadruple therapy has been assessed in chronic HCV GT4 patients failing DAAs regimens in Egypt. SVR was achieved in 97% (109/113) of patients with good tolerability and safety of the drug combination. The same group of researchers conducted another study using SOF/DCV/SMV/ RBV in a similar cohort of Egyptian patients, and reported similar efficacy (SVR 97%) and safety results. However, the studies from Egypt are limited and include relatively small patient samples. Efficacy and safety analyses of data from the Egyptian NCCVT in patients using these DAA combinations are needed. In July 2017 the Food and Drug Administration (FDA) approved a regimen consisting of a fixed-dose combination of sofosbuvir, velpatasvir and a second-generation HCV-protease inhibitor, voxilaprevir, for retreatment of HCV GT 1–6 after failure with sofosbuvir or other NS5A inhibitors without RAS testing. Again, clinical trials are needed to assess the efficacy and safety of these new generations of DAAs in Egyptian patients. Until these pangenotypic DAAs become affordable, accessible, and easily provided in Egypt’s market based on clinical proof of better safety and efficacy, a scenario that may not materialize for several years, precise identification of RAS leading to treatment failure is of value for better patient management, providing the possibility of improved quality of life.

**Consent for publication**

Written informed consent was obtained from the cases included in the study.

**Data availability**

The data used to support the findings of this study are available from the corresponding author upon request.

**Acknowledgments**

This study was supported by the Spanish Ministry of Health, Consumer Affairs, and Social Welfare, grant name: Plan Estratégico Nacional contra la Hepatitis C. This study was also funded by Instituto de Salud Carlos III, PI15/00856 and PI16/00337, cofinanced by CIBERehd (Consorcio Centro de Investigación en Red de Enfermedades Hepáticas y Digestivas), which is funded by Instituto de Salud Carlos III and Centro para el Desarrollo Tecnológico Industrial (CDTI) from the Spanish Ministry of Economy and Business, grant number, IDI-20151125. The authors thank Celine Cavallo for English language support and helpful editing suggestions.

**Disclosure**

The funder, Roche Diagnostics S.L., provided support in the form of salaries for one author (Josep Gregori), but the company did not have any additional role in the study design, data collection and analysis, decision to publish, or preparation of the manuscript. The authors report no other conflicts of interest in this work.

**References**