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HAL Id: hal-02627950
https://hal.inrae.fr/hal-02627950
Submitted on 26 May 2020

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Model of persistent foot-and-mouth disease virus infection in multilayered cells derived from bovine dorsal soft palate

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Abstract
Foot-and-mouth disease virus (FMDV) causes a highly contagious vesicular disease in livestock, with serious consequences for international trade. The virus persists in the nasopharynx of cattle and this slows down the process to obtain an FMDV-free status after an outbreak. To study biological mechanisms, or to identify molecules that can be targeted to diagnose or interfere with persistence, we developed a model of persistent FMDV infection in bovine dorsal soft palate (DSP). Primary DSP cells were isolated after commercial slaughter and were cultured in multilayers at the air-liquid interface. After 5 weeks of culture without further passage, the cells were infected with FMDV strain O/FRA/1/2001. Approximately, 20% of cells still had a polygonal morphology and displayed tight junctions as in stratified squamous epithelia. Subsets of cells expressed cytokeratin and most or all cells expressed vimentin. In contrast to monolayers in medium, multilayers in air demonstrated only a limited cytopathic effect. Integrin αvβ6 expression was observed in mono- but not in multilayers. FMDV antigen, FMDV RNA and live virus were detected from day 1 to 28, with peaks at day 1 and 2. The proportion of infected cells was highest at 24 hr (3% and 36% of cells at an MOI of 0.01 and 1, respectively). At day 28 after infection, at a time when animals that still harbour FMDV are considered carriers, FMDV antigen was detected in 0.2%–2.1% of cells, in all layers, and live virus was isolated from supernatants of 6/8 cultures. On the consensus level, the viral genome did not change within the first 24 hr after infection. Only a few minor single nucleotide variants were detected, giving no indication of the presence of a viral quasispecies. The air-liquid interface model of DSP brings new possibilities to investigate FMDV persistence in a controlled manner.
1 | INTRODUCTION

Foot-and-mouth disease (FMD), a highly contagious vesicular disease of cloven-hoofed animals, has important consequences for livestock farming. FMD influences the agricultural economy on a global scale, partly by affecting production (Rweyemamu et al., 2008), but also because a country's official FMD status determines its access to international markets (OIE, 2018). A confirmed FMD diagnosis can result in exclusion from the most profitable trade (Knight-Jones & Rushton, 2013). The aetiologic agent of FMD is a 30 nm sized, positive-sensed single-stranded RNA virus, the FMD virus (FMDV), which belongs to the genus Aphthovirus within the family Picornaviridae. The FMD virions consist of naked capsids with high antigenic variation, resulting in the existence of seven serotypes (O, A, C, SAT1, SAT2, SAT3 and Asia 1) and multiple subtypes. Both domesticated and wild even-toed ungulates can become infected. In cattle, the clinical signs are characterized by vesicular lesions in and around the mouth, on the muzzle, feet and udder, and by sudden deaths in young individuals, as well as by secondary affections such as mastitis and reproductive disturbances (Kitching, 2002). Virus shedding occurs already before clinical signs, which, in combination with high viral loads and high level of tenacity in the environment, leads to a very high transmissibility (Alexandersen, Quan, Murphy, Knight, & Zhang, 2003; Bravo de Rueda, de Jong, Eble, & Dekker, 2015). Moreover, infected cattle can carry virus in the pharyngeal area for a prolonged period, even despite preceding vaccination (Eschbaumer et al., 2016; Stenfeldt et al., 2016). Animals that harbour FMDV longer than 28 days post-infection (dpi) are defined as carriers (OIE, 2018) and, although convincing scientific evidence for spontaneous virus transmission to naïve animals is missing (Bertram et al., 2018; Bronsvoort et al., 2016; Maree et al., 2016; Parthiban, Mahapatra, Gubbins, & Parida, 2015), carriers are considered a potential virus source, either in vivo or post-mortem (Arzt, Belsham, Lohse, Botner, & Stenfeldt, 2018). This slows down the process to return to the status 'FMD free country or zone' and favours stamping-out strategies for disease control following an incursion of the virus. Persistent virus replicates in follicle-associated epithelium in the nasopharynx and soft palate (Alexandersen, Zhang, & Donaldson, 2002; Arzt, Juleff, Zhang, & Rodriguez, 2011; Stenfeldt et al., 2016) and is stored in an inactive form in the germinal centres of nasopharyngeal lymph nodes (Juleff et al., 2008; Maree et al., 2016). Reliable diagnostic methods to identify the persistently infected animals are lacking. Repeated oropharyngeal probang sampling or pharyngeal tonsil swabs are needed (Maree et al., 2016) and the probability to detect a carrier cattle in a herd rapidly decreases, both with time after an outbreak and with the age of the sampled animal (Bronsvoort et al., 2016). More robust ways to diagnose persistence, or even to treat or prevent this condition, could potentially help in the control of FMDV. However, to reach these goals, the mechanisms involved need to be elucidated.

Suggestions have already been put forward on how persistence is induced, such as through viral suppression of host immune responses (Pacheco et al., 2015; Stenfeldt et al., 2016; Stenfeldt, Eschbaumer, et al., 2017), adaptation of host cells to the virus (Martin Hernandez, Carrillo, Sevilla, & Domingo, 1994; de la Torre et al., 1988) or vice versa, through mutations in the viral genome leading to functional changes, such as immune escape (Gebauer et al., 1988), or a change in the use of receptors (O’Donnell et al., 2014). Studies have been performed in animals as well as in monolayers of pharyngeal cells or other epithelial cell lines, namely BHK-1, IBRS-2 and MDBK cells (Kopliku et al., 2015; O’Donnell et al., 2014; de la Torre, Davila, Sobrino, Ortin, & Domingo, 1985). The drawback of cells in culture is that they need to be regularly disrupted, that is passed, in order to survive. To avoid such passages, which do not occur in vivo, we developed a model of multilayer cells from the dorsal soft palate (DSP) that can be left intact for several months. The aim was to establish a model of persistent infection, to enable future studies of virus-host interactions, or the screening of molecules to detect or interfere in persistence. Hypotheses raised from results obtained in this model could be verified in live animals and, hopefully in routine, in vitro studies in the model could precede interventional in vivo studies to spare experimental animals.

2 | MATERIALS AND METHODS

2.1 | Virus

The FMDV O Clone 2.2 (Cl 2.2) used in this study is a twice-plaque-purified viral clone derived from the O/FRA/1/2001 strain (Kopliku et al., 2015) that was further propagated on BHK-21 cells (four passages). Viral batches were titrated on BHK-21 monolayers by plaque assay as described in Kopliku et al., (2015).

2.2 | Isolation of cells from the bovine dorsal soft palate

Epithelial tissue from the DSP was collected immediately after commercial slaughter of clinically healthy, conventionally reared adult male and female cattle of the Swedish red and white breed. During transport to the laboratory, the tissue was stored in incubation medium consisting of Dulbecco’s Modified Eagle’s Medium (DMEM, Lonza, Belgium), supplemented per litre with 2.5 mg amphotericin B (Sigma-Aldrich, A-9528), 1 mg deoxyribonuclease (Sigma-Aldrich, DN-25), 1 g dithiothreitol (Sigma-Aldrich, D-0632), 20 mg gentamicin...
(Sigma-Aldrich, G-1397), 20 ml 1 M HEPES (VWR, BioWhittaker BE17-737F), 10 ml 200 mM L-glutamine (VWR, BioWhittaker BE17-605F), 60 mg penicillin G sodium salt (Sigma-Aldrich, P3032) and 100 mg streptomycin sulphate salt (Sigma-Aldrich, S6501). In the laboratory, epithelial tissue was dissected and digested at 4°C overnight in incubation medium additionally supplemented per litre with 1 g protease XIV (Sigma-Aldrich, P5147).

Epithelial cells were thereafter manually scraped off the underlying tissue, filtered through a 40 μm cell strainer and incubated in cell culture flasks for 4 hr at 37°C and 5% CO₂. At this stage, the cell culture medium consisted of DMEM containing 10% gamma-irradiated, heat inactivated, foetal calf sera (FCS, HyClone™, GE Healthcare) and supplemented per litre with 31.25 KU Nystatin (Sigma-Aldrich, N6261), in addition to HEPES, L-glutamin, penicillin G and streptomycin as above.

Cells that did not adhere to the plastic were centrifuged at 200×g for 10 min at room temperature. They were thereafter frozen, thawed and propagated for three to five passages in cell culture flasks before being seeded in 12 mm diameter Corning® Transwell®-COL collagen-coated PTFE membrane inserts with 3.0 μm pores (Sigma-Aldrich, CLS3494, Figure 1) at a density of 7.5 × 10⁵ cells per insert. The cell culture medium was DMEM/Nutrient Mixture F-12 Ham (Sigma-Aldrich, D8437), containing 10% FCS and supplemented per litre with 20 μg recombinant human hepatocyte growth factor (Sigma-Aldrich, H9661), in addition to L-glutamin, penicillin G and streptomycin as above. This medium was removed from the upper compartment after five days of culture and changed in the lower compartment every two or three days (Figure 1).

2.3 | Cell characterization

The cellular expression of cytokeratin, integrin α₃β₆ and vimentin was analysed after freezing and thawing of the cells, and after three to five passages in flasks and culture in a Nunc® Lab-Tek® permcano Chamber Slide™ system (Sigma-Aldrich, C7182), as well as in cells cultured in multilayers on inserts at the air-liquid interphase for five weeks without passage. Cells were fixed in −20°C methanol for 5 min at room temperature prior to staining. The target molecules were detected by immunofluorescence microscopy (in a Nikon Eclipse Ts2R microscope) or confocal laser scan microscopy (in a ZEISS LSM700 microscope) by using mouse monoclonal antibodies against human cytokeratin (type 4, 5, 6, 8, 10, 13 and 18, clone C-11, Sigma-Aldrich, C2931, with interspecies cross-reactivity) and bovine vimentin (clone RV202, Santa Cruz Biotech, sc-32322) and mouse integrin α₃β₆ (clone 10D5, Abcam, ab77906 (Burman et al., 2006), together with rat monoclonal antibodies against mouse IgG1 or IgG2b, heavy chain, conjugated with FITC or Alexa 647, respectively (clone M1-14D12, eBioscience, 11-4015, or clone SB84a, Abcam, ab172325, respectively). The cells were mounted with ProLong® Diamond Antifade Mountant with DAPI (Life technologies corporation), according to the manufacturer’s instructions. The cytokeratin expression was further assessed by immunohistochemistry (IHC) on paraffin-embedded, FMDV-infected inserts with a pan-cytokeratin cocktail that consisted of two monoclonal mouse antibodies (clones A1/A3, DAKO, M3515, which recognize cytokeratin 1, 2, 3, 4, 5, 6, 7, 8, 10, 13, 14, 15, 16 and 19 and which recognized bovine epithelial cells). Immunohistochemistry (IHC) was performed using an automated Discovery XT (Ventana Medical Systems, Roche Diagnostics), with streptavidin-biotin-alkaline phosphatase, 5-bromo-4-chloro-3-indolyl phosphate as a substrate and nuclear fast red counterstaining.

The cell morphology was studied by light and electron microscopy. For light microscopy, after fixation in 10% buffered formalin, selected multilayers and the underlying PTFE membranes were embedded in 1.3% agarose then left in 70% ethanol overnight. They were then embedded in paraffin, routinely processed, sliced at 4 μm, stained with haematoxylin-eosin-saffron (HES) and examined by light microscopy. For electron microscopy, two control multilayers and the underlying PTFE membrane were fixed in Sorensen phosphate buffer containing 2.5% glutaraldehyde, 0.1% picric acid, 2% paraformaldehyde and 0.18 mol/L sucrose. The samples were post-fixed in 1% osmium tetroxide then washed in Sorensen buffer. They were then dehydrated in ethanol and embedded in Spürr’s low viscosity epoxy resin. Semi-thin sections were stained with Toluidine blue for light microscopy. Ultra-thin sections (60 nm) were stained with uranyl acetate and examined under a Hitachi-7100 transmission electron microscope equipped with a digital camera.

2.4 | Experimental design and FMDV infection

Dorsal SP cells were cultured on inserts for five weeks and were thereafter infected with FMDV O Cl 2.2, or a placebo. In the first

**FIGURE 1** Schematic draw of a permeable insert used to propagate multilayers of bovine dorsal soft palate cells. Insert (a); upper compartment (b); multilayer of bovine dorsal soft palate cells (c); cell culture medium (d); porous membrane (e); lower compartment (f) and well (g) of a 12-well plate.
and second experiment (Exp 1 and 2), cells originated from a male and a female, respectively, and were infected at a multiplicity of infection (MOI) of 0.01. In a third experiment (Exp 3), the cells originated from the same female individual as in Exp 2, but were infected at an MOI of 1. A different batch of virus (albeit from the same passage) was used in Exp 1, compared to Exp 2 and 3, which were performed with the same batch. The inserts were incubated for 1 hr with 500 μl of cell lysate from FMDV-infected or uninfected cell cultures that had been clarified by centrifugation. The inoculum was then left on the inserts (Exp 1 and 2) or removed (Exp 3). Immediately following infection and thereafter at a maximum interval of three days, the upper compartments were washed with 500 μl cell culture medium containing 10% FCS and, similarly, but only from 2 dpi, the medium in the lower compartments was changed (for the compartment layout and experimental design, please see Figures 1 and 2, respectively). The wash medium was harvested and frozen at −80°C, pure and diluted 1:4 in TRIzol LS (Invitrogen), for assays of infectious FMDV and FMDV RNA, respectively, as indicated in Figure 2. Moreover, inserts were regularly harvested and fixed, embedded and then cross-sectioned for the detection of FMDV antigens, until day 28 (Figure 2). Cells from 0, 1 and 28 dpi were lysed in TRIzol and stored at −80°C for viral RNA sequencing and for transcriptome analyses. In total, 90 inserts were used in the study.

2.5 | FMDV isolation

2.5.1 | Cell line

IBRS-2 swine kidney epithelial cells (CCLV-RIE 103, FLI) were grown in Earle’s minimum essential medium (MEM) with L-glutamine (Invitrogen), supplemented with 7% FCS (Eurobio, Courtaboeuf, France), 1.5% lactalbumin hydrolysate (Sigma-Aldrich), 1% penicillin-streptomycin (PS) (Invitrogen) and 25 mM HEPES (Invitrogen).

2.5.2 | Assessment of FMDV-infectivity

Samples of undiluted cell culture medium used to wash the upper compartments of the inserts were analysed for infectious virus. Ninety-six-well plates were seeded with 5 × 10^4 IBRS-2 cells per well and incubated at 37°C under 5% CO₂ for 24 hr. The monolayers were thereafter washed twice with serum-free culture medium and each sample from the DSP cultures was inoculated in a volume of 50 μl per well. After 1 hr of adsorption at 37°C under 5% CO₂, 250 μl of growth medium was added. Cells were then incubated at 37°C under 5% CO₂ and were monitored for cytopathic effect (CPE) for 48 hr. Immunofluorescent detection was performed using monoclonal antibodies raised against the capsid protein VP2 of FMDV type O (mAb 13G11), as described in Kopliku et al., 2015.

Virus titrations were performed on diluted wash-samples (from 1:10). Fifty μl of cell culture medium without serum was added in each well of 96-well plates. Ten-fold serial dilutions of virus (50 μl per well, eight replicates per dilution) were distributed, and the same volume of growth medium was added to the mock-infected control wells. About 50 μl of IBRS-2 cell suspension at 3 × 10^5 cells/ml in medium was added in each well. Plates were incubated at 37°C under 5% CO₂. Three dpi, plates were fixed with 70% ethanol for 30 min at room temperature (RT) and then stained with methylene blue (1% w/v in water). Values for 50% tissue culture infective dose (TCID₅₀) per 50 μl were calculated using the Stellmann and Bornarel table (Stellmann & Bornarel, 1971) and were translated into values per ml.

2.6 | Detection of FMDV RNA

2.6.1 | RNA extraction for FMDV RT-qPCR

Viral RNA was extracted from 50 μl of cell culture washes from the upper compartment, using the LSI MagVet Universal Isolation kit (Life Technologies SAS) and a KingFisher robotic magnetic particle processor (Thermo Scientific) according to the manufacturers’ instructions. In total, 2–6 samples were analysed per time point.

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**FIGURE 2** Experiment timeline and FMDV detection. Bovine dorsal soft palate cells were propagated in multilayers on porous membrane inserts in the air-liquid interphase and were infected with FMDV O CI2.2, or placebo. In experiment 1 (Exp 1, empty legends) and 2 (Exp 2, light grey legends), cells derived from male and female cattle, respectively, and were infected at an MOI of 0.01. In experiment 3 (Exp 3, dark grey legends) cells derived from the same female cattle and were infected at an MOI of 1. Cell culture medium and cells were analysed for the presence of FMDV, by virus isolation (triangles), RT-qPCR (circles) or immunohistochemistry (squares), at indicated days post-infection.
except for in experiment 3, when single samples were analysed at 2 and 4 hr post-infection (hpi), as well as at 5, 7, 12, 16 and 23 dpi. The RNAs were eluted with 80 μl of elution buffer provided with the kit.

### 2.6.2 | One-step duplex real-time FMDV RT-qPCR

A one-step duplex pan-FMDV real-time RT-PCR (rtRT-PCR) assay was used as previously described (Gorna et al., 2016). The assay targeted the FMDV 3D coding region and the cellular β-actin gene as an internal control. A weakly positive FMDV RNA control was included in each run as well as a negative control containing water only.

### 2.7 | Detection of FMDV antigen by immunohistochemistry

Immunohistochemistry was performed using an automated Discovery XT (Ventana Medical Systems, Roche Diagnostics), with streptavidin-biotin-peroxidase, 3′-diaminobenzidine as a substrate and haematoxylin counterstaining. The primary antibodies were rabbit polyclonal against FMDV (FMDV O1 Manisa R3 262/97, The Pirbright Institute, United Kingdom) used at 1/1000; mouse monoclonal against FMDV non-structural protein 3D polymerase (mAb 3F12, kindly provided by Dr E. Brocchi from IZSLER, Brescia, Italy) and mouse monoclonal against cytokeratin AE1-AE3 (M3515, DAKO), used at 1/500.

Positivity for FMDV was evaluated by image analysis of sections stained with polyclonal antibodies. Scanned images of two sections per insert were taken by an ImagerZ1 Zeiss microscope and an AxioCam HRc Zeiss camera. FMDV-positive cells and total number of cells (mean 600, SD 306 cells per insert) were manually counted on each section with ImageJ Cell Counter (Schneider, Rasband, & Eliceiri, 2012).

### 2.8 | FMDV genome analysis, proteomics and transcriptomics

#### 2.8.1 | RNA extraction and sequencing

For further cell characterization and for sequencing of the FMDV inoculum strain, total RNA was extracted from FMDV O Cl 2.2 and libraries were prepared as described in Pfaff et al., (2018). Briefly, total RNA was extracted from cell culture supernatant using TRIzol (Life Technologies) in combination with the RNeasy Mini Kit (Qiagen, Hilden, Germany). Double-stranded cDNA was generated using the cDNA synthesis system kit (Roche) together with random hexamer primers (Roche). Subsequently, the GeneRead DNA Library L Core Kit (Qiagen) was used to prepare libraries for sequencing on the Ion S5XL system using the Ion 530 OT2 and Chip kit (Life Technologies). For sequencing of FMDV from inserts, polyadenylated RNA was used as described previously (Pfaff et al., 2019) from infected cell lysates obtained at 24 hpi and 28 dpi in Exp 1 and 2. Briefly, total RNA was extracted as described above and mRNA was enriched using the Dynabeads mRNA DIRECT Micro kit (Invitrogen). Subsequently, whole-transcriptome libraries were prepared using the Ion Total RNA-Seq Kit v2 (Life Technologies) and sequenced using a S5XL sequencing system (Life Technologies) along with the Ion 540 OT2 and Chip kit (Life Technologies). Proteins were extracted and identified by liquid chromatography and mass spectrometry as described previously (Pfaff et al., 2019). Briefly, proteins derived from the organic TRIzol phase of samples and were separated by SDS-PAGE. In-gel digestion was performed by trypsin, followed by purification of peptides and LC-MS/MS analyses by an EASY.nLC Il coupled to a LTQ Orbitrap-Velos mass spectrometer (Thermo Fisher Scientific).

#### 2.8.2 | FMDV inoculum sequence assembly and variant analysis

In order to create a full-length FMDV consensus sequence of the inoculum, raw reads were initially mapped against the FMDV reference FRA/1/2001 (AJ633821) using the 454 Sequencing Systems Software suite (version 3.0: Roche) with default options for quality trimming. All full or partially mapping reads were then selected for further de novo assembly using the same software. The resulting full-length FMDV sequence was annotated in Geneious (Geneious, version R10, 2018 August 1) (Kearse et al., 2012) according to the NCBI RefSeq of FMDV and uploaded to the European Nucleotide Archive (ENA) under the project accession PRJEB28325. In order to call low frequency nucleotide variants in the inoculum and in the treatment samples from different time points, the raw sequence reads were quality trimmed using Trimomatic (Trimmomatic, version 0.36, 2018, August 1) (Bolger, Lohse, & Usadel, 2014), with a 5-base wide sliding window that cuts average qualities below 20 (phred33) and reads shorter than 50 nucleotides. Subsequently, the trimmed reads were mapped along the full-length inoculum sequence using Bowtie (Bowtie2, version 2.3.4.1, 2018, August 1) in ‘very-fast-local’ mode. Duplicated reads were marked and removed from the resulting alignments using SAMTools (SAMtools, version 1.7, 2018, August 1) and single nucleotide variants (SNV) were then called using LoFreq (LoFreq, version 2.1.3.1, 2018, August 1) (Wilm et al., 2012).

### 3 | RESULTS

#### 3.1 | Primary DSP cells grew in multiple layers at the air-liquid interface

Primary cells originating from the bovine DSP were isolated and propagated in multilayers on porous membranes at the air-liquid interface. After protease digestion of epithelial tissue, filtration, removal of rapidly adhering cells, and freezing and thawing, the expression of vimentin intermediate filaments was less extensive than that of cytokeratin (data not shown). After five passages in flasks and before seeding in inserts, a larger number of cells expressed vimentin than cytokeratin (Figure 3a and b). The cells were mostly polygonal or round and expressed integrin αvβ6 to varying extent (Figure 3c). Secondary antibodies alone did not generate fluorescence at similar...
conditions of confocal laser microscopy scanning or fluorescent microscopy (data not shown). Monolayers of the cells were infected with FMDV at an MOI of 0.01 and an almost complete CPE was observed 24 hpi (data not shown). Five days after seeding cells on inserts, the cell culture medium was removed from the upper compartment and the cells continued to grow in air. Five weeks after seeding, at the time of FMDV infection of inserts, the cells grew in multilayers and approximately 20% had a polygonal morphology and displayed tight junctions (Figure 4a and b). A subset of cells expressed cytokeratin (Figure 4c) and mRNA transcripts related to both tight junctions and cytokeratins were detected by transcriptomics (Figure 5). Whereas integrin $\alpha_V\beta_6$ expression could no longer be demonstrated by immunostaining or proteomics (data not shown), low levels of mRNA transcripts coding for the integrin chain $\beta_6$ were still detectable in most inserts (Figure 5). On the other hand, mRNA transcripts coding for the integrin chains $\alpha_V\beta_1$ and $\beta_3$ were abundant (Figure 5) and this was confirmed by proteomics (Figure 6).

3.2 | FMDV infection of DSP multilayers resulted in limited lysis

Foot-and-mouth disease virus (FMDV) infection of DSP multilayers in inserts resulted in limited lysis of cells in the upper cell layers, from 24 hpi to 7 dpi, as observed by light microscopy (Figure 7). At 24 hpi, infection resulted in multifocal cell detachment and loss, with foci of sparing of several layers (infection at MOI 0.01) and foci of loss of the deeper layers (infection at MOI 1), in contrast to the mock-infected membranes that did not show lysis (Figure 8). A vesicle-like structure was found in one insert at 24 h after infection with FMDV at an MOI of 1: the multilayered epithelium was characterized by intercellular edema and an image consistent with suprabasilar clefting (Figure 9). The CPE was more extensive at an MOI of 1, compared to 0.01 during the first 48 h (Figure 7). Thereafter, CPE was no more observed, only cellular debris (Figure 7). The cultures seemed to have gradually recovered between 14 and 28 dpi. At 28 dpi the morphology of cells in infected and uninfected inserts did not differ (Figure 7). Sparse cytokeratin-positive cells were detected in both infected and uninfected inserts at 28 dpi; however, by using double IHC, FMDV antigens could not be detected in these particular cells (data not shown).

3.3 | FMDV RNA and infectious FMDV persisted up to 28 dpi in DSP multilayers

Throughout the course of the experiment, cell culture washes were analysed for the presence of viral RNA by real-time RT-PCR targeting the FMDV 3D protein coding region. Whereas the internal control $\beta$-actin was detected at Ct values between 26–32 (Exp 1), 23–40 (Exp 2) and 15–39 (Exp 3), FMDV RNA was not detected in cell culture washes collected before infection, but was detected shortly after infection, at 5 hpi (Exp 2, MOI 0.01, Figure 10a), or at 2 and 4 hpi (Exp 3, MOI 1, Figure 10a). The average quantity of FMDV RNA peaked at 24 hpi (Exp 1 and 2, MOI 0.01) or 48 hpi (Exp 3, MOI 1), at almost similar levels, despite the fact that the cells originated from different animals and were infected at different MOI (Figure 10a). The peaks occurred at average Ct values of 19, 16 and 15 in experiment 1, 2 and 3, respectively. After this peak, the FMDV RNA tended to continuously decrease throughout the three experiments, although in cells infected at an MOI of 1 (Exp 3), this decrease was irregular (Figure 10a). At the termination of the experiments at 28 dpi, FMDV RNA could still be detected in cell culture medium of all infected inserts analysed (Figure 10a, n = 8, Ct values ranging between 25 and 28 for Exp 1 and 2, MOI 0.01 and between 22 and 27 for Exp 3, MOI 1), but not in controls (n = 8).

In agreement with the RT-qPCR data, live FMDV was recovered from the undiluted upper compartment wash medium by isolation on susceptible IBRS-2 cultures, from 1 dpi to 28 dpi in all experiments (Table 1). The viral titres consistently peaked on 1 dpi, at higher titres in the infections performed at an MOI of 1 (average 6.7 log$_{10}$ TCID$_{50}$/ml, Exp 3) compared to those performed at an MOI of 0.01 (average 6.3 log$_{10}$ TCID$_{50}$/ml and 6.5 log$_{10}$ TCID$_{50}$/ml in Exp 1 and 2, respectively, Figure 10b). The virus could not be titrated on 28 dpi in experiment 2 and 3 (Figure 10b), since these samples were virus negative at a dilution of 1:10, which was the
starting dilution in the viral titration assay, due to the restricted volume of wash medium. FMDV could not be isolated, neither titrated, later than 2 dpi from one insert, infected at an MOI of 1, despite presence of viral RNA until 28 dpi. This particular insert had the highest quantity of FMDV RNA on 2 dpi, compared to all inserts throughout the study. The negative samples deriving from this membrane (n = 4) were excluded from the viral titration data presented in Figure 10b, to give a better view of the titres in the remaining inserts.

3.4 | FMDV antigen persisted up to 28 dpi in a low proportion of cells

Virus antigen was detected by IHC using polyclonal antibodies, from day 1 through day 28 in the three experiments, in all inserts investigated (one per experiment and time point, as indicated in Figure 2). FMDV antigen was detected in the cytoplasm of cells at the upper, middle and lower levels of the multilayers infected at an MOI of 0.01 and at an MOI of 1 (Figure 8). The proportion of FMDV-positive cells detected by IHC using polyclonal antibodies was analysed on two sections per insert. At 24 hpi, 36% of the cells were FMDV-positive after infection at an MOI of 1, while 1 to 3% of the cells were FMDV-positive at an MOI of 0.01. At 28 dpi, the proportion of infected cells was 0.15% and 0.91% at an MOI of 0.01 (Exp 1 and Exp 2, respectively) and 2.10% at an MOI of 1 (Exp 3, Figure 11). Expression of viral non-structural protein 3D was detected by IHC in the cytoplasm of cells at all the layers at 24 hpi and 28 dpi, at an MOI of 0.01 as well as at an MOI of 1 (Figure 9). This protein was additionally detected in all the cells at the edges of the vesicle-like structure observed in one insert at 24 hpi (MOI 1; Figure 9).

3.5 | FMDV genome showed no consensus-level changes at 24 hpi

Sequencing and subsequent de novo assembly of the FMDV inoculum strain resulted in two contigs spanning the complete viral genome. Both contigs extended upstream and downstream of a long polyC stretch in the 5’UTR, whose exact length was not determined but exceeded a total number of seven cytosine nucleotides. With respect to the FMDV reference FRA/1/2001 (AJ633821), eight Ns were introduced in order to mark this uncertainty. The genome was covered by 175,339 reads in total, of which 14,775 reads were non-duplicated (compare Table 2). In comparison to FRA/1/2001 (AJ633821), the inoculum strain FMDV O CI 2.2 showed seven SNPs and a single insertion. Of these, three SNPs each were located in
RNA transcripts related to tight junctions, FMDV receptor integrins and cytokeratins. Bovine dorsal soft palate cells were isolated and propagated in multilayers on porous membrane inserts in the air-liquid interface. After five weeks of culture, membranes were infected with FMDV at MOI 0.01 or MOI 1 or with uninfected cell lysate as placebo. Cells were harvested at indicated time points, before (0 hr, 0h) and post-infection (24 hr, 24 hpi, 28 days dpi) and total RNA was extracted. Messenger RNA was thereafter detected by RNA sequencing on the Ion S5XL system using the Ion 530 OT2 and Chip kit (Life Technologies). Gene transcripts associated with a) tight junctions (CLN, OCLN, TCF3, TCF7L1), b) FMDV-associated integrin chains (ITGAV, ITGB1, ITGB3, ITGB8) and c) cytokeratin (KRT) expression are expressed as variance stabilizing transformed (vs.t) read counts. Each data point represents a technical replicate that derived from duplicate analyses of two biological replicates.

Protein expression related to FMDV receptor-associated integrins. Bovine dorsal soft palate cells were isolated and propagated in multilayers on porous membrane inserts in the air-liquid interface. After five weeks of culture, membranes were infected with FMDV at MOI 0.01 or with uninfected cell lysate as placebo. Cells were harvested at indicated time points, before and after infection. Proteins were thereafter extracted, separated by SDS-PAGE and analysed by LC-MS/MS using an EASY-nLC II coupled to a LTQ Orbitrap-Velos mass spectrometer (Thermo Fisher Scientific). ITGB6 and ITGB8 were not detected.
the 5′UTR and coding region. A single SNP and an insertion were observed in the 3′UTR. SNPs in the coding region resulted in the following amino acid substitution: A560R (1C region), E922G (1D region) and F2119I (3D region).

Reads from sequencing of FMDV O Cl 2.2 and the insert samples were mapped along the de-novo assembled FMDV O Cl 2.2 consensus sequence and SNV were called. The rate of reads matching FMDV in a certain dataset varied between <0.01% and 20.03% (Table 2) with highest and lowest rates observed within samples after 24 hpi and 28 dpi, respectively. For the inserts, no nucleotide exchanges were observed at the consensus level. Due to very low genome and overall coverage, an analysis of minor variant nucleotides below consensus level was not feasible for samples from 28 dpi. Three SNVs with variant frequencies between 0.05 and 0.06 were detected for the FMDV O Cl 2.2 that all resulted in amino acid changes in the 2C region of the polyprotein (Table 3). A single SNV at position 5,994 (T → C) that resulted in an amino acid exchange (I1355T) was also observed in three samples from the inserts taken 24 hpi at frequencies of 0.08, 0.11 and 0.13.

4 | DISCUSSION

In this work, we describe an in vitro model of FMDV persistence in multilayer cultures of DSP cells, up to 28 days post-infection. The model allows to study FMDV-specific responses of target cells, viral kinetics without influence of the adaptive immunity in vivo, or cell passage in vitro, and viral changes in absence of selective pressure from the immune system (antibodies, immune cells). The model can potentially generate complementary information on pathways and origin of some of the processes that were already identified in vivo, which have likely been based on a sum of responses from a mixture of cells, including immune cells and epithelial cells that were or were not directly exposed to the virus. Moreover, to spare animals and get a better understanding of the FMDV persistence, hypotheses raised in other in vitro models, based on cell lines or monolayers of primary cells, can be verified in the multilayer model before animal experiments are performed.

We attempted to reconstitute an epithelium with cells from the stratum basale and stratum spinosum, which together constitute the approximately nine cells deep, stratified, non-cornified, squamous epithelia in the DSP of cattle (Schley, Ward, & Zhang, 2011). The model did not reach a similar depth, but some epithelial characteristics were achieved, such as impermeability to cell culture medium from 5 days of culture on inserts, and the continuous expression of tight junctions and cytokeratin in subsets of cells.

Persistent FMDV has previously been detected post-mortem in follicle-associated epithelium in nasopharynx and the DSP, and more specifically in cytokeratin-negative and weakly positive cells in the basal cell layers (Arzt et al., 2011; Pacheco et al., 2015), as well as in highly cytokeratin-positive cells in the upper layers (Stenfeldt et al., 2016). In this study, double IHC revealed coexistence of cytokeratin-positive cells and FMDV-positive cells in the inserts at 28 dpi, but did not allow the detection of co-localization of these antigens. The infected cells were consistently detected in all the layers on 28 dpi.
and cytokeratin mRNA expression persisted, albeit at lower levels compared to at the time of infection.

The majority of cells in the multilayer model expressed vimentin, but many maintained a round, flat or polygonal morphology, which suggests they were of epithelial rather than of fibroblast origin. Though vimentin was previously considered as a static fibroblast marker, both vimentin and cytokeratin can be differentially and even simultaneously expressed by the same epithelial cell (Eriksson et al., 2009; Kasper & Stosiek, 1990; Mendez, Kojima, & Goldman, 2010; Rogel et al., 2011). Vimentin expression is commonly induced when epithelial cells are cultured in vitro (Pieper et al., 1992), possibly during cell passage, because injured epithelial cells undergo epithelial to mesenchymal transition in vivo, in a process that may be reversed (Mendez et al., 2010; Rogel et al., 2011). In the present model,
fibroblast contamination was decreased or avoided by removal of rapidly adhering cells, elimination of cultures that contained elongated cells and the addition of hepatocyte growth factor that is mitogenic to keratinocytes and inhibits fibrosis (Marchand-Adam et al., 2006).

Following infection with FMDV at an MOI of 0.01 or 1, the multilayer DSP cells showed only limited CPE and rapid recovery. Only one vesicle-like structure was found in one insert at 24 hr after infection with FMDV at an MOI of 1, with an image consistent with suprabasilar clefting described in the early vesicles in bovine (Arzt, Gregg, Clavijo, & Rodriguez, 2009). This is in agreement with the observed absence of lesions and the very restricted FMDV replication that occurs in the DSP in vivo (Stenfeldt & Belsham, 2012; Zhang & Alexandersen, 2004). It has been suggested that the low
FMDV O Cl 2.2 at an MOI of 0.01 (experiment 1 and 2), or 1 (experiment 3) 0 days post-infection (dpi).

Culture cell medium from the inserts was analysed for the presence of live FMDV by isolation on IBRS-2. The IBRS-2 cells were monitored for cytopathic effect for 48 hr and immunofluorescent detection was performed using monoclonal antibodies raised against the capsid protein VP2 of FMDV type O (mAb 13G11), as described in Kopliku et al. (2015).

susceptibility to infection in this tissue, in comparison with, for example the lung, is due to low expression of both the FMDV receptor integrin αVβ6 (Giorgakoudi et al., 2015; Monaghan et al., 2005) and of TNF superfamily genes that are linked to apoptosis and necrosis (Zhu et al., 2013). Accordingly, whereas integrin αVβ6 was expressed by subsets of DSP cells after isolation, and these cells were susceptible to FMDV-mediated lysis, integrin αVβ6 expression could not be detected in the multilayers that were more resistant. Nevertheless, the multilayer cells were also infected, maybe through integrins αVβ1, αVβ3, αVβ5, αVβ8 (O’Donnell et al., 2014; Wang, Wang, Shang, Zhang, & Liu, 2015), heparane sulphate (Bai et al., 2014; Jackson et al., 1996), or other proteins that have not yet been characterized (Baranowski et al., 2000). Our model probably mirror what happens naturally but since the virus had been passed in cell culture, it cannot be excluded that the receptor use had changed compared to that of the parental strain (Baranowski et al., 1998; O’Donnell et al., 2014).

The virus replication data based on isolations and RT-qPCR on washes of the DSP cells, correlated well with the CPE. Virus replication was observed soon after infection and declined to a minimum at 28 dpi. Whereas FMDV RNA was detected in all infected inserts throughout the experiment, live virus was not detected consistently in all inserts analysed. The detection of live virus was more successful at the lower MOI (MOI 0.01) after 2 dpi, maybe because at the higher MOI (MOI 1), more cells had been lysed and free virus

### TABLE 1  Isolation of FMDV during long-term culture of bovine dorsal soft palate cells on porous membrane inserts in the air-liquid interphase

<table>
<thead>
<tr>
<th>dpi</th>
<th>Exp 1&lt;sup&gt;b&lt;/sup&gt; MOI 0.01</th>
<th>Exp 2&lt;sup&gt;b&lt;/sup&gt; MOI 0.01</th>
<th>Exp 3&lt;sup&gt;b&lt;/sup&gt; MOI 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0/1</td>
<td>0/3</td>
<td>3/3</td>
</tr>
<tr>
<td>1</td>
<td>2/2</td>
<td>4/4</td>
<td>3/3</td>
</tr>
<tr>
<td>2</td>
<td>2/2</td>
<td>4/4</td>
<td>3/3</td>
</tr>
<tr>
<td>5</td>
<td>2/2</td>
<td>2/2</td>
<td>0/1</td>
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<tr>
<td>7</td>
<td>2/2</td>
<td>4/4</td>
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<td>9</td>
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<td>12</td>
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<td>4/4</td>
<td>1/1</td>
</tr>
<tr>
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<td>4/4</td>
<td>2/2</td>
</tr>
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<td>NA</td>
<td>4/4</td>
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<td>19</td>
<td>NA</td>
<td>3/4</td>
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</tr>
<tr>
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<td>0/1</td>
</tr>
<tr>
<td>28</td>
<td>2/2</td>
<td>3/3</td>
<td>1/3</td>
</tr>
</tbody>
</table>

<sup>a</sup>Bovine soft palate cells were propagated in multilayers in vitro, on porous membrane inserts in the air-liquid interface and were infected with FMDV O Cl2.2 at an MOI of 0.01 (experiment 1 and 2), or 1 (experiment 3) 0 days post-infection (dpi).

<sup>b</sup>Culture cell medium from the inserts was analysed for the presence of live FMDV by isolation on IBRS-2. The IBRS-2 cells were monitored for cytopathic effect for 48 hr and immunofluorescent detection was performed using monoclonal antibodies raised against the capsid protein VP2 of FMDV type O (mAb 13G11), as described in Kopliku et al. (2015).

### TABLE 2  Reads matching the FMDV genome from the sequencing of the inoculum and samples from inserts 24 hpi and 28 dpi

<table>
<thead>
<tr>
<th>Sample</th>
<th>Total reads</th>
<th>Reads mapped to FMDV O Cl 2.2</th>
<th>Rate</th>
<th>Unique Reads mapped to FMDV O Cl 2.2</th>
<th>Average coverage</th>
<th>Genome coverage</th>
</tr>
</thead>
<tbody>
<tr>
<td>FMDV O Cl 2.2 (lib02533)</td>
<td>1.389.330</td>
<td>175,339</td>
<td>12.62%</td>
<td>14.775</td>
<td>405.4</td>
<td>100%</td>
</tr>
<tr>
<td>24 hpi, male, Exp 1, I (lib02361)</td>
<td>13.634.337</td>
<td>134,098</td>
<td>0.99%</td>
<td>5.976</td>
<td>82.0</td>
<td>100%</td>
</tr>
<tr>
<td>24 hpi, male, Exp 1, II (lib02345)</td>
<td>12.959.531</td>
<td>286,935</td>
<td>2.21%</td>
<td>6.735</td>
<td>93.8</td>
<td>100%</td>
</tr>
<tr>
<td>24 hpi, female, Exp 2, I (lib02451)</td>
<td>14.295.264</td>
<td>2,511,451</td>
<td>17.57%</td>
<td>8.531</td>
<td>131.3</td>
<td>100%</td>
</tr>
<tr>
<td>28 dpi, male, Exp 1, I (lib02347)</td>
<td>10.782.214</td>
<td>2,201</td>
<td>0.02%</td>
<td>776</td>
<td>8.8</td>
<td>99.2%</td>
</tr>
<tr>
<td>28 dpi, male, Exp 1, II (lib02349)</td>
<td>12.869.179</td>
<td>975</td>
<td>0.01%</td>
<td>392</td>
<td>4.2</td>
<td>90.3%</td>
</tr>
<tr>
<td>28 dpi, female, Exp 2, I (lib02455)</td>
<td>13.745.858</td>
<td>875</td>
<td>0.01%</td>
<td>241</td>
<td>3.3</td>
<td>78.2%</td>
</tr>
<tr>
<td>28 dpi, female, Exp 2, II (lib02456)</td>
<td>15.570.465</td>
<td>468</td>
<td>&lt;0.01%</td>
<td>127</td>
<td>1.6</td>
<td>50%</td>
</tr>
</tbody>
</table>
had been removed by washing. The discrepancy between virus isolation and RT-qPCR late in the infection might be explained by a production of defective virus particles, or by inhibition of live virus, by molecules such as type I interferons, defensins, cathelicidins or interfering RNA (Stenfeldt, Arzt, et al., 2017). As described by Pfaff and colleagues in a joint publication on the transcriptome of this model, such innate responses were activated (Pfaff et al., 2019), although they were probably modulated by the virus, based on findings in vivo (Stenfeldt et al., 2016; Zhu et al., 2013). The declining quantity of FMDV over time, which is also observed in vivo, suggests that the virus can infect and replicate at a low level, but not indefinitely, not even in the absence of the adaptive immune responses that are important to clear virus (Maddur et al., 2009; Stenfeldt, Eschbaumer, et al., 2017). The persistent FMDV infection thus appears as a chronic infection that is self-limiting. The multilayer model can be kept for at least three months and long-term persistence could thereby be further investigated in future studies.

Besides being associated with host factors, persistence may also rely on genetic and antigenic changes in the virus. In our model, no such changes were apparent. The consensus sequence of the virus was consistent throughout the acute and persistent phases of infection and we did not observe an accumulation of minor nucleotide variants below the consensus. The use of a viral clone might have favoured this lack of variation, although generation of mutations was previously demonstrated with plaque-purified virus (Huang, Li, Fang, & Zheng, 2011; de la Torre et al., 1988). Similarly, non-synonymous nucleotide exchanges occur within and between hosts during natural infections (Cottam, King, Wilson, Paton, & Haydon, 2009; Morelli et al., 2013) and are likely driven by selective pressure from the adaptive immune system (Gebauer et al., 1988). It has been suggested that such changes could allow viral subpopulations to escape clearance by the immune system and facilitate persistent infection (Horsington & Zhang, 2007). Previous studies have described persistence-associated amino acid substitutions in serotype O viruses (Horsington & Zhang, 2007; Pauszek et al., 2017; Ramirez-Carvajal et al., 2018), but neither these nor any other viral determinants of persistence were identified in this study.

In conclusion, this new model of bovine DSP allows long-term culture without disruption of cells and mimics the in vivo conditions, in the way that soft palate cells grow in multilayers and are in contact with air. The model will make it possible to get a better understanding of the mechanisms underlying FMDV persistence in order to develop ways to control this infection and to improve its diagnosis.

ACKNOWLEDGEMENTS

This work was supported by the Swedish Research Council (Formas, Sweden), L'Agence Nationale de la Recherche (ANR, France) and the Federal Office for Agriculture and Food (BLE, Germany), through ANHIWA ERA NET (TRANSCRIPTOVAC project, no. FP#26). The authors would like to acknowledge Lövsta Kött, Uppsala, Sweden, for providing tissue from slaughtered animals, as well as the Cells for Life Platform, partly funded by the Infrastructure Committee at SLU, Sweden, for providing facilities and equipment. The staff at the Clinical Science laboratory at SLU are acknowledged for their technical assistance and Jeremy Adler at the Scilife laboratory, Uppsala, Sweden, is thanked for his support during work with confocal microscopy. We acknowledge Patrick Zitzow of the Institute of Diagnostic Virology at the Friedrich-Loeffler-Institut for excellent technical assistance. Emiliana Brocchi of the IZSLER is acknowledged for providing monoclonal FMDV antibodies. Many thanks to Chakib Djediat of the Museum National d’Histoire Naturelle for his expertise in electron microscopy.

CONFLICT OF INTEREST

The authors declare that they have no conflict of interest.

ETHICAL STATEMENT

The authors confirm that the ethical policies of the journal, as noted on the journal’s author guidelines page, have been adhered to. No ethical approval was required as the samples were collected after commercial slaughter.

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