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Separation of foot-and-mouth disease virus leader protein activities; identification of mutants that retain efficient self-processing activity but poorly induce eIF4G cleavage

Su Hua Guan and Graham J. Belsham*

Abstract
Foot-and-mouth disease virus is a picornavirus and its RNA genome encodes a large polyprotein. The N-terminal part of this polyprotein is the leader protein, a cysteine protease, termed Lpro. The virus causes the rapid inhibition of host cell cap-dependent protein synthesis within infected cells. This results from the Lpro-dependent cleavage of the cellular translation initiation factor eIF4G. Lpro also releases itself from the virus capsid precursor by cleaving the L/P1 junction. Using site-directed mutagenesis of the Lpro coding sequence, we have investigated the role of 51 separate amino acid residues in the functions of this protein. These selected residues either are highly conserved or are charged and exposed on the protein surface. Using transient expression assays, within BHK-21 cells, it was found that residues around the active site (W52, L53 and A149) of Lpro and others located elsewhere (K38, K39, R44, H138 and W159) are involved in the induction of eIF4G cleavage but not in the processing of the L/P1 junction. Modified viruses, encoding such amino acid substitutions within Lpro, can replicate in BHK-21 cells but did not grow well in primary bovine thyroid cells. This study characterizes mutant viruses that are deficient in blocking host cell responses to infection (e.g. interferon induction) and can assist in the rational design of antiviral agents targeting this process and in the production of attenuated viruses.

INTRODUCTION
Foot-and-mouth disease (FMD) is one of the most feared diseases of livestock animals. It is caused by FMD virus (FMDV), which can infect more than 70 different species including cattle, swine and sheep [1]. The disease can significantly impact the economy of affected countries [2]. FMDV is the prototypic member of the genus Aphthovirus within the family Picornaviridae. The genome of FMDV is a positive-sense, single-stranded RNA of about 8400 nucleotides (nt) and in the virus particle the viral RNA is enclosed within a protein shell (capsid) that includes 60 copies of each of the four structural proteins VP1, VP2, VP3 and VP4 [3].

The genome includes a single, large, open reading frame (ORF), circa 7000 nt, that encodes a polyprotein (circa 2330 residues). Translation of the RNA is initiated at two different AUG codons that are 84 nt apart, yielding two alternative forms of the N-terminal component of the polyprotein, namely the leader protease (Lpro); these are termed Lab (201 amino acids) and Lb (173 amino acids) [4, 5]. The 84 nt spacer region (encoding 28 amino acids) is not required for virus viability but could have a role that is separate from its coding function, since deletion of the entire Lab sequence is not tolerated, whereas viruses lacking the 84 nt spacer region alone or the complete Lb coding sequence are viable within baby hamster kidney cells (BHK-21) [6]. Both forms of Lpro are identical at their C-termini and appear to display the same functions [7]. Lpro cleaves itself from the nascent capsid protein precursor P1-2A [8]. In addition, Lpro also induces the cleavage of the two homologues of a host cell translation initiation factor, termed eIF4GI [9] and eIF4GII [10]; these are around 46% identical. This cleavage separates the N-terminal domain of eIF4G, which is responsible for attachment to the cap-binding protein eIF4E, from the rest of the molecule. This process results in the shut-off of host cell cap-dependent mRNA translation, a common hallmark of picornavirus infection [11]. In contrast, the initiation of translation on picornavirus RNA is achieved through a cap-independent mechanism directed by an internal ribosomal entry site (IRES) located within the 5′-untranslated region (UTR) and this process is maintained...
following cleavage of eIF4G (e.g. see [11, 12]). FMDV 3C\textsuperscript{pro} has also been shown to induce cleavage of eIF4G [13], at least in cells from certain species (e.g. BHK-21). This occurs on the C-terminal side of the site generated by the expression of the L\textsuperscript{pro} and this cleavage occurs after the L\textsuperscript{pro}-mediated cleavage is complete (see [14]). Cleavage of eIF4G induced by L\textsuperscript{pro} can be observed even in the absence of virus replication, when virus protein expression is very low [13]. Besides the FMDV protease, the L\textsuperscript{pro} of other members of the genus \textit{Aphthovirus}, namely bovine rhinitis B virus (BRBV) and equine rhinitis A virus (ERAV), also induce eIF4G cleavage [15, 16].

In addition to these activities, the FMDV L\textsuperscript{pro} has also been reported to inhibit NF-\kappaB activation [17] and to exhibit deubiquitinase (DUB) activity for certain critical signalling proteins of the type I interferon (IFN) signalling pathway [18]. Modification of the L33A/L86A motif within a SAP (for SAF-A/B, Acinus, and PIAS) domain of L\textsuperscript{pro} has been reported to abrogate the DUB activity as well as its ability to block signalling to the IFN-\beta promoter [18].

L\textsuperscript{pro} is a papain-like cysteine proteinase with a flexible C-terminal extension (CTE) of 18 amino acid residues (Asp184-Lys201); the three-dimensional (3D) structure of the FMDV Lb\textsuperscript{pro} has been determined [19]. The critical active site residues are Cys51 and His148 [20, 21]. Substitution of either of these residues ablates the L\textsuperscript{pro} catalytic activities. The L/P1 cleavage site is well characterized; cleavage occurs at either a K/G or R/G junction (in different strains of FMDV, see [22]) and the cleavage can occur \textit{in trans} [7] but also probably \textit{in cis}. In contrast, the cleavage sites within the two distinct forms of eIF4G (eIF4GI and eIF4GII) are less well understood. Studies with eIF4GI identified a cleavage site produced in vitro by purified, recombinant, L protease as ANLG/RTTL [23]; however later studies on the closely related eIF4GII identified a cleavage site as LNVG/SRRS [10]. This cleavage site is not in the equivalent position to that in eIF4GI. It is possible that the cleavage of eIF4GI and eIF4GII occurs through an indirect mechanism (i.e. mediated by another protease, see [24]). Another cellular protein, Gemin5, has also been shown to be cleaved in the presence of the FMDV L\textsuperscript{pro} [25]. Whatever the mechanism, it is apparent that the recognition of the cellular protein substrates by the FMDV L\textsuperscript{pro} has to be significantly different from the recognition of the L/P1 junction within the viral polyprotein. Indeed, there is evidence that cleavage of the L/P1 junction \textit{in vitro} can be affected without impact on eIF4G processing, and vice versa. Mutations Y168F and W159A, N46Q, D136A, D136AF137A, F137AH138A, W159A, N46Q, D136A, D136AF137A, F137AH138A, H138P, V103AW105A, F151A, D164A, D163AD164AE165A and P182AD184AP187A within L\textsuperscript{pro} had rather limited or

It has been shown that the sequence between two initiation codons of L\textsuperscript{pro} is not essential for its cleavage functions [6, 7]. Therefore, the coding region of Lb was chosen as the target for mutational analysis in this study. Despite sharing protease activities, only 44% of the residues in Lb\textsuperscript{pro} were invariant among about 100 different FMDV isolates representing each of the seven different serotypes [22]. Using site-directed mutagenesis of the FMDV cDNA, we have now substituted both surface exposed and conserved residues within Lb\textsuperscript{pro} and examined the properties of the mutant proteins that were generated using transient expression assays within cells. Selected modifications were also analysed in the context of the infectious virus and the influence of these modifications on the replication of FMDV was determined.

**RESULTS**

### Screening for amino acid substitutions that differentially affect activities of FMDV Lb\textsuperscript{pro}

As indicated above, many residues within the Lb\textsuperscript{pro} are variable between different strains of FMDV. To identify residues that modify the substrate specificity of the L\textsuperscript{pro}, the conserved amino acids and the hydrophilic residues that are exposed to the aqueous solvent on the protein were located using the known 3D structure [19]. Codons corresponding to the selected residues were modified, using site-directed mutagenesis of cDNA including the L/VP4 coding region. In total, mutations encoding 51 different amino acid substitutions (out of 173 residues in Lb) were made (Fig. 1, Table 1).

As an initial screen, to analyse the properties of the mutant Lb proteins, BHK-21 cells [infected with the recombinant vaccinia virus vTF7-3, which expresses the T7 RNA polymerase (see [28])], were transfected with the mutant Lb\textsuperscript{pro} expression plasmids together with a dicistronic luciferase reporter plasmid (pFluc/EMCV/Rluc, [29]). This plasmid expresses a dicstronic mRNA that encodes both firefly luciferase (Fluc) and Renilla luciferase (Rluc). The production of the latter is dependent on the activity of the IRES while the Fluc expression is achieved by cap-dependent translation initiation. The ratio of Fluc to Rluc activity observed in the presence of the catalytically inactive C51A mutant was set to 100% and the relative levels of Fluc in the presence of the other forms of Lb were calculated and are shown in Table 1. The C51A mutant is modified at one of the two critical active site residues within L\textsuperscript{pro} [20]. Wild-type (wt) L\textsuperscript{pro}, which induces eIF4G cleavage, strongly inhibited (to about 10% of that seen with the C51A mutant) the cap-dependent expression of Fluc, as expected. In contrast, 19 mutant plasmids that were constructed in this study encoding the substitutions K38EK39E, K38EK39T, W52A, L53S, Q58EL59A, Q58E, A149D, G158AW159A, W159A, N46Q, D136A, D136AF137A, F137AH138A, H138P, V103AW105A, F151A, D164A, D163AD164AE165A and P182AD184AP187A within Lb\textsuperscript{pro} had rather limited or
no inhibitory effect on Fluc expression (similar to that observed with the mutant C51A, see Table 1). This indicates that these modifications abrogated the negative effect of Lpro on cap-dependent translation. In contrast, the other 35 mutants of Lbpro produced a strongly negative impact on Fluc expression (less than 50 % of the Fluc activity seen with the C51A mutant, see Table 1).

To confirm and extend the analysis of the functionality of the Lpro mutants, the L/VP4 cleavage activity and the induction of eIF4GI cleavage within transfected BHK-21 cells were assessed for selected variants of the FMDV Lb. For these analyses, plasmids that express, from the T7 promoter, RNA transcripts containing the FMDV IRES (to maintain expression when eIF4G is cleaved) followed by the coding region for L/VP0 with a C-terminal FLAG epitope tag (termed L/VP0-FLAG) were used (Fig. 2a). Cells, infected with vTF7-3 as above, were transfected with the wt or mutant plasmids and cell extracts, prepared 20 h later, were analysed by immunoblotting using anti-eIF4GI and anti-FLAG antibodies. The properties of the mutant Lb proteins were compared to the wt protein and to the Lpro C51A mutant. As shown in Fig. 2(b), nearly all the tested mutant proteins (other than C51A) retained the L/VP4 self-cleavage activity since the FLAG-tagged VP0 was produced in each case. One exception was the Q58EL59A mutant, which had lost this function (n.b., in principle, the L/VP0 cleavage can occur in cis or in trans in these assays since multiple copies of each mutant protein are made in the cells). Expression of the wt Lpro resulted in complete loss of the intact eIF4GI and the production of a specific cleavage product (Cp) whereas no loss of the intact eIF4GI was observed when the C51A mutant was expressed (Fig. 2c), as expected. Mutants K38E, R44A, N46Q, V103AW105G and F151A each induced quite high levels of eIF4GI cleavage but some residual intact protein was apparent. In contrast, only low levels of the eIF4GI Cp were observed with the mutants K38EK39E, R44AH138A, W52A, L53S, Q58EL59A, A149D and W159A (Fig. 2c).

To test whether the mutations within the Lb coding region affect the L/VP0 cleavage in trans, the Lpro C51A-VP0 plasmid was co-transfected with two different amounts of the Lbpro mutant plasmids. The plasmids tested were selected because each induced a very low level of eIF4GI cleavage activity in BHK-21 cells. The results (see Fig. 2d) showed that complete cleavage of the L/VP0 junction occurred with each of the mutants tested (except for C51A) when using the higher level of the plasmid. However, when a low level of the mutant Lb expression plasmid was used, then high levels of the uncleaved L/VP0 were observed for the W52A and L53S mutants and some precursor was also apparent with each of the other mutants tested as well (except for A149D). In contrast, Lpro C51A-VP0 was cleaved completely even in the presence of the low amount of wt Lpro plasmid (Fig. 2d). Thus, these results showed the W52A and L53S amino acid substitutions impaired the in trans cleavage activity to some degree but did not block it completely, while the other selected mutants retained the ability to cleave the L/VP0 junction efficiently in trans.

**Influence of the Lpro modifications on virus replication**

To determine the importance of the modifications of Lpro activity in the context of the infectious virus, selected mutations (encoding the substitutions K38EK39E, R44AH138A, W52A, L53S, A149D and W159A) were introduced into the full-length cDNA of FMDV [strain O1 Kaufbeuren (O1K)]. Each of the variants selected retained the ability to cleave the L/VP0 junction; if this activity is lost it is highly unlikely that an infectious virus could be produced since this would prevent correct production of the virus capsid protein precursor P1-2A. Using these plasmids, full-length RNA transcripts were produced in vitro and then introduced into

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**Fig. 1.** The amino acid sequence of FMDV Lpro indicating targeted residues. The asterisks indicate the key catalytic residues (C51 and H148). Underlined residues are conserved among FMDV strains [22]. Residues in bold, italics were modified in this study. The sequence represents the complete Lab form of the protease; the Lb form lacks the N-terminal 28 amino acids.
BHK-21 cells by electroporation. As judged by the production of cytopathic effect (CPE) (passage 3), rescued viruses were obtained in each case. The virus stocks were titred and the presence of only the expected mutations within the L pro coding sequence of the rescued viruses was confirmed by sequencing [following reverse transcription PCR (RT-PCR), data not shown]. The mutant viruses were then used to infect IBRS-2 cells (note in these cells, in contrast to BHK-21 cells, the FMDV 3C pro does not induce eIF4G cleavage) [14]. Consistent with the results described above, the modifications to L pro tested here had almost no impact on self-processing at the L/VP4 junction, as shown in Fig. 3(a), since VP0 (without L) was released from the polyprotein in each case.

As expected, with the wt L pro, the cellular eIF4GI was largely cleaved at 2 h post infection (h.p.i.) and was completely processed at 4 h.p.i. In contrast, in uninfected cells or within cells infected with a mutant FMDV lacking the Lb coding region (ΔLb, [6]) the eIF4GI remained intact through to 6 h.p.i. The rescued viruses with just 1 or 2 amino acid substitutions within L pro were much less efficient at producing eIF4G cleavage than the wt virus. Although some cleavage was detectable at 2 h.p.i. with mutants K38EK39E and A149D or at 4 h.p.i. (mutants R44AH138A, W52A, L53S, and W159A), intact eIF4GI persisted throughout the course of infection (up to 6 h.p.i.) for all of the mutants (Fig. 3b).

### Table 1. Effect of wt and mutant L pro expression on cap-dependent translation in BHK-21 cells

The grey blocks indicate the modifications of L pro that allow efficient cap-dependent translation of Fluc and were selected for further analysis. Data are presented as the mean±SD from the indicated number of independent experiments.

<table>
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<th>No. of observations</th>
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<tr>
<td>D164A</td>
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BH-21 cells by electroporation. As judged by the production of cytopathic effect (CPE) (passage 3), rescued viruses were obtained in each case. The virus stocks were titred and the presence of only the expected mutations within the L pro coding sequence of the rescued viruses was confirmed by sequencing [following reverse transcription PCR (RT-PCR), data not shown]. The mutant viruses were then used to infect IBRS-2 cells (note in these cells, in contrast to BHK-21 cells, the FMDV 3C pro does not induce eIF4G cleavage) [14]. Consistent with the results described above, the modifications to L pro tested here had almost no impact on self-processing at the L/VP4 junction, as shown in Fig. 3(a), since VP0 (without L) was released from the polyprotein in each case.

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### Table 1. cont.

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**Time course of FMDV replication in pBTY cells**

Primary bovine thyroid (pBTY) cells have been shown to be the most sensitive cell type for detecting field strains of all seven serotypes of FMDV [30]. However, as shown previously [6], the presence of a functional Lb protease is required for the virus to grow efficiently in pBTY cells. In this study, some CPE was observed in wt FMDV-infected pBTY cells at 10 h.p.i., and by 23 h.p.i. most of the monolayer was affected. In contrast, using the FMDV lacking the Lb coding region (ΔLb, [6]) and also the FMDVs with L pro point mutants described above, >90% of the pBTY cells were still unaffected visually by 23 h.p.i. However, in
BHK-21 cells, each of the viruses (wt and mutants) induced 80–90% CPE by 23 h.p.i. (data not shown).

To analyse, in more detail, the growth characteristics of the mutant O1K viruses, with just one or two amino acid substitutions in the L<sub>pro </sub>, we infected, at low m.o.i. (0.01 TCID<sub>50 </sub>cell<sup>-1</sup>), two pBty cells (Fig. 4a) and, in parallel, BHK-21 cells (Fig. 4b). The replication of each virus was measured using quantitative real-time RT-PCR (RT-qPCR) assays that determine the level of FMDV RNA within the samples collected at multiple time points after the start of the infection. In BHK-21 cells, each of the viruses grew with similar kinetics but it was noted that by 10 h.p.i., the level of wt FMDV RNA accumulated to higher levels than for the L<sub>pro</sub> mutants. As shown in Fig. 4(a), wt FMDV also grew well in pBty cells. However, in contrast, little or no amplification of any of the L<sub>pro</sub> mutant viruses was observed beyond 5 h. p.i. (Fig. 4a).

**DISCUSSION**

The 3D structure of L<sub>pro</sub> includes an active site with the two key catalytic residues, C51 and H148, on the opposite sides of a cleft separating two domains, an α-helical
domain and β-sheet domain including the C-terminal region with a unique CTE (Fig. 5a). Residues 183–195 of the CTE have been shown to be important for the interaction with eIF4GI [27]. The Lbpro with only a partial CTE (residues 183–195) retained the activities of wt Lbpro, whereas deletion of the entire CTE reduced the ability to induce cleavage of eIF4G. Modification of residue C133, which is structurally close to the CTE, produced similar results in vitro [31]. Further study indicated that the conserved residues D184 and E186 within the CTE were important for this loss of function [32].

Some studies have indicated that the cleavage of eIF4G induced by the FMDV Lpro requires the interaction of eIF4G with the cap-binding protein eIF4E [33]; this translation initiation factor binds to the N-terminal region of eIF4G. Recent studies have shown that interaction of Lbpro with a short fragment of eIF4GII requires the presence of eIF4E [34], within a heterotrimeric complex, and that residue C133 and the CTE of Lbpro are required for the interaction with eIF4E. However, these experiments were based on the use of rabbit reticulocyte lysate (RRL) in vitro translation system and were focused on the CTE and residues around the active site. Here we have investigated the conserved and charged amino acid residues distributed across the surface of the Lpro structure and generated an insight into the roles of these residues in producing cleavage of eIF4G within cells.

Compared to other papain-like proteases, Lpro displays a deeper and narrower cleft (S1 subsite) with the active site at the bottom, the loop preceding the central helix α1 (N50–E61) on one side, and the β-turn connecting strands β3 and β6 (F137–S156) on the other [19]. Among the residues involved in this subsite, the amino acid substitutions W52A, L53S, Q58EL59A, and A149D each led to the loss of eIF4G cleavage activity (see Fig. 2c). The double mutant Q58EL59A also failed to release Lpro from the polyprotein. These key residues are all located in the vicinity of the active site (see Fig. 5a) and may contribute to the substrate...
recognition since (apart from the double mutant) they do not affect the L/P1 cleavage activity; thus they do not affect the catalytic activity of the enzyme per se. It is noteworthy that these residues are highly conserved in FMDV [22] and they (except W52 and A149) are also present in the equivalent positions within the L protease of ERAV but not in the L\[^{pro}\] of equine rhinitis B virus (ERBV) [16]. Unlike the ERBV L\[^{pro}\], which is unable to shut down cellular cap-dependent translation, the ERAV L\[^{pro}\] and FMDV L\[^{pro}\] both induce cleavage of eIF4G1 at very similar or identical positions [16]. Taken together, these observations are consistent with a significant role for these residues, around the S1 subsite, in the induction of eIF4G cleavage by L\[^{pro}\].

Residues W52 and A149 are also involved in the S2 subsite, a deep hydrophobic pocket [19]. A comparison between the monomeric Lb\[^{pro}\] L200F and sLb\[^{pro}\] (lacking the last six residues of Lb\[^{pro}\]) showed that residue W52 displayed a significant backbone amide shift when the C-terminal residues were removed [35], suggesting an interaction between W52 and the CTE. Moreover, W52 is adjacent to the catalytic residue C51. These properties may contribute to the loss of eIF4G cleavage observed with the mutant W52A; however the L/P1 cleavage activity was retained by this mutant so the enzyme is clearly still active (c.f. the C51A mutant). The conserved residue A149 is adjacent to the other key catalytic residue H148. Modification of A149 to the charged Asp (A149D) resulted in delayed eIF4G cleavage (Fig. 3b). This modification changes the hydrophobic environment and the acidic side chain of Asp may modify the space for the substrate to penetrate into the S2 pocket. Indeed, clear chemical shifts were observed for residues involved in the S2 pocket in the presence of a 12 amino acid peptide derived from eIF4G1 [36].

In this study, we have also shown the importance of residues K38, K39 and R44 for the Lb\[^{pro}\] -induced cleavage of eIF4G. It seems that an acidic side chain (as in Glu, E) is not tolerated in place of K38 or K39. The double substitutions K38EK39E significantly impaired the cleavage of eIF4G (Fig. 3b, c). However, this was not the case for the single-point mutants K38A and K39A (n.b. these changes introduce a neutral amino acid rather than an acidic residue), which both significantly inhibited the expression of Fluc, reflecting loss of intact eIF4G (Table 1). It could be that the double substitutions K38AK39A would have a similar effect to K38EK39E on eIF4G cleavage induced by Lb\[^{pro}\]. However, the mutant K38AK39T that has lost both the basic side chains impaired cap-dependent translation (as monitored by Fluc expression, see Table 1). The substitution K39E alone or in K38EK39T has a clear negative impact on Fluc expression (Table 1) and induced eIF4G processing albeit these effects were weaker than with the wt L\[^{pro}\]. Therefore, it appears that the double substitutions in K38EK39E change the local ionic environment leading to a reduction in the L\[^{pro}\]-induced eIF4G cleavage activity.

Amino acids R44 and H138 are conserved, charged residues within Lb. It is apparent from the 3D structure that, like K38 and K39, the side chain of R44 protrudes away from the globular domain of L\[^{pro}\] (Fig. 5b) and none of these residues appears to interact with the CTE [19]. Instead, these basic side chains are well positioned to interact with

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**Fig. 5.** Location of residues affecting induction of eIF4G cleavage. (a) 3D structure of Lb\[^{pro}\] [19] (PDB ID code 1QOL, annotated using PyMol). The two critical active site residues (C51 and H148) are marked in red. Residues coloured yellow were demonstrated in this study to play an important role in eIF4G1 cleavage. (b) Hydrophobic region. Residues in purple form a hydrophobic region with residue W159.
residues from another protein. However, the interaction may be not very strong or at least involve more than one residue. It is apparent that single mutants K39E and R44A did not reduce the eIF4G cleavage as much as the double mutants K38EK39E and R44AH138A. These results are consistent with previous observations that the mutant H138L was reduced in its ability to induce eIF4G cleavage but did not affect self-processing activity [21].

Residue W159 was investigated here due to its conservation in FMDV, BRBV and ERAV (data not shown). The modification W159A greatly decreased the eIF4G cleavage induced by Lbpro compared to wt. The cleaved product could only be observed from 6 h.p.i. in virus-infected cells (Fig. 3b). The side chain of W159 together with residues L134, F137 and V154 form a very hydrophobic region within the structure of Lbpro (Fig. 5b). Studies using NMR showed amide chemical shift changes for these residues in the presence of an eIF4GI-derived 12mer peptide [36], suggesting the possible involvement of these residues in substrate interaction. Indeed, as shown here, the substitutions in mutant D136AF137A resulted in the high cap-dependent expression of Fluc (Table 1) (i.e. reflecting poor eIF4G cleavage).

To facilitate the analysis of L/P1 cleavage activity, a FLAG-epitope tag was introduced at the C-terminus of the coding region for the VP2 capsid protein. Thus, the cDNA fragment corresponding to the IRES followed by the L/VP4/VP2 (L-VP0) coding region was amplified using primers SG1 and SG42 with the plasmid pT7S3 (as above) as template. The fragment (2007 bp) was digested with EcoRI and BamHI and ligated into similarly digested pGEM3Z, generating pSG42. Individual mutations within the Lb coding region were introduced by site-directed mutagenesis and resulted in the following substitutions: C51A, W52A, L53S, K38EK39E, Q58EL59A, A149D and W159A (following the numbering system for the Lab form of Lbpro). Briefly, primers 10PPN36 and SG57 (Table S1), and the derivatives of pSG2 carrying the mutations were used as the templates to generate megaprimers for the second round PCRs. The template in the second round of PCR was pSG42. The final constructs were sequenced throughout the L/VP2 coding region using primers SG1 and SG42.

### Transient expression assays

Plasmid DNAs were transfected, using FuGene 6 (Promega) as described by the manufacturer, into BHK-21 cells (35 mm wells, circa 90 % confluent) that had been infected with the recombinant vaccinia virus vTF7-3 [28] which expresses the T7 RNA polymerase essentially as described [29]. For the luciferase assays, the pSG2 variants with mutations in Lbpro (30 ng) were co-transfected with pPluc/EMCV/Rluc (2 µg) [this expresses both firefly and Renilla luciferase; [29]). Inhibition of cap-dependent translation was monitored by measuring Fluc expression relative to the Rluc expression. The cap-independent expression of the Rluc is directed by the EMCV IRES. At 20 h.p.i., cell lysates were prepared using the Rluc assay lysis buffer for Fluc and Rluc assays, or using buffer C [20 mM Tris-HCl (pH 8.0), 125 mM NaCl, and 0.5 % NP-40] for Western blot analysis, and extracts were clarified by centrifugation at 18 000 g for 10 min at 4 °C.

### Luciferase reporter assays

Fluc and Rluc activities were determined using the firefly and Renilla reporter assay systems (Promega), respectively, according to the manufacturer’s protocol. The Fluc activities were normalized to the measured Rluc activities and the data were collected from at least three independently conducted experiments.

### Immunoblotting

Immunoblotting was performed according to standard methods as described previously [40]. Briefly, samples were...
mixed with 2× Laemmli sample buffer (containing 25 mM DTT), separated by SDS-PAGE (4–15% polyacrylamide from Bio-Rad), and transferred to PVDF membranes (Millipore). After blocking in 5% nonfat dry milk and 0.1% Tween 20 in PBS, membranes were incubated with primary antibodies diluted in the same buffer. The following primary antibodies were used as described [41]: monoclonal mouse anti-FLAG 1:3000 dilution (Sigma); polyclonal rabbit anti-eIF4GI (1:2000 dilution, kindly provided by N. Sonenberg, McGill University, Montreal, Canada, diluted in 5% BSA); and anti-FMDV VP2 1:2000 (mouse monoclonal antibody 4B2, kindly provided by L. Yu, Harbin, P. R. China). Bound proteins were visualized using the appropriate secondary, horseradish peroxidase-conjugated antibodies (Dako). Detection was achieved using chemiluminescence reagents (Pierce ECL; Thermo Fisher Scientific) and images were captured using a Chemi-Doc XRS system (Bio-Rad).

**Rescue of viruses from full-length FMDV cDNA**

To introduce mutations into the FMDV full-length cDNA, the KpnI fragment from the pSG2-derived plasmids was ligated to KpnI-digested pT7S3-NheI [42]. The presence of the mutations was confirmed by sequencing using primers 10PPN36 and SG57 (Table S1). The plasmids containing the wt and mutant full-length FMDV cDNAs were linearized by digestion with HpaI, then purified (Fermentas PCR purification kit), and in vitro transcribed using T7 RNA polymerase (Megascript kit; Ambion) as described by the manufacturer. The RNA transcripts were analysed using agarose gel electrophoresis and then introduced into BHK-21 cells by electroporation as described previously [43] and incubated at 37 °C for 48h. The viruses were harvested following freezing, and amplified in one or two subsequent passages (P2 or P3) in BHK-21 cells. The appearance of cytopathic effect (CPE) was monitored during each passage. After these passages, when CPE was apparent, viral RNA was extracted (QIAamp RNA Blood Mini kit; Qiagen) and reverse transcribed using Ready-To-Go Prime First-Strand Beads (GE Healthcare, Life Sciences), and amplicons (900 bp) corresponding to the L/VP4 coding region were amplified by PCR using primers 10PPN36 and SG57 (Table S1). The products were sequenced to confirm the presence of the introduced mutations. Note that control reactions, lacking reverse transcriptase, were used to show that the PCR products obtained were derived from the viral RNA template. Viral titres were determined, as 50% tissue culture infectious doses (TCID\textsubscript{50}), by titration in BHK-21 cells, according to standard procedures [44].

**Infection and virus growth kinetics**

BHK-21, IBRS-2 and pBTY cells were grown in Dulbecco’s modified Eagle’s medium supplemented with 5% (BHK-21 and IBRS-2) or 1% (pBTY) foetal bovine serum. Cells were infected with either wt or mutant FMDV. After a 1 h adsorption step, additional medium was added and the infection was left to progress and stopped at pre-selected times. For immunoblotting analysis, IBRS-2 cells were infected at an m.o.i. of 2. Cells were harvested at selected times with buffer C (as above).

For virus growth kinetics, BHK-21 and pBTY cells were infected at an m.o.i. of 0.01 at 37 °C. Infections were stopped at selected times by freezing. Viral RNA was isolated using a robotic protocol (MagNA Pure LC Total Nucleic Acid Isolation kit, Roche) as described by the manufacturer and the RNA samples were eluted in 50 μl of water. The production of FMDV RNA was determined using RT-qPCR as described previously [45, 46].

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**Conflicts of interest**

The authors declare that there are no conflicts of interest.

**Ethical statement**

In these studies, no experiments were performed on humans or animals and thus no ethical approval was required.

**References**


