1	Genetic evidence of a functional linkage between the RNA-dependent RNA polymerase
2	and the highly structured S fragment located at the 5' end of the genome of foot-and-
3	mouth disease virus.
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## 26 Abstract

Secondary and tertiary RNA structures play key roles in genome replication of picornaviruses. 27 Complex, functional structures are particularly abundant in the untranslated regions, where 28 they are involved in initiation of translation, priming of new strand synthesis and genome 29 circularisation. The 5' UTR of foot-and-mouth disease virus (FMDV) is predicted to end in a 30 c. 360 nucleotide-long stem-loop, termed the short (S) fragment. This structure is highly 31 32 conserved and essential for viral replication, but the precise function(s) is unclear. Here, the validity of earlier structural predictions was strengthened by comparative genomic analyses 33 34 that confirmed structure conservation of S fragments from a wide range of field isolates. In addition, we used selective 2' hydroxyl acetylation analysed by primer extension (SHAPE) to 35 experimentally-determine the organisation of the structure of the genome. To examine the role 36 37 of S fragment stem-loop structure in virus replication, we introduced a series of deletions to 38 the distal and proximal regions. These truncations affected genome replication in a sizedependent and, in some cases, host cell-dependent manner. Furthermore, during passage of 39 viruses incorporating the largest tolerated deletion from the proximal region of the S fragment 40 stem-loop, an additional mutation was selected in the viral RNA-dependent RNA polymerase, 41 3D<sup>pol</sup>, which influenced the function of the enzyme. These data suggest that the S fragment and 42 3D<sup>pol</sup> interact in the formation of the FMDV replication complex. 43

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#### 45 Author summary

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Foot-and-mouth disease (FMD) is a highly contagious viral disease. It is a constant threat to the global livestock industry, as it remains endemic in many parts of the world. A deeper understanding of the molecular mechanisms of FMDV replication could help facilitate the development of future novel control measures for FMD. However, a number of features of the viral RNA genome remain poorly understood, including the role of a highly structured region,

52	termed the S fragment. In this study, we examined the structure of the S fragment and
53	introduced a series of deletions. These affected the ability to replicate. Furthermore, mutations
54	which arose when one of the deletions was incorporated into virus indicates that the S fragment
55	RNA and the RNA-dependent RNA polymerase enzyme interact during viral replication.
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57	

#### 59 Introduction

The *Picornaviridae* is a family of single-stranded positive sense RNA viruses with small 60 genomes approximately 7.5 - 8 kb in length. Typically the genome comprises a major single 61 open reading frame flanked by 5' and 3' untranslated regions (UTRs), however, additional small 62 open reading frames have been identified in some cardioviruses and enteroviruses (1, 2). The 63 large 5' UTRs present in most picornaviruses range from ~800 to ~1300 nucleotides (nts) and 64 65 have been predicted in silico by energy minimisation algorithms to comprise several distinct highly structured domains. Some of these RNA structures are common for all picornaviruses 66 67 and have been well characterised, for instance an internal ribosome entry site (IRES) responsible for the initiation of translation, which comprises ~450 nts for most picornaviruses, 68 although smaller IRES elements resembling those found in some flaviviruses were also found 69 70 (3–5). The number and organisation of these structural domains varies amongst picornaviruses. 71 For example, a small stem-loop, the cis-active replicative element (*cre*), which is essential for replication, is found in the 5' UTR of Aphthoviruses, but within the coding region in 72 Enteroviruses (6-8). All picornavirus genomes include a structured domain at the 5' end, 73 although the size, conformation and sequence varies in viruses from different genera. 74 Enterovirus 5' UTRs terminate in a ~80 nucleotide cloverleaf structure, while in Aphthoviruses 75 and Cardioviruses the 5' end forms an extended stem-loop (9-11). The size of this stem-loop 76 varies markedly; 40 nts in Hepatoviruses and ~80 nts in Cardioviruses, whereas the largest 77 (~360 nts) is present in the Aphthoviruses (12). The functions of some of these RNA structures 78 remain unknown. 79

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Here, we focus on the long 5' terminal stem-loop (termed S fragment) of foot-and-mouth disease virus (FMDV), a pathogen of cloven-hoofed animals, which belongs to the genus *Aphthovirus*. FMDV is the causative agent of foot-and-mouth disease, a highly contagious and

economically important infection posing a constant thread to the global livestock industry. In 84 85 endemic countries FMD is controlled by vaccination and by movement restrictions, while movement restrictions and slaughter were used when outbreaks occurred in non-endemic 86 countries (13). Control by vaccination is complicated by high antigenic variability and the 87 occurrence of asymptomatic carrier animals (14-16). FMDV has an 8.5 kb RNA genome 88 organised into L, P1-2A, P2 and P3 coding regions, flanked by the 5' UTR described above 89 90 and a short 3' UTR. L is a protease responsible for separating itself from the polyprotein and cleaving key cellular proteins, P1 encodes the structural (capsid) proteins, while P2 and P3 91 92 encode non-structural proteins involved in genome replication. The latter include the RNAdependent RNA polymerase (RdRp, also termed 3D<sup>pol</sup>) and the protease 3C<sup>pro</sup> as well as less 93 well-characterised proteins including 2C, a potential helicase (17-19). At approximately 1.3 94 kb, the 5' UTR of FMDV is  $\sim 1/7^{\text{th}}$  of the entire genome and is the longest 5' UTR within the 95 96 family *Picornaviridae* (10). Its sequence comprises at least five structurally and functionally distinct domains. The S fragment stem-loop is located at the 5' terminus and is followed by a 97 long poly-C-tract of variable length (about 70 to 250 nts), 2-4 tandem repeated sequences 98 predicted to form pseudoknots, the small stem-loop cre involved in uridylation of the 99 replication primer peptide VPg and finally the IRES element responsible for protein translation 100 initiation (20). The functions of several of these domains are poorly understood (21, 22), 101 102 although we have recently described the importance of a sequence in the pseudoknot region for 103 virus assembly (23).

104

105 Whilst the S fragment is known to be essential for replication, its precise function remains 106 unclear. The secondary structure of the S fragment appears to be conserved, although its 107 sequence and length varies between different clades of FMDV (10, 24). Truncated S fragments 108 have been observed for some field isolates, each time retaining its long stem-loop structure 109 (10). It has been reported that while distal portions of this stem-loop structure are not essential for genome replication, they play a role in the modulation of host-cell innate immune responses 110 (25, 26). A corresponding structure present at the 5' end of poliovirus (PV) RNA, a highly 111 structured region called cloverleaf, is involved in the switch from protein translation to RNA 112 replication, in addition to recruiting host and viral proteins which contribute to RNA 113 circularisation during genome replication (9, 27). It is possible that the S fragment in FMDV 114 115 has similar roles and it is known to interact with the 3' UTR, suggesting a role in genome circularisation (28). 116

We used selective-2' hydroxyl acylation analysed by primer extension (SHAPE) to provide 117 118 direct evidence to complement comparative bioinformatic analyses and in silico predictions of 119 the structure of the S fragment. We confirm that substantial deletions can be made to the distal region of the S fragment stem-loop without seriously compromising in vitro replication, 120 121 although host cell differences were observed. We also show that proximal deletions are less well tolerated, which correlates with our observation that the proximal part of the S fragment 122 shows higher conservation of nucleotide pairings. Furthermore, viruses reconstructed to 123 include the maximal viable proximal deletion can, during sequential passages, select a 124 compensatory mutation in the 3D<sup>pol</sup>. This mutation is located in a highly conserved position 125 126 known to interact with template RNA, and granted faster replication of a mutant replicon carrying this proximal S fragment deletion. 127

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#### 129 Material and Methods

#### 130 Cells lines.

Baby hamster kidney (BHK)-21 and Madin-Darby bovine kidney (MDBK) cells were obtained
from the American Type Culture Collection (ATCC) (LGC Standard) and maintained in
Dulbecco's modified Eagle's Medium with glutamine (Sigma-Aldrich) supplemented with
10% foetal calf serum (FCS), 50 U/ml penicillin and 50 µg/ml streptomycin as previously
described (29).

136

## 137 Plasmid construction.

138 The FMDV ptGFP replicon plasmids along with the equivalent 3D<sup>pol</sup> knock-out controls 3D<sup>pol</sup>

-GNN have already been described (30, 31).

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A variant of the ptGFP replicon was produced to allow for easy mutagenesis of the S-fragment. An *EagI* site was introduced at each end of the S-fragment by PCR mutagenesis, altering the wild-type sequence from AAAGGGGGGCATTA to AAACGGCCGATTA at the 5' end and GCGCCCGCCTTT to GCGCGGCCGTTT at the 3' end. S-fragment sequences containing deletions were chemically synthesised (Thermofisher) and inserted into the replicon vector using complementary *EagI* sites.

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Mutation of residue I189L of 3D<sup>pol</sup> in the ptGFP replicon was achieved by PCR mutagenesis
resulting in A7023C substitution. All primer sequences are available on request.

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# 151 **Protein purification.**

152 The I189L mutation was introduced into the His-tagged 3D<sup>pol</sup> expression clone pET28a-3D,

and recombinant protein was expressed and purified as previously described (32, 33).

# 155 In vitro transcription.

In vitro transcription reactions were performed as described previously (34, 35). Briefly, 5 µg 156 of replicon plasmid was linearised with AscI (NEB), purified by phenol-chloroform extraction, 157 ethanol precipitated, redissolved in RNase-free water and used in a T7 in vitro transcription 158 reaction. Reactions were incubated at 32°C for 4 hours, treated with 1.25 units of RQ1 DNase 159 for 30 minutes at 37°C and RNA recovered using an RNA Clean & Concentrator-25 spin 160 column kit (Zymo Research), following manufacturer's instructions. All transcripts were 161 quantified by NanoDrop 1000 (Thermo Scientific) and RNA integrity assessed by MOPS-162 formaldehyde gel electrophoresis. 163

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## 165 **Replication assays.**

BHK-21 and MDBK cell based replicon replication assays were performed in 24-well plates 166 with 0.5  $\mu$ g/cm<sup>2</sup> of RNA using Lipofectin transfection reagent (Life Technologies) as 167 previously described (35). For every experiment, transfection was performed in duplicate and 168 169 experiments biologically repeated as indicated. Replicon replication was assessed by live cell imaging using an IncuCyte Zoom Dual colour FLR, an automated phase-contrast and 170 fluorescent microscope within a humidifying incubator. At hourly intervals up to a defined end 171 point, images of each well were taken and used to enumerate the ptGFP positive cell count per 172 well, using our established protocols (34, 35). Data are shown at 8 hours post transfection 173 (when maximum replication was observed) on a linear scale. 174

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For ribavirin sensitivity assays, BHK-21 cells were pre-treated with ribavirin (Sigma) for 7hours prior to transfection.

## 179 SHAPE analysis.

RNA transcripts representing the entire 5' UTR of FMDV (nucleotides $1 - 1581$ ) were				
prepared as above. A sample containing 12 pmol of transcribed RNA was heated to 95°C for				
2 minutes before cooling on ice. RNA folding buffer (100 mM HEPES, 66 mM $MgCl_2$ and				
100 mM NaCl) was added to the RNA and incubated at 37°C for 30 minutes. The folded				
RNA was treated with 5 mM <i>N</i> -methyl isatoic anhydride (NMIA) or DMSO for 50 minutes at				
37°C. The chemically modified RNA was ethanol precipitated and resuspended in 0.5x Tris-				
EDTA (TE) buffer.				
Hex or FAM fluorescent primers were bound to modified RNA by heating the reaction to 85°C				
for 1 minute, 60°C for 10 minutes and 35°C for 10 minutes in a thermocycler. Reverse				
transcription was continued using Superscript III (Invitrogen) following manufacturers				
protocol with incubation at 52°C for 30 minutes.				
Post-extension, cDNA:RNA hybrids were disassociated by incubation with 4M NaOH at 95°C				
for 3 minutes before neutralisation with 2M HCl. Extended cDNA was ethanol precipitated				

and resuspended in deionized formamide (Thermo Fisher). Sequencing ladders were similarly
produced using 6 pmol of RNA with the inclusion of 10 mM ddCTP in the reverse transcription
mix and using a differentially labelled fluorescent primer (either Hex or FAM). A sequencing
ladder was combined with NMIA or DMSO samples and dispatched on dry ice for capillary
electrophoresis (Dundee DNA seq) (23).

200

NMIA reactivity was used as constraints for RNA secondary structure prediction of the S
fragment using the Vienna RNA probing package and based on sequence of the replicon used
in this study (36).

## 205 Construction of recombinant viruses.

Replicons used here are based on plasmid pT7S3 which encodes a full-length infectious copy 206 of FMDV O1 Kaufbeuren (O1K) (37). To generate infectious viral genomes the reporter 207 sequence was removed from replicons by digestion with flanking PsiI and XmaI restriction 208 enzymes and replaced with the corresponding fragment from pT7S3 encoding the capsid 209 210 proteins. Full length viral RNA was transcribed using a T7 MEGAscript kit (Thermo Fisher Scientific), DNase treated using TurboDNase (Thermo Fisher Scientific) and purified using a 211 MEGAclear Transcription Clean-Up kit (Thermo Fisher Scientific). RNA quality and 212 213 concentration were determined by denaturing agarose gel electrophoresis and Qubit RNA BR 214 Assay Kit (Thermo Fisher Scientific).

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### 216 Virus recovery.

BHK-21 cells were transfected in 25 cm<sup>2</sup> flasks with 8  $\mu$ g per flask of infectious clone-derived RNA using TransIT transfection reagent (Mirus) as described previously (35). At full cytopathic effect (CPE) or 24 hours post-transfection (whichever was earlier) cell lysates were freeze-thawed and clarified by centrifugation. The clarified lysate (1 ml) was blind passaged onto naïve BHK-21 cells and this was repeated for five rounds of passaging.

222

## 223 Sequencing of recovered virus.

Recovered viruses at passage 3, 4 and 5, were sequenced with an Illumina Miseq (Illumina) using a modified version of a previously described PCR-free protocol (38). Total RNA was extracted from clarified lysates using TRizol reagent (Thermo Fisher Scientific) and residual genomic DNA was removed using DNA-free DNA removal Kit (Thermo Fisher Scientific)
prior to ethanol precipitation. Purified RNA was used in a reverse transcription reaction as
previously described (38, 39). Following reverse transcription, cDNA was purified and
quantified using a Qubit ds DNA HS Assay kit (Thermo Fisher Scientific) and a cDNA library
prepared using Nextera XT DNA Sample Preparation Kit (Illumina). Sequencing was carried
out on the MiSeq platform using MiSeq Reagent Kit v2 (300 cycles) chemistry (Illumina) and
paired-end sequencing.

234

FastQ files were quality checked using FastQC with poor quality reads filtered using the Sickle 235 algorithm (40). Host cell reads were removed using FastQ Screen algorithm and FMDV reads 236 assembled *de novo* into contigs using IDBA-UD (41). Contigs that matched the FMDV library 237 (identified using Basic Local Alignment Search Tool (BLAST)) were assembled into 238 consensus sequences using SeqMan Pro software in the DNA STAR Lasergene 13 package 239 240 (DNA STAR) (42). Finally, the filtered fastQ reads were aligned to the de novo constructed 241 consensus sequences using BWA-MEM algorithm incorporated into Burrows-Wheeler Aligner (BWA), and mutations were visualised using the Integrative Genomics Viewer (IGV) (43-46). 242

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#### 244 Cell killing assays.

Virus titre was determined by plaque assay on BHK-21 cells as described before (47). BHK-21 cells were seeded with 3 x10<sup>4</sup> cells/well in 96 well plates and allowed to settle overnight. 21 cell monolayers were inoculated with each rescued virus at MOI of 0.01 PFU for 1 hour, 22 inoculum was removed and 150  $\mu$ l of fresh GMEM (supplemented with 1% FCS) was added 23 to each well. Appearance of CPE was monitored every hour by live cell microscopy (contrast 24 phase) using the IncuCyte S3. CPE was observed as rounding of the cells and progress of infection was monitored as a drop in confluency compared to the mock treated cells (i.e.,
treatment with uninfected cell lysate). Serial images of cells were analysed using IncuCyte S3
2018B software.

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#### 255 Sym/Sub polymerase activity assays

Sym/Sub RNA with sequence GCAUGGGCCC was synthesised (Sigma Aldrich) and 5' end labelled using  $\gamma^{32}$ P UTP (Perking Elmer) and T4 polynucleotide kinase (NEB) following manufacturer's protocol. Labelled RNA was purified by ethanol precipitation and resuspended in nuclease free water.

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261 0.5  $\mu$ M of end labelled RNA was incubated in polymerisation buffer (30 mM MOPS pH 7.0, 262 33 mM NaCl, 5 mM MgAc) and heated to 95°C for 2 minutes before chilling on ice. 2  $\mu$ M of 263 recombinant 3D<sup>pol</sup> was added to the RNA and incubated at 37°C for 10 minutes to promote 264 annealing. After annealing, 50  $\mu$ M of rNTP was added to the reaction, addition of nucleotide 265 varied depending on experiment. Aliquots were taken periodically, and reactions stopped by 266 addition of 2x TBE-urea RNA loading dye (Thermo Fisher).

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Extension products in loading dye were heated to 70°C for 5 minutes before loading on a 23% denaturing polyacrylamide gel containing 7 M urea. Samples were electrophoresed until sufficiently separated. After electrophoresis gels were fixed for 30 minutes in fixative solution and exposed onto a phosphoscreen (48, 49).

272

## 273 **Bioinformatic analysis**

Full genome sequences of 118 FMDV isolates, representing all seven serotypes(Supplementary Table S1), were downloaded from GenBank. The sequences were chosen

(based on the region encoding the VP1 protein) to represent the known genomic diversity of
FMDV across all seven serotypes. VP1 is the most variable part of the FMDV genome and
frequently used to calculate phylogenetic relationship of FMDV isolates.

279 The RNA structure of the S fragment was predicted as described before using the RNAalifold 280 program implemented in the ViennaRNA package (36, 50). For the covariance analysis, also calculated using RNAalifold program, only those isolates were included which contained 281 complete sequence at the 5' end of the genome. Data representing covariance were 282 superimposed onto the schematics of the S fragment RNA structure and visualised using Forna 283 284 tool implemented in the ViennaRNA Web Services (51). The sequence logo representing amino acid conservation within positions 185 – 194 of the 3D<sup>pol</sup> was prepared using a WebLogo 285 3.7.4 server and using 1123 FMDV 3D<sup>pol</sup> sequences obtained from fmdbase.org (an FMDV 286 287 sequence database generated by the FAO World Reference Laboratory for FMD at The 288 Pirbright Institute) (52, 53).

## 290 **Results**

## 291 SHAPE analysis of the S fragment.

Based on computational folding, such as mFOLD algorithms, the S fragment is predicted to form a single large hairpin stem-loop comprising approximately 360 nucleotides. Here, we use selective 2' hydroxyl acylation analysed by primer extension (SHAPE) to further investigate the structure of this part of the FMDV genome. This approach relies on the formation of 2'-Oadducts in single stranded and accessible regions of the RNA, which can then be detected by reverse transcription.

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RNA transcripts representing the full FMDV 5' UTR were folded before incubation with 5 mM 299 300 NMIA. The modified RNAs were then used in reverse-transcription reactions containing 301 fluorescently end-labelled primers. Reverse transcription is terminated when the enzyme reaches a nucleotide with a NMIA adduct, thus creating a series of cDNA fragments of different 302 lengths. These cDNA fragments were analysed by capillary gel electrophoresis alongside a 303 sequencing ladder to identify the sites of termination, indicating the locations of unpaired and 304 accessible nucleotides. The proximity of the poly-C-tract to the 3' end of the S fragment limited 305 306 potential primer binding sites, resulting in unreliable data for the last 60 nucleotides of the S fragment and such was excluded from the analysis (shown in grey in Figure 1A). The rest of 307 308 the S fragment was well covered and the SHAPE reactivities from 6 independent experiments 309 were used to complement the mFOLD algorithm analyses to generate a structural prediction of 310 this region (Figure 1).

In general, the SHAPE mapping data fitted well with the *in silico* predictions with the most 312 reactive residues coinciding with bulges within the predominately double stranded stem-loop 313 structure (Figure 1A). The locations of reactive residues can be seen in the NMIA reactivity 314 graph, where groups of nucleotides within bulges showed high reactivity (Figure 1B). However, 315 some predicted bulges showed little NMIA reactivity, suggesting steric hindrance possibly due 316 to higher order tertiary structure. Similarly, some predicted base-paired nucleotides were 317 318 NMIA reactive. These mostly occurred at the tops and bottom of bulges and may result from nucleotides becoming transiently available for modification during 'breathing' (54). The 5' 319 320 terminal two nucleotides (uridines at position 1 and 2) were highly reactive and therefore likely to be non-base paired. 321

322

# The effects of deletions to the distal or proximal regions of the S fragment stem-loop on replicon replication.

The SHAPE data and the structural prediction described in Figure 1 were used to design a 325 series of truncations to the S fragment, and the modified sequences were introduced into a 326 replicon. The replicon was based on an infectious clone of FMDV O1K in which the P1 region 327 328 of the genome is replaced with a ptGFP reporter (55) (Figure 2A). Deletions were designed to end at either the top or bottom of bulges, helping ensure that the stem-loop hairpin structure 329 330 was maintained. To facilitate replacement of the S fragment with modified sequences EagI 331 restriction sites were introduced at each end of the S fragment sequence. These sites required minimal modification of the original sequence and showed no predicted change in RNA 332 structure, and the modified genome sequence is hereafter referred to as WT. Addition of the 333 334 *Eag*I site at the 5' end of the S fragment resulted in a large drop in replication, however when 335 paired with the 3' EagI site, restoring base pairing, replicative potential was drastically

improved, although still below that of the original O1K WT sequence (Figure S1). Deletions 336 of 70, 148, 246 and 295 nucleotides were introduced at the distal end (top) of the S fragment 337 stem-loop (constructs named D-70, D-148, D-246 and D-295, respectively) and deletions of 338 48, 97, 195 and 271 nucleotides were made to the proximal end (bottom) of the S fragment 339 (constructs referred to as P-48, P-97, P-195 and P-271, respectively) (Figure 2B). The truncated 340 sequences were chemically synthesised and introduced into the WT replicon via the EagI sites, 341 meaning that the highly conserved 5' and 3' regions and essential UU nucleotides at positions 342 1 and 2 were maintained in all truncation mutants. The eight new replicon clones were 343 344 transcribed into RNAs and transfected into BHK-21 or MDBK cells, alongside WT and a 3D<sup>pol</sup>-GNN negative control. The 3D<sup>pol</sup>-GNN replicon cannot replicate and so the ptGFP signal 345 produced represents translation of the input RNA. BHK-21 and MDBK cells are continuous 346 347 cell lines known to support replication of FMDV: BHK-21 cells are used for FMDV vaccine production, whereas MDBK cells originate from a natural host of FMDV. Replication was 348 monitored by measuring reporter expression using an IncuCyte ZOOM live cell imaging 349 system, with analysis by our established protocols (performed 8 hours post transfection, when 350 replication has reached its peak (Figure S2)) (Figure 2). 351

352

As described previously and consistent with sequence information from natural virus isolates (10, 26), the D-70 mutation caused no significant drop in replication in either cell line. However, the replicon bearing the D-148 deletion, which removes the distal half of the S fragment stemloop, showed reduced replication in BHK-21 cells and none in MDBK cells (Figure 2C,D). Innate immunity is compromised in BHK-21 cells but intact in MDBK cells (56, 57), suggesting that the distal portion of the S fragment stem-loop plays a role in modulating a competent host immune response, in agreement with previously published data (25, 58). Replicons with the largest distal deletions, D-246 and D-295, did not replicate in either cellline.

362

While the consequences of deleting sequences from the distal region of the S fragment stem-363 loop have been reported (25), the effects of proximal deletions have not been investigated. The 364 two smallest proximal deletions (P-48 and P-97), showed modest but significant decreases in 365 replication compared to WT (1.7 and 1.4-fold, respectively) (Figure 2E,F. However, the larger 366 deletions, P-195 and P-271, completely ablated replication and reporter gene expression was 367 reduced to a level comparable to the GNN control. Interestingly, there was no significant 368 difference between the replication of the P-48 and P-97 deletions in either BHK-21 or MDBK 369 370 cells (Figure 2). Overall, the proximal part of the S fragment stem-loop appears to be more sensitive to deletion in a cell type-independent manner, with truncations as small as 48 371 nucleotides having a significant effect on replication. 372

373

374 Earlier studies indicated that the proximal part of the S fragment stem-loop is more conserved than the distal part (10). In recent years, advances in high throughput sequencing (HTS) have 375 led to a great increase in the number of FMDV full genome sequences available on public 376 domains, including sequences of viruses of SAT serotypes which were previously 377 underrepresented. This increase in the range of sequences available prompted us to revisit the 378 subject of the S fragment variability and we carried out a covariance analysis based on 118 379 380 FMDV isolates representing all seven serotypes. Figure 3 shows that there is more base-pairing conservation in the proximal than in the distal parts of the S fragment stem-loop (Figure 3). 381 382 The base-pairing conservation in the proximal part of the otherwise very variable S fragment stem-loop (low nucleotide identity of approximately 47%) suggests evolutionary pressure to 383

maintain the structure of this portion of the stem-loop and complements our observations from
the replicon experiments (Figure 2) (10, 59).

386

# 387 Viruses with proximal deletions of the S fragment replicate more slowly than WT and 388 can select for a mutation in 3D<sup>pol</sup> during serial passage.

We investigated the consequences of proximal deletions to the S fragment stem-loop further 389 by introducing them into an FMDV O1K infectious clone. This enabled evaluation of 390 replication in the context of the entire genome and investigation of the possible selection of 391 392 compensatory or adaptive mutations during serial passage. Replicons bearing proximal deletions of the S fragment stem-loop were converted into infectious clones by replacing the 393 ptGFP reporter sequence with the original O1K structural protein sequence. RNA transcripts 394 395 of the truncated S fragment clones and WT were transfected into BHK-21 cells. Supernatants containing recovered virus were harvested and used to infect naïve BHK-21 cells for 5 396 continuous passages. Recovery of infectious virus from each construct was assessed by the 397 ability of 5<sup>th</sup> passage samples to induce CPE. As expected, CPE was induced and virus 398 recovered from the WT construct. Infectious virus was also recovered from P-48 and P-97 but 399 400 no virus was recovered from either P-195 or P-271 constructs.

401

The consequences of the P-48 and P-97 S fragment deletions for the dynamics of virus replication were investigated by assessing the time taken to induce CPE in BHK-21 cell monolayers after infection with the mutated viruses or WT at a low MOI of 0.01 PFU. The integrity of the cell monolayers was assessed at hourly intervals over 63 hours using live cell imaging monitored by Incucyte S3 (Figure 4A). Both P-48 and P-97 viruses developed CPE

more slowly than WT, as anticipated from the earlier replicon experiments (Figure 4A). For
comparison, while a D-148 virus bearing a larger deletion from the distal end of the S fragment
stem-loop replicated at a slower rate than the WT virus, D-148 replicated significantly faster
than viruses carrying deletions at the proximal end of the S fragment (i.e., P-48 and P-97;
Figure 4A); further confirming replicon experiments. Ability of the viruses to form plaques
was also assessed. Both, P-48 and P-97 mutants generated significantly smaller plaques when
comparing to WT and/or D-148 mutant virus (Figure 4B).

414

Sequencing of recovered viruses revealed one isolate with mutations in 2C, which were not 415 investigated further. In addition, a single mutation in the 3D<sup>pol</sup> sequence was found in a high 416 proportion of the viral progeny of one of three P-97 viral replicates. The A7203C change 417 resulted in I189L mutation in the 3D<sup>pol</sup> protein amino acid sequence. The evolution of this 418 mutation was investigated further by sequencing viruses from earlier passages. The proportion 419 420 of the I189L mutation in the population increased from 40% at passage 3 to 51% at passage 4, 421 and 61% at passage 5, suggesting it confers a selective advantage in the context of the proximal S fragment stem-loop deletion and so is sequentially enriched. With four exceptions (two 422 isolates containing valine and two isolates containing threonine at this position) out of 1123 423 sequences of 3D<sup>pol</sup> available, all known FMDV isolates have isoleucine at this position (Figure 424 S3), making the amino acid substitution for leucine novel. . This residue is known to interact 425 with template RNA (Figure 5A), playing a crucial role during viral replication (60). No such 426 mutations were observed in the WT or the P-48 mutant viruses after passage. 427

428

# 430 Effect of the 3D<sup>pol</sup> I189L compensatory mutation on replicon replication.

The 3D<sup>pol</sup> I189L mutation was introduced into WT, GNN and P-97 replicons by site-specific mutagenesis and the consequence for replication assessed by reporter gene expression following transfection into BHK-21 cells, as above. Introduction of 3D<sup>pol</sup> I189L into the WT replicon had no significant effect on replication, however, a small but significant increase in replication was observed when it was introduced into the P-97 replicon (Figure 5B). As expected, I189L was unable to restore the activity of the GNN negative control.

437

# 438 **I189L modifies 3D**<sup>pol</sup> activity.

The influence of the 3D<sup>pol</sup> I189L mutation on polymerase function was investigated using 439 Sym/Sub assay; a highly sensitive method for interrogating polymerase activity. Sym/Sub 440 assays have been used to study PV replication and enable extension of a radiolabelled template 441 to be measured at the single nucleotide level (48). The method uses a 10-nucleotide template 442 oligonucleotide which base-pairs to create a small double stranded region with a four 443 444 nucleotide 5' overhang, thus allowing 3' nucleotide addition. In the presence of rNTPs and recombinantly expressed 3D<sup>pol</sup>, the addition of single or multiple nucleotides can be assessed 445 over time by separating input template and elongation products by PAGE (Figure 6A). Initially, 446 447 an assay using ATP only was undertaken to probe the ability of the enzyme to add a single nucleotide. WT and I189L 3D<sup>pol</sup> proteins behaved similarly in the assay with no significant 448 differences in the rate of addition of the single nucleotide (Figure 6B & C). 449

450

451 3D<sup>pol</sup> activity was examined further by including all four rNTPs into the Sym/Sub assay. The
452 design of the Sym/Sub oligonucleotide allows for elongation by a maximum of +4 nucleotides

to produce a fully double stranded product (Figure 7A). Both the WT and the I189L  $3D^{pol}$ enzymes produced the expected +4 product and, in addition, a longer product equivalent to +12 nucleotides was produced by both polymerases (as calculated by Rf value for migration travelled) (Figure 7B). This is, reminiscent of the larger products seen in recombination assays reported in the investigation of PV  $3D^{pol}$  (48, 61). Although there was no measurable difference in rate of addition of the four nucleotides by WT or mutant enzyme, I189L  $3D^{pol}$  produced the larger +12 product in greater quantity and faster (Figure 7C & D).

460

The functions of WT and I189L 3Dpol enzymes were dissected in finer detail by examining 461 incremental addition of rNTP combinations to Sym/Sub assay in the order ATP, UTP, GTP 462 and CTP. The reactions were incubated for 300 seconds (as above) before separation by 463 electrophoresis. Addition of ATP alone produced a distinct +1 band, as previously described 464 (Figure 8A). The addition of ATP and UTP together produced a +2 product in addition to lower 465 466 intensity bands corresponding to +3 and +4 nucleotides. Inclusion of ATP, UTP and GTP 467 resulted in a strong band equivalent to +3 product in addition to a minor +4 band. All four rNTPs produced a +4 band, as expected (Figure 8A). The appearance of the additional bands 468 in reactions where some rNTPs were missing indicates misincorporation into the growing 469 strand. We examined the error-rates of WT and I189L 3D<sup>pol</sup> by measuring the relative amount 470 of misincorporated +4 product and found no significant difference between the two enzymes 471 472 (Figure 8B).

473

474 Differences in misincorporation by the WT and I189L mutant polymerase were investigated
475 further by assessing the effects of ribavirin on replicons bearing either of the two enzymes as
476 increased misincorporation rates should result in increased sensitivity to the nucleoside

analogue. However, no differences were seen in the sensitivity of the WT, P-97 and P-97 3D<sup>pol</sup>
I189L replicons in the presence of up to 1 mM ribavirin (Figure 8C). Therefore, it appears that
differences in error rate were unlikely to be responsible for the production of increased amounts
of the larger +12 product by the I189L 3D<sup>pol</sup> (Figure 7C and Figure 8D).

481

#### 482 **Discussion**

483 In comparison to other picornaviruses, the 5' UTR of FMDV is uniquely large and complex, comprising several distinct structural domains, the functions of most of which are poorly 484 understood. The first c. 360 nucleotides are predicted to form a single, long stem-loop called 485 486 the S fragment (22). Although supported by functional studies, to our knowledge the secondary 487 structure of the S fragment has not been determined biochemically (25). Here, using SHAPE chemistry applied to a sequence of a well-studied FMDV isolate (O1 Kaufbeuren) (35), we 488 489 confirm that the S fragment folds into a single, long stem-loop containing several bulges formed by unpaired nucleotides. The importance of some of these bulges for viral viability was 490 determined previously (25). 491

492

493 Despite the high sequence and length variability of the S fragment observed in FMDV field isolates, most viruses maintain the full-length stem-loop (10). This suggests a strong 494 evolutionary pressure to maintain the structure. The S fragment is known to interact with viral 495 496 and cellular proteins and the 3' UTR, likely facilitating circulation of the viral genome during replication (64–66). However, there is evidence that some deletions to the distal part of the S 497 fragment are tolerated and a number of unrelated field isolates from different FMDV serotypes 498 499 (e.g., A, C and O) have been shown to carry deletions in their S fragments. These deletions 500 arose independently and range in length from few to 76 nucleotides (10, 26, 62). Unlike

501 deletions found in other parts of the FMDV genome (e.g., deletions found in the region encoding 3A), those found in the S fragment do not appear to be host specific, with viruses 502 carrying deletions within the S fragment being isolated from both cattle and pig (10, 26, 62, 503 63). A more recent study showed that deletion of 164 nucleotides from the distal part of the S 504 fragment can produce a viable, although attenuated virus, suggesting a role in evasion of innate 505 immunity, at least partially explaining the evolutionary trend to maintain the full structure of 506 507 the S fragment by field viruses (25). Here, we show that although deletion of 246 nucleotides or more from the distal region on the S fragment is not tolerated mutants carrying deletions D-508 509 70 and D-148 remain replication-competent.

510

511 The proximal part of the S fragment stem-loop shows higher sequence and nucleotide pairing conservation compared to the distal region (Figure 3). Surprisingly, deletions to the proximal 512 part of the stem-loop (i.e., up to 97 nucleotides) are also tolerated, although to a lesser extent 513 514 than distal deletions, resulting in both attenuated replicons and viruses (cf P-48 and P-97 515 mutants versus D-148 mutant, Figure 2 and 4) (10). Interestingly, while the replication of D-148 mutant replicon was more impaired in MDBK cells than in the BHK-21 cells, this cell-516 517 specific difference was not seen for the viable replicons carrying deletions to the proximal part of the S fragment stem-loop. Since, unlike BHK-21 cells, MDBK cells appear to have a 518 functional interferon pathway this might suggest that, although required for viral replication, 519 this proximal part of the S fragment stem-loop does not play a part in the evasion of the innate 520 immunity (56, 57). 521

522

523 Of three viral isolates carrying the maximum viable proximal deletion to the S fragment (P-524 97), two resulted in selective enrichment of a compensatory mutation over serial passages. In

both cases, the mutations showed a trend towards fixation. One isolate developed a number of 525 mutations in 2C (data not shown), while another developed a single I189L mutation in a highly 526 conserved site of 3D<sup>pol</sup> known to form hydrophobic contact with a nucleotide base within 527 template RNA during viral replication (60). Given the essential role of isoleucine 189 in viral 528 RNA replication, we investigated the effect of the I189L mutation on replication and RNA 529 strand synthesis. In fact, introduction of I189L into the P-97 replicon enhanced its replication, 530 531 suggesting an advantageous effect of this mutation. Although further biochemical analysis of the I189L 3D<sup>pol</sup> mutant did not show an altered rate of nucleotide incorporation when compared 532 533 to the WT polymerase, it did produce more of a 12-nucleotide long product. Such long products were observed in similar reactions for the 3D<sup>pol</sup> of PV and were the result of a nucleotide mis-534 incorporation and template switch (48, 67, 68). While the 12-nucleotide long product most 535 likely occurred due to nucleotide misincorporation, there is no evidence for altered fidelity of 536 the I189L mutant. Whether this product occurred as the result of strand slippage, or a template 537 switch remains to be investigated. Nevertheless, development of the compensatory mutation 538 over a serial passage in a virus carrying maximal viable proximal deletion to the S fragment 539 structure, and the advantageous effect of this mutation on replican replication suggests the 540 possibility of (direct or indirect) interaction between the S fragment and the viral polymerase 541 (or its precursor). Further studies are required to verify this interplay. 542

543

In conclusion, using biochemical methods, we confirmed previously proposed secondary structure of the S fragment of FMDV. Despite its higher sequence and nucleotide pairing conservation, small deletions to the proximal part of the S fragment stem-loop are viable, although resulting in attenuated replicons and viruses. An advantageous compensatory mutation within a highly conserved site of 3D<sup>pol</sup> appeared during serial passage of a viral isolate

549 bearing the largest viable mutation to the proximal part of the S fragment, suggests potential

550 interplay between the S fragment and the viral polymerase.

551

552

553

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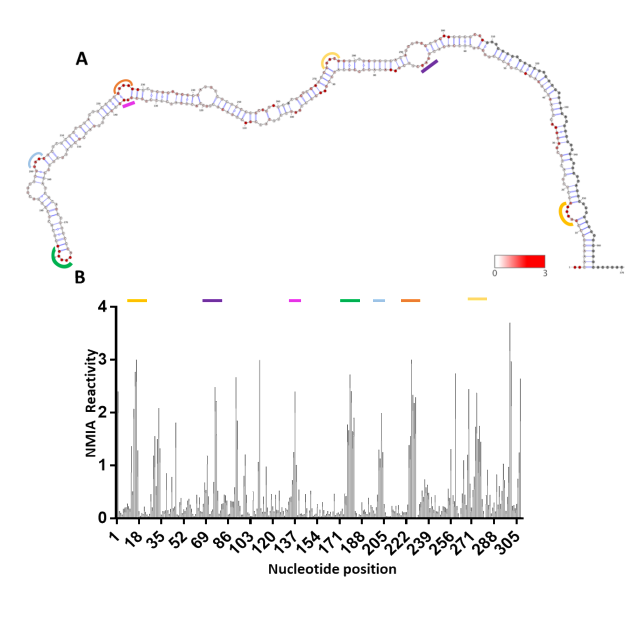
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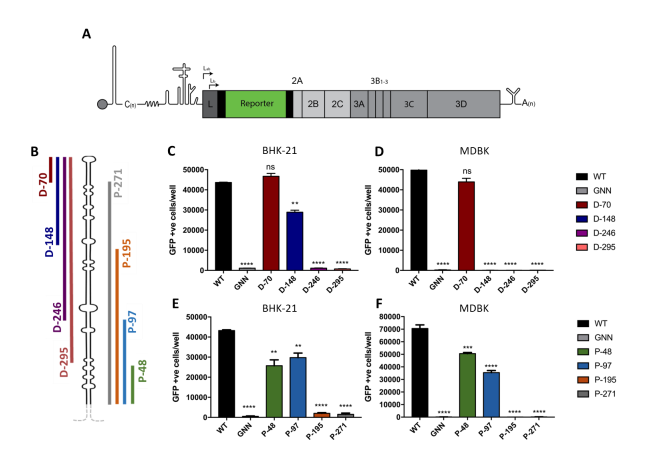


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761 Figure 1. SHAPE analysis of the S fragment. (A) Superimposition of SHAPE reactivity on the in silico predicted secondary structure WT S fragment. Secondary structure was predicted using the Vienna 762 763 RNA probing package and visualised using VARNA. NMIA reactivity is overlaid and represented on a 764 colour scale from low (white) to high (red). Nucleotides for which there is no data are represented as grey. Coloured regions corresponding to the peaks in NIMIA reactivity are shown for ease of 765 766 interpretation and localisation. (B) Individual nucleotide NMIA reactivity as analysed by SHAPE reactions and capillary electrophoresis. High reactivity indicates high probability of single-stranded 767 regions i.e. non base-paired nucleotides. Data was analysed using QuSHAPE. Corresponding coloured 768 769 regions are shown for some regions across A and B for ease of interpretation and localisation of SHAPE data to RNA structure. (n = 6, error bars represent SEM). 770

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# 775

776 Figure 2. Truncations to the distal and proximal regions of the S fragment can impair replication. 777 (A) Cartoon schematic of FMDV O1K replicon with the region encoding the structural proteins replaced 778 with a GFP reporter. (B) Schematic representation of S fragment deletions. The maintained 3' and 5' 779 proximal (P) regions are represented by the dotted grey line and was constant in all deletions (C-D) 780 Replication of replicons with 70, 148, 246 and 295 nucleotides removed from the distal (D) region of 781 the S fragment was measured following transfection into BHK-21 cells or MDBK cells. Replication was monitored by GFP expression using an IncuCyte, shown at 8 hours post-transfection alongside WT and 782 783 3D<sup>pol</sup> inactivating mutant 'GNN' acting as a positive and negative control, respectively. (E-F) Replication 784 of replicons with 48, 97, 195 and 271 nucleotides removed from the proximal region of the S fragment and measured as in (C). (n = 3, error bars represent SEM) \*\* P<0.01, \*\*\*\* P<0.0001. 785

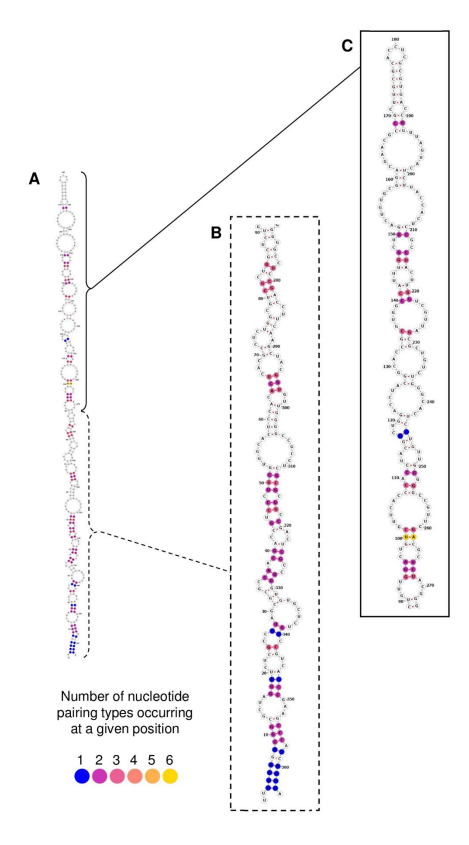
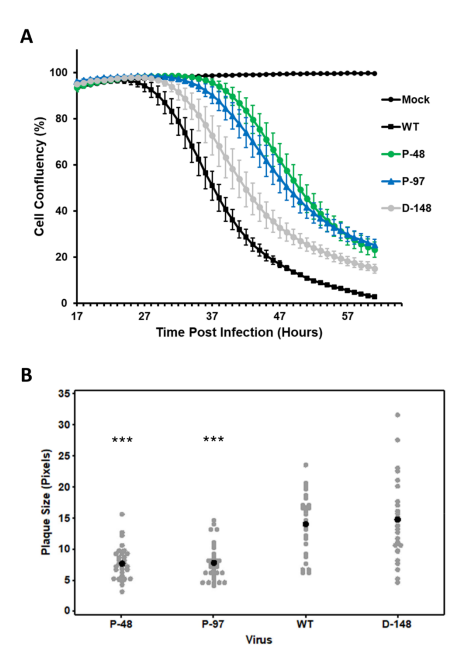


Figure 3. The extent of nucleotide and pairing conservation within the S fragment of the FMDV. (A)
 A schematic representation of conserved RNA structure of the S fragment in all 118 FMDV isolates.
 Nucleotide positions which form conserved pairing in 116/118 FMDV isolates were colour-coded
 according to number of pairing types (purple for 1 nucleotide pairing type at a given position and
 yellow for 6 nucleotide pairing types; see the colour matrix included at the left bottom corner of the

- figure). There are six possible nucleotide pairings: A-U, G-C, G-U, U-A, C-G and U-G. Nucleotide pairings
- which were not conserved in three or more FMDV isolates, remained white. Due to the length of the
- 594 S fragment stem-loop, the resolution of the image was not sufficient for detailed view of the individual
- nucleotide pairings and so the S fragment was artificially divided into two parts, proximal and distal, and the images of their schematic RNA structures enlarged (**B-C**). (**B**) The proximal part of the S
- fragment included nucleotide positions 1 to 90 and 272 to 364, **(C)** while the distal part included
- 798 nucleotide positions 90 272. Numbers represent nucleotide positions of the S fragment sequence.



802 Figure 4. Fitness of FMDV containing a truncation at the proximal end of the S fragment. (A) 803 Recovered WT, P-48, P-97 and D-148 virus populations were used to infect BHK-21 cells at an MOI of 804 0.01 PFU. CPE was monitored over time using live an IncuCyte S3 live cell imaging and is shown as a 805 drop in cell confluency; (n =3 error bars represent SEM). (B) Plague sizes of the WT, P-48, P-97 and D-148 variants grown on BHK-21 cells. Each wells containing plaques was scanned and the plaque size 806 807 was estimated in pixels. All plaques were counted to avoid bias in plaque selection. Grey dots represent individual plaques, while black dot represents the mean; \*\*\* P < 0.001 when comparing to 808 809 plaque size induced by the WT variant.



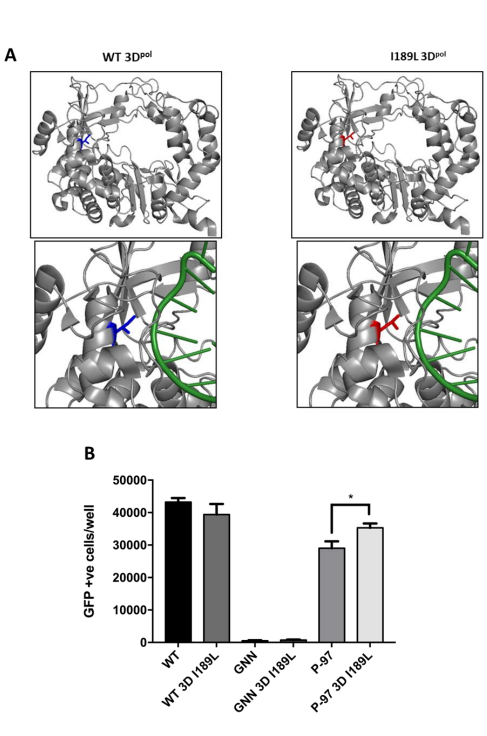


Figure 5. Compensatory mutation I189L in the structure of  $3D^{pol}$  and its effect on replication of FMDV replicons. (A) WT  $3D^{pol}$  crystal structure with I189 residue highlighted in blue, shown with (bottom, left) and without bound RNA (top, left). The I189L residue mutation was modelled onto the WT  $3D^{pol}$  crystal structure and highlighted in red (right). Structure shown with (bottom, right) and without (top, right) bound RNA. PDB file 1WNE. (B) The  $3D^{pol}$  I189L mutation was introduced into WT, GNN and P-97 replicons, termed WT I189L, GNN I189L and P-97 I189L respectively. Replicon RNA was transcribed and transfected into BHK-21 cells. WT and  $3D^{pol}$  GNN replicons were included as controls, with the latter for the level of input translation (n =3 error bars represent SEM, \* P < 0.05).

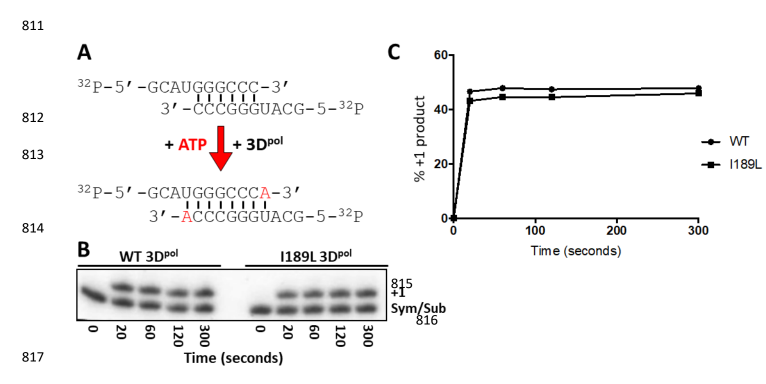


Figure 6. Single nucleotide addition to a Sym/Sub substrate by WT and I189L 3Dpol. (A) Schematic 818 of the Sym/Sub experimental protocol. Radioactively end labelled RNA oligonucleotides are annealed 819 before addition of rNTPs and recombinant 3D<sup>pol</sup>. (B) Extension of the 10mer Sym/Sub template with a 820 single nucleotide (ATP) using either WT or I189L FMDV 3D<sup>pol</sup>. Aliquots of the reactions were taken at 821 822 20, 60, 120 and 300 seconds, RNA fragments separated by electrophoresis and visualised using a phosphoimager. (C) Densitometry of +1 product is plotted as rate of addition of a single nucleotide 823 824 shown over time, %+1 product refers to the total amount of input Sym/Sub template that was 825 extended by 1 nucleotide.

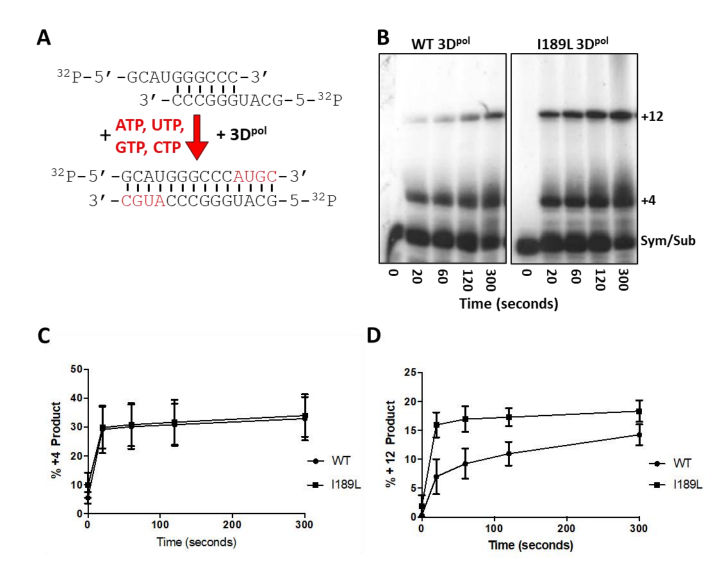
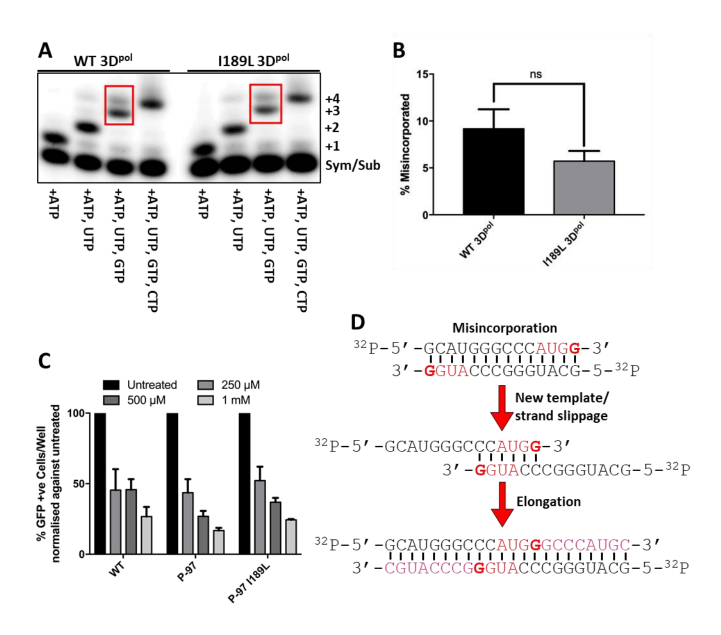


Figure 7. Sym/Sub assays showing differences between WT and I189L 3D<sup>pol</sup> in the production of a ~ +12 nucleotide product. (A) Schematic of the Sym/Sub experimental protocol using all four rNTPs. (B) Extension of the 10mer Sym/Sub template with all four nucleotides using either WT or I189L recombinant FMDV 3D<sup>pol</sup>. Aliquots of the reactions were taken at 20, 60, 120 and 300 seconds, RNA fragments separated by electrophoresis and visualised using a phosphoimager. (C) Densitometry of the production of the +4 product by either WT or I189L 3D<sup>pol</sup> over time. (D) Densitometry of the production of the +12 product by either WT or I189L 3D<sup>pol</sup>. Data are shown as the % of the input Sym/Sub template elongating to the +4 or +12 products (n = 3, error bars represent SEM). 

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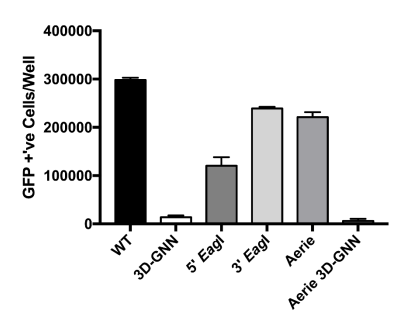


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Figure 8. Misincorporation by WT and I189L 3D<sup>pol</sup>. (A) Sym/Sub experiments were repeated with the 841 addition of either ATP, ATP and UTP, ATP and UTP and GTP or all four rNTPs together. Reactions were 842 continued for 300 seconds before electrophoresis and visualization. Misincorporation of GTP to 843 844 generate a +4 product is highlighted by the red box. (B) Densitometry of the misincorporated GTP shows no difference in the rate of misincorporation by WT or I189L 3D<sup>pol</sup>. (C) WT, P-97 and P-97 I189L 845 846 replicons were transfected into BHK-21 cells in the presence of up to 1 mM of ribavirin. Replication 847 was monitored by expression of the GFP reporter and shown here at 8 hours post-transfection. (D) Schematic for potential mechanism for the generation of a +12 product. Since misincorporation can 848 849 occur at the fourth position allowing for the addition of a second guanosine residue, if templates were to separate and reanneal by either strand slippage or annealing of a new template, extension could 850 result in the addition of 12 nucleotides (n = 3, error bars represent SEM). 851

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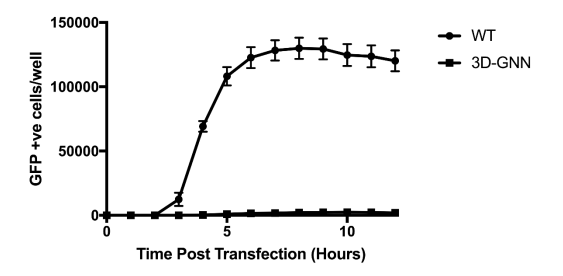
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Supplementary Figure S1. Effect of introduction of *Eagl* sites on replication. Number of GFP positive
cells/well detected at 8 hours post-transfection of BHK-21 cells. WT represents the original O1K
replicon sequence. 3D-GNN is the polymerase defective negative control. 5' *Eagl* replicon possesses
mutations to the sequence to create an *Eagl* site at the 5' end of the S fragment. 3' *Eagl* possesses
mutations to the sequence to create and *Eagl* site at the 3' end of the S fragment. Aerie shows replicon
with both 5' and 3' sites modified and Aerie 3D-GNN comprises both *Eagl* sites and 3D-GNN mutation.
n=2, error bars represent SEM.

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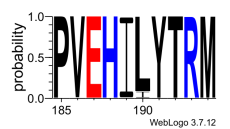


**Supplementary Figure S2. Time course of FMDV replicon replication.** Number of GFP positive 867 cells/well detected at hourly time points post-transfection of WT and 3D-GNN replicons in BHK-21

868 cells. n=2, error bars represent SEM.

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#### 870



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- 873 Supplementary Figure S3. Conservation of the isoleucine at the position 189 (I189) within the 3D<sup>pol</sup>
- 874 protein of FMDV. Sequence logo based on 1123 sequences of FMDV 3D<sup>pol</sup> available on fmdbase.org
- and prepared using WebLogo 3 server. The x-axis represents amino acids positions of the 3D<sup>pol</sup> protein
- 876 of FMDV, while the y-axis shows probability of a particular amino acid being present at a given position.
- 877 Amino acids are colour-coded according to their charge.

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# Supplementary Table S1

FMDV sequences selected from GenBank

Serotype:	Number of isolates:	GenBank accession numbers:
A	19	AY593788, MH053305, JF749843, HM854024, HQ832580, MH053306, KM268896, AY593802, KJ608371, MH053307, AY593751, AY593754, AY593761, AY593764, AY593766, AY593767, HM854022, AY593791, AY593794
Asia 1	12	AY593795, AY687334, DQ533483, DQ989306, DQ989315, DQ989319, EF149010, EF614458, HQ632774, JF739177, KM268898, MF782478
С	6	MH053308, KM268897, MH053309, AJ133357, MH053310, AJ007347
0	21	AY593819, MH053313, MH053311, MH053312, KF112885, KJ206909, HQ632769, HQ632771, KU291242, KR401154, GU384683, KF694737, AJ539140, MH053315, JX040491, MH053317, MH053318, MH053316, KJ560291, DQ404170, KU821591
SAT 1	21	AY593838, AY593845, MH053319, AY593844, JF749860, MH053321, AY593846, AY593839, AY593842, AY593841, AY593840, MH053322, AY593843, KM268899, MH053323, MH053324, MH053325, MH053326, MH053327, MW355668, MW355669
SAT2	19	MH053330, MH053332, MH053328, MH053329, JX014255, MH053333, AY593849, JX014256, MW355670 - MW355673, AY593847, MH053335, KM268900, JF749862, MH053336, MH053337, KU821592,
SAT3	20	AY593853, AY593851, MH053339, MH053340, MH053344, MH053343, AY593850, KJ820999, MH053341, MH053351, KX375417, KM268901, MH053350, MW355674 - MW355680