Murine norovirus virulence factor 1 (VF1) protein contributes to viral fitness during persistent infection

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Abstract
Murine norovirus (MNV) is widely used as a model for studying norovirus biology. While MNV isolates vary in their pathogenesis, infection of immunocompetent mice mostly results in persistent infection. The ability of a virus to establish a persistent infection is dependent on its ability to subvert or avoid the host immune response. Previously, we described the identification and characterization of virulence factor 1 (VF1) in MNV, and demonstrated its role as an innate immune antagonist. Here, we explore the role of VF1 during persistent MNV infection in an immunocompetent host. Using reverse genetics, we generated MNV-3 viruses carrying a single or a triple termination codon inserted in the VF1 ORF. VF1-deleted MNV-3 replicated to comparable levels to the wildtype virus in tissue culture. Comparative studies between MNV-3 and an acute MNV-1 strain show that MNV-3 VF1 exerts the same functions as MNV-1 VF1, but with reduced potency. C57BL/6 mice infected with VF1-deleted MNV-3 showed significantly reduced replication kinetics during the acute phase of the infection, but viral loads rapidly reached the levels seen in mice infected with wildtype virus after phenotypic restoration of VF1 expression. Infection with an MNV-3 mutant that had three termination codons inserted into VF1, in which reversion was suppressed, resulted in consistently lower replication throughout a 3 month persistent infection in mice, suggesting a role for VF1 in viral fitness in vivo. Our results indicate that VF1 expressed by a persistent strain of MNV also functions to antagonize the innate response to infection. We found that VF1 is not essential for viral persistence, but instead contributes to viral fitness in mice. These data fit with the hypothesis that noroviruses utilize multiple mechanisms to avoid and/or control the host response to infection and that VF1 is just one component of this.

INTRODUCTION
With figures upwards of 19 million cases annually, human norovirus (HuNoV) is increasingly becoming the leading causative agent of gastroenteritis [1–4]. Unlike rotaviruses, there are currently no vaccines available against HuNoV, largely due to long-standing difficulties in culturing the virus in vitro and the lack of a robust animal model that together have impeded the identification of the correlates of protection. With the identification of murine norovirus (MNV) in 2003 [5], the gap in knowledge on the norovirus lifecycle has been steadily shrinking [6, 7]. MNV has been isolated from wild and laboratory mice, with some research facilities reporting 50–70% seroprevalence [8, 9]. The establishment of reverse genetics systems, the ability to replicate in cultured cells with a tropism for macrophages and dendritic cells, and the availability of a robust homologous small animal model have made MNV a model of choice for studying norovirus pathogenesis [5, 10–12]. MNV-1 was initially discovered as an acute but lethal infection in immunocompromised (STAT1 knockout) mice [5]. Many other strains have since been identified, showing significant differences in their phenotype both in vitro and in vivo [5, 8, 13, 14]. While some strains cause an acute infection that is cleared within 7 days in immunocompetent mice, other strains are able to persist for up to 9 months [15]. MNV has therefore proven invaluable as a tool to dissect host and viral factors that contribute to viral persistence and
Host innate immunity is essential in controlling viral replication in a lethal model of MNV-1 infection [19]. Immunocompetent mice and contributed to viral virulence revealing its function as an innate immune antagonist [19]. To disrupt the innate immune response and establish a persistent infection in vivo [21]. Similarly, cyctic choriomeningitis virus infection required the immune response to establish a persistent lympho - 

during infection can dictate the outcome of a persistent infection, placing substantial pressure on viruses to adapt and subvert relationship between the host and pathogen determines whether these defences through a myriad of mechanisms. This dynamic for viral replication in cell culture but as VF1 expression was yet to be fully elucidated. In vivo, VF1 expression was found to contribute to replication during acute infection in immunocompetent mice and contributed to viral virulence in a lethal model of MNV-1 infection [19]. Host innate immunity is essential in controlling viral replication, placing substantial pressure on viruses to adapt and subvert these defences through a myriad of mechanisms. This dynamic relationship between the host and pathogen determines whether a persistent infection can be established, or if an acute infection is rapidly cleared. Recent studies have shown that early events during infection can dictate the outcome of a persistent infection and the ensuing immune response, as escape of the innate immune response was required to establish a persistent lymphocytic choriomeningitis virus infection in vivo [21]. Similarly, the hepatitis C virus (HCV) protein NS3-4A ablates signalling through RIG-I and MDA5 by cleaving MAVS, permitting HCV to disrupt the innate immune response and establish a persistent infection [22].

In this study, we sought to dissect a potential role for VF1 in viral persistence. To this aim, we examined how the lack of VF1 expression affected the ability of MNV to establish persistent infections in immunocompetent mice. While the introduction of a single stop codon in ORF4 to prevent VF1 expression led to significant reduction in shed virus, expression was restored in infected mice, concomitant with restoration of VF1 expression by the introduction of genetic mutations. Following the introduction of three stop codons in VF1 to suppress this reversion, a consistently reduced level of shedding was observed throughout a 3 month persistent infection in vivo. Our data confirm that while VF1 is not required for persistence, it nevertheless contributes to viral fitness during persistent MNV infections.

**METHODS**

**Cell culture and plasmid constructs**

The murine microglial BV-2 cell line was kindly provided by Jennifer Pocock (University College London). The cells were maintained in Dulbecco's modified Eagle's medium (DMEM; Gibco) supplemented with 10% FBS (Biosera), penicillin (100 units ml⁻¹), streptomycin (100 µg ml⁻¹), 2 mM l-glutamine and 0.075% sodium bicarbonate (Gibco). Murine macrophage RAW264.7 cells and immortalized murine embryonic fibroblasts (MEFs) were maintained in DMEM supplemented with 10% FBS, penicillin (100 U ml⁻¹) and streptomycin (100 U ml⁻¹) at 37°C with 5% CO₂. Baby hamster kidney cells expressing T7 polymerase (BSR-T7, kindly provided by Karl-Klaus Conzelmann, Ludwig Maximillian University) were maintained in DMEM, supplemented with 10% FCS, penicillin (100 units ml⁻¹), streptomycin (100 µg ml⁻¹) and 0.5 mg/ml G418. Bone marrow-derived macrophages (BMDMs) were prepared from BALB/c mice as previously described [23]. The previously described full-length MNV-3 cDNA clone (pT7:MNV-3) [11], which contains the genome under the control of a T7 promoter, was used for this study. cDNA clones of VF1-mutant viruses that included either a single or a triple stop codon in ORF4 were generated by PCR mutagenesis using KOD hot start DNA polymerase (Novagen). The MNV-3 M1 mutant was generated by inserting a stop codon, through the change of T to A at position 5118. The MNV-3 M3 mutant was generated by inserting three stop codons by the introduction of the mutations T5118A, C5198A and T5207A. In all cases, the VF1 mutations did not affect the protein coding sequence in the VP1 reading frame.

**Immunoblotting**

Unless otherwise stated, BV-2 cells were infected with MNV at an m.o.i. of 5 TCID₅₀ per cell, and 12 h post-infection the cells were harvested and lysed in RIPA buffer [50 mM Tris- 

HCl (pH 8.0), 150 mM NaCl, 1 mM EDTA, 1% Triton X-100, 0.1% SDS]. Cell lysates were resolved by SDS-PAGE and then probed using anti-GAPDH (AM4300; Ambion), anti-NS7, anti-ATP5α (ab14748; Abcam), or a mouse monoclonal anti-VF1 (4K5; Abmart) antibodies. The anti-NS7 antibody was described previously [24]. Goat anti-rabbit (sc-2030; Santa
Fig. 1. In vitro characterization of MNV-3 VF1. (a) A schematic representation of the MNV genome, showing the four ORFs that encode the non-structural proteins NS1/2–7, VP1, VP2 and VF1. (b) Western blot analysis of VF1 expression following a 12 h infection with MNV-1 (1-WT) and MNV-3 (3-WT) in RAW264.7 cells (m.o.i.=5). Mock inoculated samples were treated with Dulbecco’s modified Eagle medium (DMEM). (c) An alignment of VF1 protein sequences from MNV-1 (acute, lab- adapted strain), MNV-3 (persistent, lab- adapted strain), CR6 (persistent, lab- adapted strain) and WM6 (persistent, wild strain). VF1 encoded by MNV-3 and MNV-1 are 89% identical. (d) Confocal microscopy of MEFs transfected with plasmids expressing enhanced green fluorescent protein (EGFP), or EGFP fused to the N or C terminus of MNV-3 VF1. Green indicates EGFP, red indicates MitoTracker and blue denotes DAPI. (e) Biochemical fractionation of BV-2 cells infected with MNV-1 (1-WT) and MNV-3 (3-WT) (m.o.i.=5). Mock inoculated cells were treated with DMEM. Lysates were collected 12 h post-infection and mitochondrial (mito.) and cytoplasmic (cyto.) fractions were enriched as described in the text.
The homogenate was extracted and added to 250 µl of 70% fialified across the VF1 coding region prior to use.

Reverse genetics

VF1 mutant viruses were rescued using the DNA-based reverse genetics system as previously described [10, 11]. Briefly, BSR-T7 cells were infected with recombinant fowlpox virus expressing T7-polymerase at an m.o.i. of ~5, and then transfectioned with 1 µg of the respective cDNA clone using Lipofectamine 2000 (Invitrogen). At 48 h post-transfection, cells were freeze-thawed and clarified lysates were titrated by measuring the 50% tissue culture infectious dose (TCID₅₀), measured by looking for signs of a cytopathic effect at 4 days post-infection. The remaining lysates were used to infect BV-2 cells at an m.o.i. of 0.05 TCID₅₀ per cell to generate high titre passage 1 or 2 stocks. In all cases, stocks were sequence verified across the VF1 coding region prior to use.

In vivo studies

Three- to 4-week-old male C57BL/6 mice were inoculated with 100 µl of virus containing 1000 TCID₅₀ by oral gavage. Mock-infected mice were inoculated with DMEM. Mice were weighed, and faecal and tissue samples were taken at various times post-infection. Mock-infected mice were euthanized, and tissue samples harvested on day 29 only. Tissue samples included mesenteric lymph nodes (MLN), spleen, duodenum, ileum, caecum and colon, which were stored in RNA Later (Ambion). This work was carried out in accordance with regulations of The Animals (Scientific Procedures) Act 1986 [25] and the ARRIVE guidelines [26]. All procedures were approved by the University of Cambridge Animal Welfare and Ethical Review Body (AWERB) and the UK Home Office and carried out under the Home Office project licence PPL 70/7689.

RNA extraction and qRT-PCR

Faecal pellets were accurately weighted, then resuspended and homogenized in PBS at a final concentration of 100 mg of faecal pellet ml⁻¹ PBS. Following centrifugation for 5 min at 4000 g, viral RNA (vRNA) was extracted from 100 µl of supernatant using the GenElute mammalian total RNA kit (Sigma-Aldrich), according to the manufacturer’s instructions.

Tissue samples (including MLN, spleen, duodenum, ileum, caecum and colon) were sliced into 1 mm squares with sterile disposable scalpels. Sterile silica beads and 250 µl lysis buffer were added and tissues were homogenized using the Fast-Prep-24 homogenizer (MP Biomedicals) at 4 m s⁻¹ for 1 min. The homogenate was extracted and added to 250 µl of 70% ethanol, and RNA was extracted using the GenElute mammalian total RNA kit (Sigma-Aldrich).

Extracted vRNA from faecal and tissue samples was reverse transcribed using M-MLV RT (Promega) and random hexamer primers (Roche). Viral loads were determined by TaqMan qPCR, as previously described [11]. Briefly, cDNA was added to a mastermix containing 2× Precision MasterMix (Primer Design) and primers (sense: 5’-CCGCAGGAACGCTACGAG-3’, anti-sense: 5’-GGCTGAATGGGGACGGCCTG-3’, and TaqMan probe: 5’-ATGAGTGATGGCGCA-3’). Reactions were subjected to 50 cycles consisting of denaturation at 95 °C for 15 s, followed by annealing and elongation at 60 °C for 1 min, using a ViiA7 qPCR machine (AB Applied Biosystems). Viral genome copy number was calculated by interpolation from a standard curve fit generated from known quantities of vRNA.

Titration of infectious virus from faecal samples

Viral titres were determined with an endpoint dilution assay. Faecal samples collected from inoculated mice were accurately weighed and homogenized in PBS at a final concentration of 100 mg ml⁻¹. Resuspended faecal material was then centrifuged at maximum speeds for 5 min and 100 µl of supernatant was extracted and subsequently centrifuged at maximum speeds for a further 5 min to remove any traces of faecal debris. Purified samples were titrated by TCID₅₀ in BV-2 cells as above.

Caspase activation assay

The activation of caspases 3 and 7 was determined using the Caspase-Glo 3/7 assay kit (Promega), according to the manufacturer’s protocol.

ELISA

The IFN-β ELISA protocol has been previously described [27]. A monoclonal rat anti-mouse IFN-β antibody (sc57201; Santa Cruz) was used for capture, a polyclonal rabbit anti-mouse IFN-β antibody (32401-1; R and D Systems) was used for detection, and a goat anti-rabbit-HRP (7074; Cell Signaling Technology) was used as a secondary antibody. Recombinant mouse IFN-β (12400-1; R and D Systems) was used as a standard.

Confocal microscopy

MEFs were seeded on cover slips overnight and were transfected with the appropriate plasmids using Lipofectamine 2000 (Invitrogen), according to the manufacturer’s protocol. The cells were incubated for 30 min in labelling solution (MitoTracker Red CMXRos; Invitrogen) 24 h after transfection, washed three times with PBS-T, fixed in 4% paraformaldehyde, washed three times with PBS-T and then washed twice in PBS. The cover slips were then mounted on slides with Mowiol (Sigma) containing the DAPI nuclear stain. The cells were visualized on a Leica SP5 confocal microscope (Leica Microsystems), and data were analysed with ImageJ (National Institutes of Health).

Sequence alignment

Alignment of VF1 protein sequences was carried out using Clustal Omega [28], with sequences from MNV-1.CW1 (accession number: DQ285629.1), MNV-3 (accession number:
cases, error bars indicate standard error of the mean. In all studies, and confocal imaging of GFP-tagged VF1 in transfected cells [19]. Given the high sequence similarity between VF1 proteins from different MNV strains (Fig. 1c), with 89% similarity between VF1 encoded by MNV-3 and MNV-1, we hypothesized that VF1 expressed in MNV-3-infected cells will also localize to mitochondria. To test this, N-terminal or C-terminal MNV-3 VF1 GFP-fusions were constructed and transfected into MEFs. As shown in Fig. 1d, both the N- and the C-terminal MNV-3 VF1 GFP-fusions demonstrated a mitochondrial localization pattern, akin to that observed for MNV-1 [19]. To examine the localization of MNV-3 VF1 expressed in the context of viral infection and to rule out any artefacts due to fusion to GFP, we carried out biochemical fractionation on infected murine macrophage cultured cells, that it localizes to the mitochondria, and that it antagonizes the induction of IFN-β [19]. To confirm the expression of VF1 in MNV-3-infected cells, RAW264.7 cells were either mock-infected or infected with MNV-1 or MNV-3 at a high m.o.i., and harvested at 12 h post-infection. As expected, western blot analysis showed that VF1 expression was readily detected in both MNV-1 and MNV-3 infected cells (Fig. 1b).

The mitochondrial localization of MNV-1 VF1 was previously determined through biochemical fractionation studies, and confocal imaging of GFP-tagged VF1 in transfected cells [19]. Given the high sequence similarity between VF1 proteins from different MNV strains (Fig. 1c), with 89% similarity between VF1 encoded by MNV-3 and MNV-1, we hypothesized that VF1 expressed in MNV-3-infected cells will also localize to mitochondria. To test this, N-terminal or C-terminal MNV-3 VF1 GFP-fusions were constructed and transfected into MEFs. As shown in Fig. 1d, both the N- and the C-terminal MNV-3 VF1 GFP-fusions demonstrated a mitochondrial localization pattern, akin to that observed for MNV-1 [19]. To examine the localization of MNV-3 VF1 expressed in the context of viral infection and to rule out any artefacts due to fusion to GFP, we carried out biochemical fractionation on infected murine macrophage cultured cells. VF1 was detected in the mitochondrial fractions but not in the cytoplasmic fractions in both MNV-1 and MNV-3 infected cells (Fig. 1e), indicating a mitochondrial localization. The host proteins GAPDH and ATP56 were used to confirm enrichment of cytoplasmic and mitochondrial fractions respectively. Taken together, these data indicate that like MNV-1 VF1, MNV-3 VF1 localizes to the mitochondria in transfected and infected cells and suggest that its mitochondrial localization does not require any other viral component.

**RESULTS**

**VF1 is expressed during replication of MNV-3 in tissue culture**

The MNV-1.CW1 acute and attenuated strain (MNV-1) originated from the brain of immunocompromised mice and was subsequently plaque purified three times, whereas the persistent MNV-3 strain was isolated from a persistently infected murine research colony [5, 8]. We have previously demonstrated that VF1, encoded by the MNV-1 ORF4 (Fig. 1a), is expressed by 9 h post-infection of murine macrophage cultured cells, that it localizes to the mitochondria, and that it antagonizes the induction of IFN-β [19]. To confirm the expression of VF1 in MNV-3-infected cells, RAW264.7 cells were either mock-infected or infected with MNV-1 or MNV-3 at a high m.o.i., and harvested at 12 h post-infection. As expected, western blot analysis showed that VF1 expression was readily detected in both MNV-1 and MNV-3 infected cells (Fig. 1b).

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**VF1 is not required for MNV-3 replication in cell culture**

In order to assess a potential role for the VF1 protein in MNV-3 replication, an MNV-3 infectious clone was mutated to introduce one or three stop codons into the VF1 coding frame to produce the mutants M1 or M3 respectively (Fig. 2a, b). MNV-3 M1 has a single nucleotide substitution (T5118A) that truncates the VF1 protein after 17 aa, whereas MNV-3 M3 has three nucleotide substitutions at T5118A, C5198A and T5207A. VF1 protein is encoded in an alternative reading frame to ORF2, which encodes the major capsid protein VP1, so all mutations were designed to be non-synonymous for VF1 without affecting the overlapping capsid coding sequence (Fig. 2b). The yields of infectious MNV-3 M1 and M3 mutants, determined using the previously described DNA-based reverse genetics system [10], were comparable to the WT MNV-3 titres (Fig. 2c). To investigate if VF1 was required for MNV-3 replication in vitro, replication of MNV-3 M1 and M3 in BV-2 cells was compared to that of WT MNV-3 (Fig. 2d). The lack of VF1 had no impact on the ability of MNV-3 to replicate in the immortalized microglial BV-2 cell line, as we have previously seen with MNV-1 [19].

**The MNV-3 VF1 protein inhibits virus-induced apoptosis and IFN induction**

We have previously shown that in the absence of VF1, MNV-1-induced apoptosis is increased with enhanced caspase 3/7 cleavage observed in infected cells, indicating that VF1 expression delays apoptosis [19]. However, the induction of apoptosis occurs late in the MNV life cycle, and therefore the impact of the lack of VF1 on virus-induced apoptosis is seen only in the latter stages of the infection [19]. To assess a potential role for VF1 in apoptosis induction during MNV-3 infection, we compared the effect of the lack of VF1 expression on caspase 3/7 activity in MNV-1 and MNV-3 infected cells. We found that the absence of VF1 expression in MNV-3 M1 had only a marginal, but reproducible, impact on the levels of active caspase 3/7 when compared to WT MNV-3 (Fig. 3a, top panel). This was in contrast to the more substantial difference observed when comparing WT MNV-1 and MNV-1 M1-infected cells. This impact was not due to any inherent differences in the overall replication rate of MNV-1 and MNV-3 as NS7 expression was comparable (Fig. 3a, bottom panel).

While VF1 from MNV-1 has been shown to inhibit IFN-β induction during infection [19, 29], whether this function is present in MNV-3 is unclear, as reports using VF1 from a variant of MNV-3 that does not persist in immunocompetent mice failed to inhibit RIG-I-dependent induction of a luciferase reporter under the control of the IFN-β promoter [29]. Preliminary data from our group, however, suggest that VF1 can, under certain conditions, inhibit the expression of co-transfected transgenes (data not shown), thus complicating the use of reporter-based assays to investigate VF1 function. We therefore explored the role of VF1 in antagonism of IFN-β production during infection using cytokine
ELISA. Infection with MNV1 M1 mutant led to significantly increased secretion of IFN-β compared to WT MNV1, as expected (Fig. 3b). A similar observation was made with MNV-3, confirming that the VF1 protein from MNV-3 also inhibits the innate response. We also noted that the levels of IFN-β produced from WT MNV-3-infected cells were considerably higher than those infected with WT MNV-1 (Fig. 3b).

**Loss of VF1 incurs a fitness cost in MNV-3 replication in vivo**

Although the MNV-3 VF1 protein was not essential for viral replication in tissue culture, its importance may be more apparent during infection of its natural host. Indeed, some viral accessory proteins that modulate the host response are dispensable for replication in cell culture, but play a role in pathogenicity in vivo [30]. To explore this, we orally...
inoculated immunocompetent mice with sequence-verified WT MNV-3 and MNV-3 M1. As expected based on previous studies [11], no significant effect of WT MNV-3 or MNV-3 M1 infection on weight was observed throughout the duration of the 28-day-long experiment (Fig. 4a).

Quantification of viral genome copies from stool revealed that MNV-3 M1 was secreted at significantly lower levels throughout the first 3 days of infection compared to WT MNV-3 (Fig. 4b). The greatest difference was observed at day 3 post-infection where wildtype titres peaked at $10^6$, but MNV-3 M1 titres were two orders of magnitude lower at $10^4$. These differences began to subside from day 5 onwards, when shedding for MNV-3 decreased and MNV-3 M1 increased. Sequence analysis of the secreted virus at day 5 revealed introduction of a tryptophan at position 17 in VF1, in place of the introduced stop codon. This position is a leucine in WT MNV3, so this mutation represents a phenotypic, non-genotypic reversion of MNV-3 VF1, which occurred in 75% of mice by day 5 (Fig. 4b, c). In these mice, the reverted virus was shed at comparable levels to WT MNV-3. Conversely, the mice shedding the lowest levels of MNV-3 M1 at day 5 did not have the reverted virus, indicating that absence of MNV-3 VF1 compromises viral fitness and replication in mice, resulting in selective pressure to restore its expression. However, coinciding with the phenotypic reversion that occurs from day 5 that restores VF1 expression, comparable levels of MNV-3 M1 and WT MNV-3 were detected in all tissues with the exception of MLN, where MNV-3 M1 kinetics were still somewhat delayed even after day 5 post-infection (Fig. 5a–f).

MNV-3 VF1 is not required for viral persistence

Advances in the last decade have demonstrated a role for IFN responses in MNV persistence [16]. Treatment with type III IFNs was sufficient for clearance of a persistent strain of MNV [31, 32], and an acute strain of the virus appeared to persist in IFN-λ receptor-deficient [33] and conditional type I IFN receptor-deficient mice [34]. Since VF1 is able to antagonize IFN responses, we predicted that it would play a role in viral persistence. The reversion of MNV-3 M1, which had a single stop codon in ORF4, indicates that VF1 probably plays a role in viral fitness, as its absence compromised viral replication in the infected host. However, it also presented a limitation in examining the potential role of VF1 on the establishment of persistence. To prevent the restoration of VF1 expression, we used the triple stop codon mutant MNV-3 M3 characterized

To determine the replication kinetics of MNV-3 M1 in tissues, we harvested tissue samples from the MLN, spleen, duodenum, ileum, caecum and colon of mice infected with WT MNV-3 or MNV-3 M1 for viral genome quantification. As shown in Fig. 5a–f, the highest viral copies for both WT MNV-3 and MNV-3 M1 were observed in the MLN at day 7, and in the ileum, caecum and colon at day 5, while the spleen and duodenum were significantly lower. Similar to the pattern observed for shed virus, MNV-3 M1 was found at significantly lower levels than WT MNV-3 at days 1–3 in MLN, ileum, caecum and colon (Fig. 5a, d, e and f). However, coinciding with the phenotypic reversion that occurs from day 5 that restores VF1 expression, comparable levels of MNV-3 M1 and WT MNV-3 were detected in all tissues with the exception of MLN, where MNV-3 M1 kinetics were still somewhat delayed even after day 5 post-infection (Fig. 5a–f).
in Fig. 2a–c. Immunocompetent mice were orally inoculated with sequence-verified WT MNV-3 and MNV-3 M3, and stool samples were collected over a 3-month period to quantify viral genome copies with qRT-PCR (Fig. 6a). As before, peak shedding was observed at 3 days post-infection with WT MNV-3, while MNV-3 M3 was shed at 10-fold lower levels. Shedding of MNV-3 WT and MNV-3 M3 continually decreased over the duration of the study, with MNV-3 M3 consistently shed lower, close to the detection limit. Similar patterns were observed with infectious viral titres from the stool samples, confirming that there was shedding of infectious virus throughout the duration of the study (Fig. 6b). Sequence analysis verified the triple stop codons were intact throughout the course of infection, suggesting that VF1 was not necessary for viral persistence but instead contributed to viral fitness during a persistent infection of an immunocompetent host.

DISCUSSION

The balance between control of infection by host immunity and various mechanisms of subversion by viruses can determine how rapid viral clearance occurs (reviewed in [35, 36]). Since the initial discovery of the acute MNV-1 virus strain in 2003, a number of other MNV strains have been identified that have been shown to persist, including MNV-3 [5, 8, 37]. While MNV-1 is normally cleared in a matter of days, MNV-3 has been shown to persist for up to 9 months [15]. In the context of acute MNV-1 infections, VF1 has been shown to be an innate immune modulator and is important for virulence. In this study we utilized in vitro culture systems, reverse genetics and in vivo challenge models available for MNV to compare characteristics and functions of VF1 originating from acute or persistent MNV strains. We showed that the MNV-3 VF1 protein displays the same functions in vitro as MNV-1 VF1 and revealed that it is important for viral fitness during persistent infections.

The persistent strain of MNV used in this study, MNV-3, encodes VF1 that is 89% identical to that of the MNV-1 strain at the amino acid level. Sequence variation from the two strains did not affect the mitochondrial localization of VF1, suggesting that its mitochondrial targeting sequence is conserved, although this has yet to be defined. However, we found differences in the ability of VF1 to delay apoptosis and antagonize innate signalling in vitro, with MNV-3 VF1 less potent in both of these activities. Expression levels of VF1 remained similar in assays to compare these functions, so these disparities are likely to be functional. Although the mechanism of VF1 immune antagonism has yet to be elucidated, the decreased potency of VF1 from MNV-3 may be due to weak binding interactions with an as yet unidentified interacting partner, or inherently decreased functional activity. A comparison of VF1 from multiple MNV strains is therefore warranted and may be informative in identifying key residues important for VF1 function.

In this current study, we have observed a surprisingly higher induction of IFN in cells infected with the persistent MNV-3, compared to those infected with the acute MNV-1. Whether this is due to a difference in growth kinetics is unclear. Indeed, we have previously shown that MNV-1 grows at slightly higher titres in RAW264.7 cells compared to MNV-3 [38], and this may in turn potentially engender a greater abundance of VF1 and other viral inhibitors of IFN induction. On the other hand, previous studies comparing acute and persistent strains of MNV show strain-specific differences independent of any difference in growth. For instance, cells infected with the persistent S99 strain induce a significantly attenuated response compared to the acute
CW3 strain in the absence of any differences in growth [39], whereas a higher induction of IFN was observed in cells infected with a variant of MNV-3 that does not persist but features a slightly faster growth kinetics compared to those infected with CW3 [29]. Our data thus add to a growing body of work that portrays interesting strain-specific differences in the IFN response against MNV that require further work to disentangle.

Alternatively, the lower titres previously observed in MNV-3 compared to MNV-1 in RAW267.4 cells [38] could be as a result of the difference in IFN induction, although there could also be a strain-specific evasion of IFN signalling that precludes this effect. For instance, the persistent CR6 MNV strain has been shown to induce a robust type I IFN response, but little or no type III IFN response both in vivo and in vitro [31, 40]. At the same time, CR6 favours infection of a

**Fig. 5.** Replication kinetics in tissues. C57BL/6 mice were orally gavaged with 1000 TCID\(_{50}\) of WT MNV-3 (3-WT) or MNV-3 M1 (3-M1). Mock inoculated mice were treated with DMEM. vRNA was extracted from tissue samples, including (a) MLN, (b) spleen, (c) duodenum, (d) ileum, (e) caecum and (f) colon, and quantified using qPCR. Statistical analyses were carried out using an unpaired t-test. For WT MNV-3, \(n=20\); for MNV-3 M1, \(n=20\); and for mock, \(n=5\). Shaded area shows the time points from which reversion was seen. *\(P \leq 0.05\); **\(P \leq 0.01\). LOD, limit of detection.
subtype of intestinal epithelial cells (Tuft cells) \textit{in vivo} [41] that marginally responds to type I IFNs [42], and encodes the secreted version of NS1 that inhibits signalling downstream of type III IFN receptors [43]. It is possible that MNV-3 also has a mechanism of evading downstream IFN signalling, although there is currently no direct experimental evidence showing this.

Phenotypic differences attributed to VF1 expression were seen in the \textit{in vivo} model of MNV-3 infection that were not observed \textit{in vitro}. Whilst viral replication remains unaffected in tissue culture following deletion of VF1, \textit{in vivo} viral replication is severely compromised during the acute phase of infection, indicating the presence of greater selective pressures imposed on MNV-3 M1 in infected mice. Our data indicate that at 5 days post-infection, selection of a revertant virus occurred, probably due to selective pressures by the host immune system. The same revertant mutation arose in all the mice between 5 and 14 days post-infection, reverting from a stop codon to tryptophan, which probably restored VF1 expression. While this mutation was not a genotypic reversion, the revertant virus replicated to comparable levels as the WT virus. We believe this is indicative of a role for VF1 in \textit{in vivo} viral fitness.

To examine a potential role for VF1 in the establishment of persistence, we reduced the likelihood of VF1 restoration by introducing three stop codons in ORF4 to generate the MNV-3 M3 mutant, which replicated at the same levels as WT MNV-3 in tissue culture. Consistent with our findings with the single stop codon mutant MNV-3 M1, there was a significant reduction in replication of MNV-3 M3 in infected mice, compared to the WT virus. VF1 expression remained suppressed throughout the 3 month study as restorative mutations did not occur. At the same time, viral fitness of MNV-3 M3 remained consistently low, as replication levels were maintained comparatively lower than WT MNV-3. Although replicating close to the detection limit, MNV-3 M3 was never cleared from all the mice examined. These findings provide evidence that despite VF1 contributing to viral fitness during persistent infections, it is not required for viral persistence per se.

Considerable progress has been made in elucidating both host and viral determinants of persistence in MNV infection. First, comparative studies between the persistent CR6 strain and the acute CW3 strain revealed a genetic determinant of persistence due to a single amino acid change within the 5' domain of the non-structural protein NS1/2 [44]. Specifically, a change from aspartic acid to glutamic acid at position 94 allows the acute CW3 strain to establish a persistent infection, with the reverse being true for CR6. Second, there is evidence of a critical role played by IFN responses in MNV persistence [16]. Treatment with type III IFNs was sufficient for clearance of a persistent strain of MNV for instance [31, 32], and the acute CW3 strain of the virus persists in IFN-λ receptor (IFNLR1)-deficient mice [33], as well as in CD11c-Ifnar1-/- mice that have conditional type I IFN receptor-deficient dendritic cells [34]. Lastly, the presence of commensal bacteria in the gut also promotes MNV persistence, as treatment of mice with broad-spectrum antibiotics to clear the microbiome also prevents establishment of intestinal persistence of CR6 in a manner dependent on IFNLR1 [45, 46]. There is therefore a complex interplay of viral factors, innate immune defences (especially type III IFNs), and commensal bacteria that determine persistence, although the exact mechanistic details are still not completely understood. Our data show that inhibition of IFN responses by VF1 does not translate into impairment of persistence but affects viral fitness. The effect of VF1 on type III IFN induction has yet to be determined, in contrast to NS1/2 for which there are preliminary data that suggest it mediates evasion of type III IFN responses in a strain-specific manner [33]. Nevertheless, the findings described in this paper may open new avenues in investigating norovirus immunopathology through persistence in mice.

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C.B., A.S.J., L.T., F.S., D.B. and I.G., were all involved in the conceptualization of the study, the interpretation of the results and writing of the manuscript. C.B., A.S.J., F.S. and D.B., were directly involved in the experimental work. I.G., was responsible for obtaining funding to support the work.

Conflicts of interest
The authors declare that there are no conflicts of interest.

References


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