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INTRODUCTION

HCV infection remains an important health threat, with 71 million chronically infected people, resulting in 400 thousand deaths yearly. The use of direct-acting antivirals (DAA) targeting the viral nonstructural 3 protease (NS3P), NS5A and NS5B-polymerase increased cure rates to >90%.1 However, emergence of resistance-associated substitutions (RAS)1,2 compromises efficacy of DAA regimens.1,4 Treatment failures have been widely reported and will increase as patients are treated worldwide.1

HCV shows extensive genetic diversity with eight major genotypes and >90 subtypes.1,2 Genotype 4 represents ~8% of infections worldwide, being...
Significance of this study

How might it impact on clinical practice in the foreseeable future?

► Our study provides a unique overview of the evolutionary landscape underlying the emergence of RASs conferring viral resistance to DAAs in cell culture, which provides valuable information for strategies to avoid emergence and spread of DAA-resistant variants for prevention of future treatment failures in the clinic.

► The highly efficient genotype 4a infectious system provides a key tool for development and testing of vaccine candidates, a key intervention to eliminate HCV at a global scale.

highly predominant in the Middle-East and North and Central Africa. Particularly, ~93% of >5 million HCV infections in Egypt were caused by genotype 4. This genotype is highly heterogeneous with >18 recognised subtypes. In addition, due to human migration and increasing transmission among intravenous drug users and individuals with high-risk sexual practices, the prevalence of genotype 4 is currently increasing in Europe. Subtype 4a is common, particularly in Egypt, the origin of prototype 4a strain ED43. Approved DAA regimens are highly efficient against genotype 4 However, high rates of treatment failure with preexisting and emergent RASs were recently reported in subsets of genotype 4-infected patients. In Egypt alone, ~2.4 million patients have been treated with DAAs, in particular, various NS5A inhibitors combined with the polymerase-inhibitor sofosbuvir. Although the sustained virologic response (SVR) rates were >90%, treatment failures were reported. Antiviral resistance will not necessarily alter the current treatment guidelines, but it could potentially affect treatment options in the future. Although DAA-resistance occurs at low prevalence, the extensive cross-resistance between the same molecular classes of drugs could limit future treatment options, especially since no additional antivirals are being developed for the treatment of HCV. Thus, generating new knowledge on viral resistance to DAAs is of great importance to prevent treatment failure in the future and to avoid the emergence and transmission of DAA-resistant viruses to highly exposed populations. This effort will require detailed understanding of the mechanisms underlying emergence of RASs.

In addition, prophylactic HCV vaccines will be essential for preventing transmission globally. Efficient infectious cell culture systems representing the major HCV genotypes can play an important role in the development and testing of vaccine candidates. Vaccine candidates based on inactivated whole-virus-particles are dependent on the efficient production of the virus in cell culture, which for HCV can only be achieved after virus adaptation, thus, understanding such processes is fundamental to generate relevant candidates. In addition, evaluation of the ability of vaccines to induce broad cross-genotype neutralising antibodies requires the establishment of culture systems representing the genetic heterogeneity of HCV, and here infectious full-length systems are most relevant since they recapitulate the entire viral life cycle.

Efficient full-length infectious culture systems have been developed for selected strains of genotypes 1a, 2a, 2b, 3a and 6a after complex adaptation processes. For genotype 4, a full-length cell culture system has recently been reported, but with limited propagation in Huh7.5 cells, thus, a high titre system is still required for most studies on the viral life cycle, antivirals and vaccine development. Here, we aimed at developing a robust and efficient (high infectivity titre) full-length infectious system for HCV genotype 4a, permitting in-depth analysis of the emergence of DAA-resistance and assessments of the efficacy and barrier to resistance of clinically relevant DAA regimens.

MATERIALS AND METHODS

Construction of HCV genotype 4a clones

The in vivo infectious strain ED43 clone was described. The chimeric genome comprising ED43 Core-NS5A (C5A) and JFH1-NS5B and—untranslated regions (UTRs) was generated by replacing the 5’UTR from ED43 5’UTR-JFH1 recombinant with the corresponding JFH1 sequence. Mutations were introduced by QuickChange site-directed mutagenesis kit (Agilent) or by fusion PCR. The HCV sequences of final plasmid preparations were confirmed by Sanger sequencing (Macrogen). The nucleotide and amino acid (aa) numbers refer to the ED43 full-length recombinant sequence.

Production and analysis of culture viruses

Viability of HCV recombinants was tested by transfection of RNA transcripts into Huh7.5 cells using Lipofectamine 2000 (ThermoFisher). Cells were subcultured every 2–3 days and viral passage was performed as described. Harvested cellular pellets were centrifuged at 2000 rpm for 5 min, washed 2–3 times with sterile phosphate buffered saline (PBS) (Sigma-Aldrich) and stored in 1 mL of Trizol (ThermoFisher). Infectivity titres were determined in triplicate and reported as log10 focus-forming units per milliliter (log10 FFU/mL).

Genome analysis of recovered viruses

Next-generation sequencing (NGS) was performed as described. Briefly, RNA was extracted, and reverse transcription (RT)-PCR performed to obtain complete HCV open reading frame (ORF) amplicons. For RT, primer 5'-CTAAGGTCCGGATGTGTAAGC-3', and for PCR, primers 5’-TGCTGCTGATAGGGTGCTTGCG-3’ and 5’-AGGGTGCCGATGTGTAAGCTGCC-3’ were used. PCR amplicons were processed with NEBNext Ultra II FS DNA Library Prep Kit (New England Biolabs). In order to sequence up to 500 bp to cover NS3P, NS5A domain I or the NS5B-palm domain (up to 167 aa), we performed size selection. Sequencing was carried out in-house by Illumina MiSeq using 500 cycles v2 kit. Data were analysed for single-nucleotide polymorphism (SNP). The haplotypes were reconstructed and plotted using GraphPad Prism version 6.

For ORF analysis, the linkage and haplotype reconstruction could not be applied in one read pair. Therefore, the frequency development of SNP variants over time was used. For specific samples, PCR amplicons were subcloned into TOPO-XL2 vector (ThermoFisher), allowing ORF linkage analyses. Each clone was sequenced by Sanger and aligned to build phylogeny and ancestral reconstruction.

For ED43(C5A) recombinants, recovered viruses were analysed by Sanger. For viral 5’UTR sequences, we used a 5’RACE procedure on culture supernatants. The PCR products were analysed by Sanger.

Virus stocks and treatment assays

Inhibitors (Acme Bioscience) were diluted in dimethyl sulfoxide. Escape and treatment assays were conducted using

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Inhibitors (Acme Bioscience) were diluted in dimethyl sulfoxide.
the 5th, 7th and 10th passages of ED43-20m (figure 1A,B), and not the final ED43cc virus, in order to optimise the timing of the different experiments. For the reverse-genetic testing of RAS, the ED43-31m/+A1973T/-Q2931R (named ED43-31m_opt) recombinant was used (figure 1A).

Treatments were initiated upon virus spread (≥90% HCV antigen-positive cells) and the inhibitors were added every 2–3 days when cells were subcultured. For concentration-response assays using described methods, 6,7,14,27 stocks of nontreated and of single-treatment escape viruses were prepared. To prepare virus stocks from escape viruses, supernatants recovered from the treatment experiments were used to infect naive Huh7.5 cells that were then cultured without inhibitors until virus spread. Sequences of virus stocks were confirmed by NGS. The half-maximal effective concentration (EC50) value was calculated using GraphPad Prism version 6.

For combination treatments with indicated DAA concentrations, escape stock viruses were used to infect naive Huh7.5 cells and followed until virus spread. Viral supernatants were collected at treatment initiation (day 0) and frequently hereafter, and HCV sequences were confirmed by NGS. Unless otherwise stated, the viruses were treated 28–30 days. Afterwards, cultures were followed without drugs for 14 days; infection was defined as eradicated, if no HCV antigen-positive cells were detected. 3,4

RESULTS
Development of HCV genotype 4a full-length infectious cell-culture systems
Like most prototype HCV clones, the full-length clone of genotype 4a prototype strain ED43 (pED43) is infectious in chimpanzees, but not in Huh7.5 cells. Therefore, we aimed at identifying adaptive substitutions permitting culture of this full-length clone. It has been reported that the JFH1-NS5B has high replication activity in cell culture. 35,36 Taking advantage of this, we showed that for various genotypes, cell culture adaptation of recombinants with genotype-specific Core-NS5A (C5A), JFH1-NS5B, and JFH1-UTRs resulted in identification of mutations that permitted replication of the corresponding full-length genome. 31 Thus, we first generated JFH1-based ED43-4C5A recombinants for this same purpose.

Culture-efficient ED43(C5A)-adapted recombinant
RNA transcripts from ED43(C5A)-clones with or without A1672S(NS4A), required for culture of genotypes 1a, 2a and 2b strains, yielded no HCV antigen-positive Huh7.5 cells during 30 days of follow-up. However, ED43(C5A)-2m (figure 1A) with A1786V(NS4B), previously used for adaptation of the ED43 5’UTR-NS5A recombinant, 30 and A1672S(NS4A) spread at day 52 (4.0 log10FFU/mL) and acquired five additional mutations associated with viability in cell culture (online supplemental figure 1A). Following transfection, ED43-18m was nonviable, but ED43-20m spread at day 41 (2.6 log10FFU/mL). The infectivity titres increased during consecutive passages, reaching −5.0 log10 FFU/mL at passage 10 (figure 1B and table 1).

To identify substitutions evolving during serial passage of ED43-20m in Huh7.5 cells, we analysed extracellular viral RNAs using NGS. The substitutions that emerged with similar patterns were grouped as shown in figure 1B. L1466M(NS3) and K2597N(NS5B) were found in >80% of the viral population at second passage (figure 1B). However, the fitness of a recombinant with these substitutions (ED43-22m) remained low and the virus acquired additional changes (online supplemental figure 2).

Thus, we investigated other substitution patterns (figure 1B). As shown in figure 1B, two different viral populations evolved that showed relatively equal prevalence in passage 6. However, one population became dominant in passage 10. The stepwise acquisition of substitutions was confirmed by further analysis of 6th and 10th passage viruses (figure 1C), by Sanger sequencing analysis of TA clones of amplicons. Clones were analysed by linkage and ancestral reconstruction based on phylogeny. L1466M and K2597N were observed in all clones, followed by the emergence of two different viral populations. Interestingly, we found that the mutation patterns were similar between cell-free and cell-associated viruses (figure 1B and online supplemental figure 3).

Based on the evolution during ED43-20m adaptation, we constructed a recombinant harbouring all dominant 10th passage substitutions, except for A1973T(NS5A) (figure 1A,B). We also introduced F1572L, detected in the ED43(C5A)-7m recovered virus and emerging during the first five passages of ED43-20m (figure 1A,B and online supplemental table 1). The resulting ED43-31m spread fast and produced −4.3 log10FFU/mL (figure 1A,D). We did not detect other substitutions at >5% of the viral population in the second passage (table 1). However, the titres of this virus did not match those observed at passage 10 of ED43-20m, and thus there was room for further optimisation.

Among the original 20 m substitutions, Q2931R and C2992Y had partly reverted (table 1), and R2931Q was found to increase ED43-31m titres after transfection (figure 1D). In addition, titres of ED43-31m further increased after introducing A1973T(NS5A) (detected in the ED43-20m 10th passage virus but not initially included in ED43-31m) (figure 1D). Thus, we generated an ED43-31m recombinant containing A1973T and the reversion of Q2931R (ED43-31m/+/A1973T/-Q2931R; or ED43-31m_opt), which showed fast spread kinetics and slightly higher −4.6 log10FFU/mL post-transfection titres (figure 1E).

Finally, we found that a recombinant carrying the G38A mutation in the 5’UTR (ED43-31m_opt with 5’UTR G38A, designated ED43cc, GenBank accession: MW531222), which was detected in the 10th passage of ED43-20m, produced the highest titres mimicking the ED43-20m virus 10th passage, −4.7 and 5.1 log10FFU/mL in transfection and second passage, respectively. We did not detect additional substitutions emerging at >5% of the viral population in the second passage virus (table 1).
Figure 1  Full-length HCV genotype 4a infectious cell-culture system. (A) Schematic overview of the culture adaptation process of strain ED43. Substitutions introduced into the full-length recombinant ED43-20m are shown in black. Adaptive substitutions identified during passage of the ED43-20m virus are indicated by colours from panels B and C. (B) HCV infectivity (bars) determined by FFU assays and shown by mean of triplicates±SEM (left y-axis; break indicates the cut-off of the assay), and emergence of substitutions for ED43-20m (lines) during adaptation following transfection (T; samples not available from days 0 to 27) and serial passages (P1, 2, 3, etc.). Only substitutions with SNP frequencies >20% at any time-point are shown (right y-axis). For substitutions that emerged with similar patterns, means±SEM are shown. The substitutions with similar SNP frequencies are grouped and shown with the same colour. (C) Clonal ORF sequence analysis of ED43-20m virus obtained from the 6th and 10th passages; phylogeny was generated as described.31 Subsequently, ancestral reconstruction was performed based on alignment and phylogeny to predict the appearance of respective substitutions. Only substitutions indicated in (B) are shown. (D,E) Infectivity titres (y-axis) of specific ED43 recombinant viruses at indicated days (x-axis) after transfection of Huh7.5 cells. J6/JFH1 was included as a control. #The data were obtained from a separate experiment with similar J6/JFH1 titres. Statistical significance (p<0001, two-way ANOVA) is carried out for indicated samples and shown with asterisks (***). ANOVA, analysis of variance; FFU, focus-forming unit; ORF, open reading frame; SNP, single nucleotide polymorphism; SEM, standard error of the mean.
Table 1  Non-synonymous ORF mutations in ED43 recombinant viruses during culture adaptation

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<th>AA pos</th>
<th>AA change</th>
<th>AA pos</th>
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Only mutations with a frequency of >5% of the viral population in NGS analysis are shown. Shaded background indicates frequencies of engineered mutations.

*Nucleotide G at position 38 in 5'UTR was changed to A.
†HCV infectivity titres at indicated passage and day (parentheses) are shown as log₁₀FFU/mL.
‡These recombinant viruses had A at position 9132 instead of G, and this nt was maintained.
AA, amino acid; NGS, next-generation sequencing; NT, nucleotide; NT pos and AA pos, nucleotide and amino acid positions; SNP, single nucleotide polymorphism.
Overall, we revealed unique evolutionary details on HCV culture adaptation and developed full-length 4a recombinants that propagated robustly in HuH7.5 cells. The ED43ccc harbouring 32 changes compared with the consensus ED43 clone, including 5, 4 and 13 coding changes in NS3/4A, NS5A and NS5B, respectively (online supplemental figure 1B).

HCV evolutionary genetic networks resulting in DAA-resistance for genotype 4a

The recommended DAA-based regimens for patients with chronic genotype 4 infection include NS3 protease (grazoprevir, paritaprevir and glecaprevir), NS5A (elbasvir, ledipasvir, ombitasvir, velpatasvir and pibrentasvir) and NS5B polymerase (sofosbuvir) inhibitors.\(^\text{11–13}\) The evolutionary features underlying the emergence of viral resistance are not well characterised. The HCV genotype 4a infectious culture system can serve as a valuable model to explore determinants of virus escape from DAAAs. We, thus, performed NGS and linkage analysis permitting detailed investigation of emerging RASs during culture escape experiments and examined the evolution of putative fitness compensating substitutions throughout the HCV genome using reverse genetics.\(^\text{3,11–12}\)

Evolutionary pathways underlying emergence of RASs during treatments with protease inhibitors

To induce viral escape from protease inhibitors (PIs), we performed long-term treatments of the ED43 virus with paritaprevir, grazoprevir or glecaprevir at concentrations equivalent to 8xEC\(_{50}\).\(^\text{3,11–12}\) (figure 2A–C). For paritaprevir, NS3P-D168E emerged at day 7 after treatment initiation and became a major RAS when the virus escaped at day 16. Under a higher inhibitor concentration (128xEC\(_{50}\)), D168E decreased in prevalence and different RASs singly or in combination emerged. Y56H+D168A appeared at low frequency at day 21 but became dominant following viral escape. The escape virus (PAResc) showed cross-resistance to all tested PIs compared with the original virus with ≥64-fold increase in EC\(_{50}\) (figure 2F; online supplemental figure 4). However, an ED43 recombinant harbouring Y56H+D168A was highly attenuated (figure 2F) and engineered RASs reverted after a second drug-free passage (table 2). Therefore, compensatory substitutions might be required for stability of these RASs, which were maintained during second passage of the escape virus (figure 2A). NGS analysis of the complete viral ORF during treatment showed complex patterns of substitutions (online supplemental figure 5A). T1566S[NS3-H] and K2317T(NS5A) emerged together with Y56H/D168A, RASs developed, including A156T/V (figure 5A). T1566S[NS3-H]

Evolution induced by NS5A inhibitors

Under ombitasvir, elbasvir and ledipasvir treatments (concentrations equivalent to 100xEC\(_{50}\)), the main RASs responsible for ED43 escape emerged at day 5 (figure 3A–C).\(^\text{3}\) Following viral escape from ombitasvir (OMBesc), elbasvir (ELBesc), and ledipasvir (LEDesc), RASs conferring high-level resistance (figure 3G and online supplemental figure 6) became dominant (figure 3A–C). The main NS5A RASs L28V (ombitasvir), L30H+M31V (elbasvir) and L30P+Y93H (ledipasvir) did not result in a high loss of fitness when introduced in the ED43 recombinant and were maintained after second passage (figure 3H and table 2).

For pibrentasvir, we did not observe viral escape with 100xEC\(_{50}\) and performed an experiment at 10xEC\(_{50}\). The major RASs of L30F+M31V were not acquired as rapidly as for ombitasvir, elbasvir and ledipasvir, suggesting higher barrier to resistance (figure 3D). These RASs were dominant in escape viruses until day 28. Here, the concentration was increased to 100xEC\(_{50}\), resulting in diversification of the viral population, associated with increased viral suppression (figure 3D). L28M+L30F+M31V emerged as a major population, which led to viral escape (VELesc) and high resistance levels to NS5A inhibitors (except pibrentasvir) (figure 3EG and online supplemental figure 6). The ED43 L28M+L30F+M31V recombinant was highly fit and maintained the RASs after second passage (figure 3H and table 2).

Genome-wide NGS showed that substitutions outside NS5A domain I emerged in viruses escaping ombitasvir, elbasvir, ledipasvir and pibrentasvir (online supplemental figure 7A–D). Pibrentasvir 10xEC\(_{50}\) treatment resulted in viral eradication. However, 5xEC\(_{50}\) treatment led to escape by day 33 (figure 3E). Interestingly, we primarily observed NS5A RAS L30Δ (deletion) (figure 3E), and no viral suppression was observed at increased concentrations (10xEC\(_{50}\) and 100xEC\(_{50}\)), indicating that L30Δ conferred high resistance. Indeed, the escape virus (Pibesc) showed high levels of resistance to all NS5A inhibitors (figure 3FG and online supplemental figure 6). In contrast to other NS5A-RASs, the ED43-L30Δ recombinant was highly attenuated and acquired additional substitutions after second passage (figure 3H; table 2). One of these substitutions, T2047I (T75I, NS5A-numbering), also found in the second drug-free passage of Pibesc, increased fitness of the L30Δ recombinant (figure 3EH and online supplemental figure 7E).

Evolution induced by sofosbuvir

The ED43 virus treated with sofosbuvir was initially suppressed at 2xEC\(_{50}\) but escaped at day 77 (figure 4A). NGS analysis of HCV-ORF during treatment highlighted a complex network of coselection of substitutions underlying the gradual emergence of single NS5B-RAS S282T (online supplemental figure 8), coexisting with S282T+V322A (figure 4A). Interestingly, the S282T+V322A population increased during drug-free passages (figure 4A). The SOFesc virus had decreased sensitivity to sofosbuvir (figure 4B). Strikingly, the ED43 recombinants harbouring D2689G(NS5B;>5%) and
Figure 2  Substitution networks emerging in NS3P domain under protease inhibitor treatments. (A–C) ED43 full-length cultures were treated with protease inhibitors paritaprevir (A), grazoprevir (B) and glecaprevir (C) and analysed by NGS. The percentage of HCV-antigen positive cells was determined by immunostaining (line graph, left y-axis). The distribution of haplotypes (bar graphs, right y-axis) was determined by linkage analysis and indicated by different colours. Treatments were initiated with concentrations equivalent to 8x-EC_{50}, then concentrations were increased to 128x-EC_{50} at day 16 (paritaprevir), 70 (grazoprevir) and 63 (glecaprevir). Only haplotypes constituting ≥2% of the viral population are shown. Supernatant samples from the last treatment time-points were used for passage. P1 and P2: the first and second passages. (D) Efficacy of glecaprevir against indicated ED43 full-length DAA escape viruses. Values are means of triplicates±SEM. (E) EC_{50} values and 95% CI of indicated escape viruses were calculated from data shown in (D) and online supplemental figure 4. Fold changes in EC_{50} values were compared with the original ED43. (F) Fitness of ED43 recombinant virus harbouring NS3P RASs. For details, see figure 1D,E legend. The corresponding drug targets and protein-specific numbers are indicated for RASs. Other substitutions are shown with polyprotein specific numbers. #The data were obtained from a separate experiment with similar titres of the original virus. NGS, next-generation sequencing; RAS, resistance-associated substitution; SEM, standard error of the mean.
NGS analysis of recombinant ED43 viruses harbouring RASs recovered from the second passage culture

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- All recombinant viruses harbouring RASs were generated using the ED43-31m_opt backbone as shown in figures 2–4. Shaded background indicates frequencies of mutations that were introduced into this recombinant. Only non-synonymous (coding) mutations that represented >5% of the viral population are shown.*Target proteins in which the recombinants harbouring RASs were numbered.
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- Only non-synonymous (coding) mutations that represented >5% of the viral population are shown.*Target proteins in which the rec...
Figure 3  Substitution networks emerging in NS5A domain I under treatments with NS5A inhibitors. (A–E) ED43 full-length cultures were treated with ombitasvir (A), elbasvir (B), ledipasvir (C), velpatasvir (D) and pibrentasvir (E). (A–C): Inhibitor concentration: 100x-EC_{50}. (D) Initial concentration: 10x-EC_{50} increased to 100x-EC_{50} at day 28. (E) Treatment initiated at 5x-EC_{50}, then increased to 10x- and 100x-EC_{50}, at days 33 and 40, respectively. (F) Efficacy of pibrentasvir against ED43 full-length DAA escape viruses. Values are means of triplicates±SEM. (G) EC_{50} values and 95% CI of indicated escape viruses were calculated from data shown in (F) and online supplemental figure 6. (H) Fitness of ED43 recombinant viruses harbouring NS5A RASs. The data were obtained from the experiment shown in figure 2F. For details, see figure 2 legend. RAS, resistance-associated substitution; SEM, standard error of the mean.
enginedered S282T with or without A1309P(NS3H), which developed following the emergence of S282T (online supplemental figure 8), were relatively fit (figure 4C) and S282T was maintained after two consecutive drug-free passages (table 2).

**Efficacy of clinically relevant DAA combinations against genotype 4a**

Recommended DAA combinations include paritaprevir/ombitasvir, grazoprevir/elbasvir, ledipasvir/sofosbuvir, velpatasvir/sofosbuvir and glecaprevir/pibrentasvir.11 13 These regimens were efficient in suppressing the original ED43 virus (figures 5 and 6). Except for treatment with LED(5xEC50)/SOF(1xEC50), where we consistently detected HCV-positive cells, the original virus was eradicated in all treatments. Next, we tested whether DAA combinations remained efficient against their corresponding single drug escape ED43 variants.

DAA regimens based on PIIs (paritaprevir, grazoprevir or glecaprevir) and NS5A inhibitors (ombitasvir, elbasvir or pibrentasvir) were inefficient against viruses that had escaped one of the included drugs, and thus harboured RASs at baseline (figure 5A–C).4 31 Only glecaprevir/pibrentasvir was able to control the GLDesc and PIDesc viral infections during treatment (figure 5C). In contrast, paritaprevir/ombitasvir and grazoprevir/elbasvir combinations were inefficient to suppress infections with PI or NS5A-resistant viruses (figure 5A,B), which escaped after 2 weeks of treatment.

NGS and linkage analysis of viruses escaping from these combination treatments showed that in addition to the RAS at baseline, they all acquired RASs in the new target (figure 5D–F and online supplemental figure 9). Moreover, additional substitutions outside the drug targets emerged after combination treatments (figure 5D–F). Particularly, I2841V emerged in the GRAesc virus containing NS3P-A156M.

Similarly, we tested the DAA combinations of NS5A (ledipasvir or velpatasvir) and NS5B (sofosbuvir) inhibitors against the respective LEDesc, VELesc and SOFesc viruses and found that those viral infections were not eradicated (figure 6A,B). The treatment with velpatasvir/sofosbuvir resulted in high suppression and delayed spread of the SOFesc virus as compared with ledipasvir/sofosbuvir.

After combination treatments, the original LEDesc and VELesc viruses maintained the NS5A RASs (figure 6C,D and online supplemental figure 10). The original SOFesc virus acquired NS5A L28M, L30H and L30S, while maintaining NS5B-S282T, and acquiring additional substitutions throughout the ORF (figure 6C,D).

**Glecaprevir/pibrentasvir as a retreatment option for HCV genotype 4a with baseline resistance in culture**

Our data demonstrated that DAA combinations could not eradicate infections with viruses resistant to one of the included inhibitors. Clinically, glecaprevir/pibrentasvir has been investigated as a retreatment option for HCV-infected patients who failed DAA-containing regimens.37 Thus, we investigated its effectiveness against PAResc, GRAesc, OMDesc, LEDesc, ELDesc, VEDesc and SOFesc viruses. Infected cultures were treated with 4xEC50 glecaprevir/5xEC50 pibrentasvir. The infection was suppressed by day 7 of treatment and all except for PAResc and GRAesc viruses were eradicated after 28 days of treatment (figure 7A). NGS analysis of glecaprevir/pibrentasvir escape viruses showed that the GRAesc virus mainly harboured NS3P-A156M (combined with NS3P-A151V) and NS5A-L30P+P32L (figure 7B). The PAResc virus maintained NS3P-Y56H+D168A/V and acquired NS5A-L30A+T75I (as a minor population) (figure 7B and online supplemental figure 11). Among substitutions emerging outside NS3P and NS5A domain I, we also found I2841V(NS5B) in the GRAesc virus (figure 7B). Thus, baseline PI resistance compromised the effectiveness of glecaprevir/pibrentasvir.

**DISCUSSION**

We unravelled viral adaptation leading to the development of a high-titre full-length infectious culture system for HCV genotype...
Figure 5  Efficacy of DAA combinations containing PIs and NS5A inhibitors against ED43 escape viruses. (A–C) For each DAA combination, concentrations of 5×EC₅₀ of NS5A inhibitors were used in combination with either 2×, 4× or 8×EC₅₀ of corresponding PIs. Black dashed lines indicate the time when the treatments were stopped. (D–F) NGS analysis of complete ORF sequences of viruses that escaped from DAA combinations. Only SNP of non-synonymous mutations with frequencies >20% in pre- and/or post-treatments are shown. These SNPs showed a frequency of <20% in the original virus that underwent treatments with single inhibitors (online supplemental figures 5 and 7). The putative RASs are shown in red. The corresponding drug targets and protein specific numbers are indicated for RASs. NGS, next-generation sequencing; ORF, open reading frame; PIs, protease inhibitors; SNP, single nucleotide polymorphism.

4, a major cause of chronic liver diseases in the Middle East and North/Central Africa. Using cell-culture-adapted ED43(4a) viruses and detailed NGS combined with haplotype reconstruction analysis, we showed that complex networks of RASs and other substitutions outside the drug targets evolved under DAA treatments, which resulted in positive selection of RASs inducing high levels of resistance. We further demonstrated that glecaprevir/pibrentasvir remained efficient as a retreatment option against viruses that had escaped NS5A inhibitors or sofosbuvir, in culture. This is highly relevant, since most genotype 4 patients have been treated with an NS5A inhibitor combined with sofosbuvir.

Recently, an ED43 infectious system was reported by Watanabe et al.29 This system was developed by using substitutions previously identified in ED43 replicons,38 and the final ED43 virus could produce infectivity titres of ~3.5 log₁₀FFU/mL after 39 days of infection.29 Here, we report a different strategy, which is based on the use of substitutions conferring both viral
replication and propagation combined with high infectivity titres. Previously, we had succeeded in adapting genotypes 3a and 6a to efficiently grow in culture by initially combining substitutions identified in their corresponding C5A recombinants with changes in NS5B, which consisted of modifications of the sequence to reflect the consensus as compared with other genotypes 3a and 6a sequences, respectively.3 4 However, for the original ED43, the NS5B sequence in the clone already reflected the consensus when compared with other deposited genotype 4a sequences.39 Instead, we focused on the 10 amino acids of ED43-NS5B that differ from conserved sequences of other HCV genotype strains for which efficient infectious culture systems had been developed (online supplemental figure 1A)21 and found that a subset of these changes facilitated adaptation of the full-length ED43 clone. Our ED43cc virus was highly efficient and produced titres of ~5.1 log10FFU/mL after 6 days of infection (table 1). It should be noted that the ED43cc virus contains numerous substitutions identified through an extensive adaptation process. Thus, we cannot rule out that these adaptive substitutions could impact selection of viral resistance to DAAs. Among substitutions identified in our study and in Watanabe et al.,29 substitutions at positions 271 (V271F/G) and 2413 (D2413G) were commonly observed, suggesting that these changes might be important for the culture viability of this particular virus strain. It is worth mentioning that selection of viral resistance to DAAs and development of whole-virus vaccine candidates require highly efficient viruses. Genotype 4a ED43 replicon systems have been developed.38 40 Nevertheless, the cell culture adaptive mutations identified in this study could be an alternative to generate even more efficient genotype 4 subgenomic replicons, as recently demonstrated for strain DBN3a of genotype 3a.41 Such replicons with or without RAS, recapitulating only the intracellular replication of the virus, are useful tools to study the effect of antivirals on replication but cannot be used to understand genomic-wide mutation networks.22

Infectious cell culture systems can be important tools for vaccine development. The efficient growth of the ED43cc virus, with high infectivity titres, might permit the production of enough virus to generate inactivated whole virus vaccine candidates for preclinical testing. Alternatively, further cell culture adaptation can be achieved through serial passage of ED43cc, as described previously for another recombinant.23 Thus, the highly adapted ED43cc virus could contribute to the production

Figure 6  Efficacy of DAA combinations containing NS5A inhibitors and sofosbuvir against ED43 escape viruses. (A,B) For each DAA combination, concentrations of 5x-EC50 of NS5A inhibitors were used in combination with either 1x- or 2x-EC50 of sofosbuvir. (C,D) NGS analysis of complete ORF sequences of viruses after combination treatments. For details, see figure 5 legend. NGS, next-generation sequencing; ORF, open reading frame.
of HCV virions needed in whole virus particle vaccine studies. Nevertheless, we must acknowledge a putative influence of the cell culture-adaptive substitutions needed to grow ED43 in culture in the overall viral sensitivity to neutralising antibodies, which could influence vaccine-induced immune responses. Particularly, C458R(E2) has been shown to induce viral escape from host-immune responses. Furthermore, adaptive substitutions might also influence viral sensitivity to DAAAs and facilitate viral escape; however, as the study of HCV in culture is dependent on adaptive mutations this is a universal limitation of cell culture systems.

We showed that heterologous ED43 viral populations containing different RASs evolved under various DAA treatments, which resulted in positive selection of RASs conferring high levels of resistance (figures 2A–C, 3A–E and 4A). Also, the emergence and number of RASs did not depend only on the initial potency of the drug (EC50). At a concentration of 8xEC50, A156T/V/M emerged during treatments with grazoprevir and glecaprevir, but not with paritaprevir, suggesting higher selection pressure of glecaprevir and grazoprevir during long-term treatment despite similar potency (figure 2A vs 2B and C). For NS5A inhibitors, a similar effect was observed, since despite exhibiting similar potency, ombitasvir selected a single RAS, while ledipasvir and elbasvir selected double RAS during long-term treatment (figure 3A vs 3B and C). The observed RASs L28V and L30R+Y93H also developed in genotype 4-infected patients after treatment failures with ombitasvir and ledipasvir containing regimens, respectively.

Additionally, our data suggest that viral escape heavily relied on the fitness of the corresponding RAS-containing ED43 variant. Most NS3P RASs were detrimental for viral fitness and reverted to wildtype, as also shown previously for other genotypes in culture. For A156M, although it was maintained when introduced singly in ED43, compensatory substitutions were required to improve fitness (figure 2F). Clinically, this RAS has been detected in HCV-infected patients failing grazoprevir/elbasvir, including in genotype 4 infections.

The only NS5A RAS with a high fitness cost was NS5A-L30Δ, which could partly be compensated by NS5A-T75I (figure 3H). In line with the loss of fitness of L30Δ, we previously showed that an NS5A-P32 deletion led to low fitness, but it conferred high levels of resistance to all clinically relevant NS5A inhibitors in genotype 1 viruses. In genotype 1-infected patients failing glecaprevir/pibrentasvir, NS5A-P32 deletion is also observed.

The NS5B-S282T is usually associated with reduced fitness, thus, it is rarely detected at baseline in HCV-infected patients. However, it was reported that this RAS could be selected in culture under sofosbuvir treatment. We showed that S282T was gradually selected under sofosbuvir treatment of genotype 4a and maintained without drug pressure (figure 4A). In fact, compared with other genotypes, S282T is more frequently found in genotype 4-infected patients after DAA failures. This relative fitness advantage observed for genotype 4 could enable the S282T virus to accumulate additional substitutions for facilitating its long-term persistence after treatment failures. If the same occurs in patients, it could consequently decrease the barrier of resistance of sofosbuvir-containing regimens.

DAA combination treatments for genotype 4 have not been investigated in detail in culture. Our data showed that recommended DAA regimens were highly efficient against the original genotype 4 virus (figures 5A–C and 6A,B). Similarly, these regimens are highly efficient in the clinic. Nonetheless, viral resistance to DAA combinations remains an issue, which could hamper treatment. In Egypt, treatment failures occur in 3%–5% of patients after treatment failures. As treatment failure due to antiviral resistance is universally linked to NS5A inhibitor resistance, a valid option for a salvage DAA regimen should include the pan-genotypic NS5A inhibitor pibrentasvir, which exhibits higher potency against most NS5A-resistant variants. Indeed, we showed that glecaprevir/pibrentasvir remained efficient against the 4a virus harbouring NS5A RASs (figure 7A). This effectiveness was likely due to the high barrier to resistance of pibrentasvir, as shown by resistance profile testing (figure 3F). Moreover, the virus harbouring NS5B-S282T was also eradicated by this combination. This finding has important implications for patients failing regimens containing an NS5A inhibitor combined with sofosbuvir, which have been used for the treatment of a high number of infected individuals in Egypt. In addition, since in this study the biggest loss of fitness in the ED43 virus was only associated with the introduction of substitutions at NS3-156 and
NS5A-L30Δ, glecaprevir/pibrentasvir exhibits a high barrier to resistance. In fact, it was reported that patients failing treatment with an NS5A inhibitor and sofosbuvir were retreated with glecaprevir/pibrentasvir, resulting in >90%SVR. Importantly, most patients had baseline NS5A RASs before retreatment.37

In our study, the viruses with NS3P RASs conferring high-level glecaprevir resistance could not be eradicated by glecaprevir/pibrentasvir (figure 7A). Possible retreatment options for these viruses could include triple combinations of velpatasvir/sofosbuvir/voxilaprevir or glecaprevir/pibrentasvir with the addition of sofosbuvir and/or ribavirin, which have shown great efficacy in patients.48 49 Therefore, it would be relevant to test these combinations against PI escape viruses in future studies, using developed infectious full-length culture systems of genotypes 1a, 2a, 2b, 2c, 3a, 4a and 6a.21 28 50

In summary, we developed a highly efficient full-length HCV genotype 4a infectious culture system. Besides its use to improve our understanding about DAA resistance, this system could serve as a useful tool for the development of an HCV vaccine, which is urgently needed for control of HCV worldwide.21 Here, we performed an extensive analysis of all clinically relevant DAA that are currently being used for the treatment of genotype 4 infections. NGs and linkage analysis revealed complex dynamics operating in the selection of different RASs during treatments. The relatively high fitness and stability of NSSB-S282T observed in ED43 recombinants could have implications for the persistence of this RAS in genotype 4 infections after treatment with sofosbuvir-containing regimens. However, we showed that glecaprevir/pibrentasvir might be a promising salvage DAA regimen for the retreatment of genotype 4 after failure with sofosbuvir/NS5A inhibitor-containing regimens, as also shown recently for genotype 2 using full-length culture systems.30 The detailed understanding of the evolutionary mechanisms underlying emergence of RASs generated here can contribute to efforts directed at avoiding the emergence and transmission of DAA-resistant viruses and thus to prevent treatment failure in the future.

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