A Crimean-Congo hemorrhagic fever (CCHF) viral vaccine expressing nucleoprotein is immunogenic but fails to confer protection against lethal disease

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Crimean-Congo Hemorrhagic Fever (CCHF) is a severe tick-borne disease, endemic in many countries in Africa, the Middle East, Eastern Europe and Asia. Between 15–70% of reported cases are fatal with no approved vaccine available. In the present study, the attenuated poxvirus vector, Modified Vaccinia virus Ankara, was used to develop a recombinant candidate vaccine expressing the CCHF virus nucleoprotein. Cellular and humoral immunogenicity was confirmed in 2 mouse strains, including type I interferon receptor knockout mice, which are susceptible to CCHF disease. Despite the immune responses generated post-immunisation, the vaccine failed to protect animals from lethal disease in a challenge model.

Introduction

Crimean-Congo hemorrhagic fever (CCHF) virus causes a severe and frequently fatal hemorrhagic disease in people, with a mortality rate of approximately 30%.1 CCHF virus has the most extensive geographical distribution of the medically important tick-borne viral diseases.2 It is the second most widespread of the medically important viral hemorrhagic fever viruses, after dengue virus,3 and is described as an emerging virus; CCHF is distributed over much of Asia, the Middle East, Africa and expanding areas of south-eastern Europe. The continued spread of the tick vector and reservoir (Hyalomma species) through climate change and modern farming practices, has resulted in the virus becoming established in territories where it was not previously endemic; its introduction to Turkey, Greece and, more recently, Spain being testament to this.4 CCHF virus is recognized as a possible agent of bioterrorism.5 In Iraq, it was studied as a potential biological weapon,6 and the virus has also been shown to be potentially disseminated via aerosolisation.7

Recognized antiviral compounds or vaccines have not been proven to be effective against CCHF virus in controlled trials. A vaccine developed in Bulgaria and used there since 1974 is based on CCHF virus derived from suckling mouse brain and inactivation by chloroform.8 The vaccine elicited both cell-mediated and humoral immunity, but multiple doses were required before neutralisation activity was observed; even then the activity was low.9 To date, there are no controlled efficacy studies and the vaccine is unlicensed by the European Medicines Agency or the US Food and Drug Administration. Due to its crude preparation, it is unlikely to gain widespread international regulatory approval.

Recent vaccine approaches for CCHF include a DNA-based vaccine expressing the glycoprotein-encoding region of the virus, which induced neutralising antibodies in approximately half of vaccinated mice.10 Another vaccine candidate used transgenic tobacco leaves expressing the CCHF viral glycoproteins, which were fed to mice and induced both IgG and IgA.11 However, neutralisation activity was not tested and neither vaccine approach has been tested for protection against lethal disease using a challenge model, so efficacy has not been assessed. The most promising CCHF vaccine candidate published to date is a Modified Vaccinia Ankara (MVA) vector expressing the full-length glycoproteins which induced humoral and cellular immunity, along with protection in an adult small animal model of CCHF virus infection.12

The genome of CCHF virus is distributed over 3 RNA segments: small (S), medium (M) and large (L) which encode the viral nucleoprotein (NP), glycoprotein and RNA polymerase, respectively. While all the vaccine reports published to date have...
focused on the M segment, there is compelling evidence that a vaccine based on the S segment would be a feasible alternative. The NP is recognized as the predominant antigen, inducing a high immune response in most Bunyavirus infections; it is also highly conserved between strains. Additionally, the NP has been used as an antigenic target for vaccines that have demonstrated protective effects in a range of viral diseases (Table 1). Of particular interest is the protective effect that the NP antigen has shown against 2 other viruses of the same Bunyaviridae family of which CCHF virus is a member: Hantavirus and Rift Valley fever virus.

The NP of CCHF virus consists of a large, globular domain, plus a protrusion that contains a conserved caspase-3 cleavage site. The globular region is responsible for RNA binding, while the role of the caspase-3 cleavage site is currently unclear. It has been shown that the nucleoprotein is cleaved in apoptotic cells at later stages of infection, and that it may play a regulatory role as RNA polymerase is increased when cleavage is disrupted. NP formation is essential for virus multiplication and therefore represents a potential vaccination target.

This report documents the incorporation of the CCHF virus S segment in a Modified Vaccinia virus Ankara (MVA) vector. The vaccine candidate was then tested for immunogenicity and efficacy using murine models.

Results

In vitro expression of MVA-NP constructs

In order to verify the proper expression of the inserted CCHF NP, MVA-NP3010 was used in a Western blot assay. Using an anti-V5 antibody to ascertain the molecular size of the inserted protein, a band of approximately 62.5 kDa was observed. This was consistent with the estimated size of the CCHF NP of 52 kDa, plus the V5 tag and tPA regions.

Immunogenicity of MVA-NP

Effects of type-1 interferon receptor deficiency on vaccine induced immunity

Immunogenicity studies used A129 and 129Sv/Ev mouse strains to represent a susceptible CCHF host and the parent wild-type strain, respectively. To assess whether the type-1 interferon receptor deficiency possessed by the A129 mice affected the vaccine-induced immune responses, both strains of mice were immunised with MVA-NP3010 vaccine. As observed in Figure 1, MVA-NP3010 induced similar numbers of IFN-γ secreting cells specific to peptides derived from CCHF NP in both strains of mice (P > 0.05, Mann-Whitney statistical test). Responses to the tPA and V5 tags were similarly low in both groups, demonstrating the specificity of the response to the inserted protein. Furthermore, as shown in Figure 2, Western blot analysis revealed that all (5/5) A129 mice developed an antibody response specific to NP as did all 129Sv/Ev mice (5/5). Therefore, a deficiency in the type-1 interferon receptor did not affect the induction of humoral or cell-mediated immune responses of the MVA-NP3010 vaccine in these assays.

Immunogenicity of MVA-NP10200

To determine the efficacy of the vaccine, a challenge model developed for infection with strain IbAr10200 of CCHF virus was planned. Therefore, to

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Table 1. Summary of the vaccines against viral diseases reported that have the viral nucleoprotein as the sole target antigen

| Pathogen          | Vaccine construct                        | Protection effects                                | Ref |
|-------------------|-----------------------------------------|--------------------------------------------------|-----|
| Ebola virus       | Venezuelan equine encephalitis virus replicons | Protection in C57BL/6 mice                        | [20]|
|                   | Cytomegalovirus                         | Protection in mice                                | [21]|
| Hantavirus        | Recombinant vaccinia virus               | Partial protection in Mongolian gerbils           | [22]|
|                   |                                        | Protection in mice                                | [23]|
| Influenza virus   | DNA prime and recombinant adenovirus boost | Protection in mice                                | [24]|
|                   | Recombinant adenovirus                  | Protection in mice                                | [25]|
| Lassa virus       | Recombinant vaccinia virus               | Protection in mice                                | [26]|
|                   |                                        | Protection from encephalitis in rats              | [27]|
| Measles           | Recombinant vaccinia virus               | Partial protection of mice against lethal challenge | [28]|
| Pichinde virus    | Recombinant vaccinia virus               |                                                  |     |
| Rabies virus      | Raccoon poxvirus                        |                                                  |     |
| Rift Valley fever virus | DNA vaccine                              |                                                  |     |

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Figure 1. IFN-γ ELISpot responses from A129 and 129Sv/Ev mice vaccinated with MVA-NP3010 (black), MVA-1974 (gray) or saline (white). Splenocytes from vaccinated mice were restimulated with peptides derived from the CCHFv nucleoprotein split into 2 pools and summed, or a single pool containing peptides from the tPA and V5 fusion partners. Mean ± SEM is plotted.
develop a homologous vaccine the NP insert was changed to the equivalent region of this strain. To ensure that the new construct conferred similar immune responses in vaccinated animals, A129 mice were vaccinated with 2 doses separated by 2 weeks, and then 2 weeks after the final dose 3 animals were culled. As shown in Figure 3, a NP-specific T-cell response was detected in all of the vaccinated animals. Additionally, as demonstrated in Figure 4, NP-specific antibodies were detected in all of the animals tested. Therefore, MVA-NP10200 was taken forward for testing in efficacy studies.

**Efficacy of MVA-NP**

**Survival**

A129 mice (n = 9 per group) were vaccinated with 2 doses of MVA-NP10200 spaced at 2 weeks intervals. At a time point 2 weeks after the final vaccination, they were challenged with a lethal dose of CCHF virus. As seen in Figure 5, despite the induction of NP-specific immunity prior to challenge, all mice succumbed to the lethal infection between days 4 and 5 post-challenge.

**Viral load**

To assess whether the MVA-NP10200 vaccine gave any reduction in viral load in the target sites of CCHF viral replication, viral load analysis using RT-PCR was conducted. Blood, spleen and liver were collected 4 d post-challenge from animals challenged with MVA-1974 and MVA-NP10200. As seen in Figure 6, there were no observable differences between animals challenged with the empty vector versus those that were immunised with the CCHF vaccine candidate.
Histopathology

In the spleen, patchy to diffuse infiltration of parenchyma by macrophages was noted, primarily involving the red pulp, with varying degrees of effacement of white pulp (Table 2). In addition, lymphocyte loss and apoptosis were observed, with the latter characterized by the presence of tingible body macrophages and apoptotic bodies. Using immunohistochemistry, positively staining cells, consistent with macrophages, were observed, diffusely scattered throughout the parenchyma, primarily within the red pulp.

Table 2. Severity of spleen and liver lesions in challenged mice: distribution in treatment groups

| Group        | Microscopic lesion          | Severity | Saline  | MVA-1974 | MVA-NP10200 |
|--------------|-----------------------------|----------|---------|----------|-------------|
| Spleen:      | Infiltration of red and white pulp by macrophages. | Normal | 0       | 0        | 0           |
|              | Lymphocyte apoptosis/necrosis. | Minimal | 1       | 1        | 0           |
| Liver:       | Focal hepatocyte necrosis   | Minimal | 0       | 0        | 0           |
|              | Mixed inflammatory cell infiltrate | Mild    | 2       | 0        | 1           |
|              |                             | Moderate | 3       | 3        | 2           |

In the liver, changes comprised multifocal, hepatocyte necrosis, characterized by cytoplasmic eosinophilia and nuclear pyknosis and accompanied frequently with a mixed inflammatory cell infiltration, mainly of polymorphonuclear leukocytes (Table 2). Positively stained hepatocytes, detected by immunohistochemistry, were observed scattered throughout the parenchyma.

Discussion

To elicit an immune response against the CCHF viral NP, a Modified Vaccinia virus Ankara (MVA) viral vector was used. MVA is one of the most advanced recombinant poxviral vaccine vectors used in human clinical trials, and elicits both humoral and cellular immune responses. This latter point is particularly pertinent as there is no defined correlate of protection against CCHF virus so the priming of both arms of the immune system may offer the best opportunity to observe protective effects. The results of these studies confirmed the induction of both antibody and cell-mediated immunity and it is noteworthy that the Bulgarian vaccine based on suckling mouse brain (inactivated by chloroform, heated at 58°C, and absorbed on aluminum hydroxide) also induced T-cell and humoral immunity, but efficacy of this vaccine has yet to be tested in animal models. For our studies, a homologous prime-boost approach was undertaken as we have demonstrated that this induced increased numbers of antigen-specific T cells compared to a single dose (data not shown). This finding is in line with others who have used similar approaches with MVA-based vaccines against Mycobacterium tuberculosis, and human immunodeficiency virus (HIV). Repeat administration with MVA allows reboosting of responses, despite induction of cellular and humoral immune responses against the vector. This has been reported in Phase I/II therapeutic cancer vaccine trial, as well as in a Phase I HIV vaccine trial.
This use of NP is in contrast to others who have opted to use the envelope glycoprotein as the candidate antigenic target. While the external location of the glycoprotein makes it a favorable target for the induction of neutralising antibody, it has been recognized that there is not a strict correlation between in vitro neutralisation and in vivo protection of CCHF virus-specific antibodies. The NP was considered to be an appropriate vaccine antigen, due to several characteristics. The NP in Bunyavirus infection has been recognized as the predominant antigen, inducing a high immune response, and after challenge with CCHF, it has been shown that most antibodies are directed to NP.

Despite the induction of humoral responses for recent CCHF vaccine candidates, neither vaccine approach was tested for efficacy, presumably due to the lack of a suitable animal model at that time. In 2010, 2 murine models susceptible to CCHF virus were published, with deletions in either STAT-1 or the type-I interferon receptor. The STAT-1 knockout mice exhibit signaling defects in their response to all 3 major types of interferon (type I, IFN-α and IFN-β; type II, IFN-γ; and type III, IFN-λ) that leads to a complete abolishment of the intracellular interferon response. For the efficacy studies with the MVA-NP10200 vaccine candidate, mice deficient in the type-I interferon receptor were used since these have less immune deficiency. This was considered essential for testing vaccination approaches as IFN-γ is a major cytokine involved in the adaptive immune response. For the immunogenicity of the MVA-NP vaccine, we compared the immune responses in mice with the type-I interferon receptor deficiency and the parental wild-type strain. No differences were observed in either the antibody or cell-mediated response, demonstrating that the knockout mice elicited similar responses. This was unsurprising, as others have also reported similar findings in studies with dengue virus. The type-I IFN receptor knockout mice are also valuable for studying the efficacy of vaccines, as has been shown with those against Chikungunya virus, Bluetongue virus, Vaccinia virus, and African Horse Sickness virus. Therefore, our finding that the CCHF MVA-NP vaccine did not demonstrate any protective effects seems unlikely to be a consequence of the animal model utilised.

In the present studies, antibody-induced responses using Western Blot analysis using virus infected cell lysate and recombinant CCHF viral nucleoprotein as antigen were assessed. While this did not allow us to quantify antibody levels or elucidate the subclass of immunoglobulin, it did demonstrate specific antibody recognition of the protein target. Due to the NP being internally located, its main effects in viral immunity are through T lymphocytes. However, although it recognized that antibodies against viral NP are often poor at neutralisation, others effects include complement-mediated cytolysis, increased T cell responses associated with enhanced dendritic cell function, and reduced viral replication in culture. The protective role of CCHF immunoglobulin is currently unknown, as although immune globulin therapy has been administered on several occasions its efficacy has still not been assessed in a randomized clinical trial. Should the vaccine have demonstrated protective effects, further work would have been warranted in deciphering the immune response by depletion or transfer experiments. Similarly, while the T-cell response looked at IFN-γ recall responses after stimulation with overlapping peptide pools, further work to identify the responding cell phenotypes, function and cytokine/chemokine secretory patterns could have been conducted had the vaccine shown any positive effect.

To our knowledge, this is the first report of a vaccine against CCHF virus based on the nucleoprotein. The MVA-NP vaccine candidate demonstrated antigen-specific immunogenicity in mice, but failed to exert any protective effects upon challenge with CCHF virus. This demonstrates that with the lack of any immune correlates of protection, vaccines against CCHF virus will need to demonstrate protection in a lethal dose model before protective efficacy can be established.

**Materials and Methods**

**Cells**

BHK-21 cells (American Type Culture Collection, USA) were cultured in modified essential eagle medium (Sigma, UK) supplemented with 10% foetal bovine serum, 2 mM L-glutamine, 100 U penicillin and 0.1 mg/ml streptomycin (Sigma). Chick Embryonic Fibroblast (CEF) cells (Institute for Animal Health, UK) were cultured in Dulbecco’s modified eagle medium (Sigma) supplemented as above. SW13 and VeroE6 cells (European Collection of Cell Cultures, UK) were maintained in Leibovitz’s L-15 medium containing Glutamax (Life Technologies, UK) supplemented with 10% foetal bovine serum (Sigma).

**Viruses**

MVA strain 1974/NIH clone 1 (kindly supplied by Prof B. Moss, NIH) was used for the vaccine construct. Virus titer was determined by plaque assay in BHK-21 cells. CCHF virus strain IbAr10200 was prepared from suckling mouse brain homogenate. Titer was determined by TCID₅₀ in VeroE6 cells.

**Animals**

A129 (IFN-α/βR⁻/⁻) or 129Sv/Ev (both from B&K Universal, UK) aged 5–8 weeks were used. Animal studies were approved by the ethical review process of Public Health England, UK and the UK Home Office, via project licenses. All work involving animals was performed in accordance with the United Kingdom Animals (Scientific Procedures) Act 1986.

**Construction of plasmids**

Plasmid pLW-44 (kindly provided by Prof B. Moss, NIH) encoded the green fluorescence protein (GFP) reporter gene under control of the p11 promoter and an expression cassette controlled by the artificial promoter, mH5, which allowed constitutive expression of heterologous genes. The mH5 promoter demonstrates increased stability. Plasmid pDEST44-TPA-V5 was derived by inserting a cassette of pLW-44. The cassette contained Gateway system attR recombination sequences (Life Technologies, UK) flanked by the human tissue plasminogen activator.
(tPA) leader sequence to increase secretion and neutralising antibody induction,\(^{62, 63}\) and a C-terminal V5 tag between the XmaI and SalI restriction sites for in vitro immunodetection.

The NP open reading frames from the S segments of CCHF strains 3010 (Accession number DQ099335) and IbAr10200 (Accession number NC_005302) were used in this work to generate plasmid pENTR-NP (pENTR-NP3010 and pENTR-NP10200, respectively).

Plasmid pDEST44-TPA-V5 was recombined with pENTR-NP using Gateway technology\(^{64}\) to generate plasmid pTP-NP. The resulting plasmids encoded the respective tPA-NP-V5 fusion proteins downstream of the poxvirus mH5 promoter.

Generation and characterization of recombinant MVA expressing CCHFv nucleoprotein

BHK-21 cells were infected with MVA at a multiplicity of infection of 0.05. Infected cells were transfected with pTP-NP using Lipofectamine (Life Technologies) as directed by the manufacturer. Resulting recombinant MVA-NP was serially plaque-purified 4 times in BHK-21 cells, based on GFP expression. MVA-NP was amplified on BHK-21 and CEF cells, purified by sucrose cushion centrifugation,\(^{65}\) and titrated by plaque assay on BHK-21 cells, prior to use in in vivo studies. Plaques were visualised using GFP fluorescence or by immunostaining.\(^{66}\) with rabbit anti-Vaccinia antibody (AbD Serotec, UK) and Vectastain Universal ABC-AP kit (Vector Laboratories, USA). Genomic DNA from infected cells was extracted using the Wizard SV genomic DNA purification system (Promega, UK) and used as a template for PCR with AccuPrime Taq DNA polymerase High Fidelity (Life Technologies, UK). Quality control testing to ensure expression of the insert proteins was confirmed in the resulting MVA-NP3010 and MVA-NP10200 by and Western Blotting and quantity of MVA evaluated using plaque assays in BHK-21 cells.

MVA-NP vaccination

Groups of 5–12 mice were injected into the caudal aspect of the proximal hindlimb musculature with 10⁷ plaque-forming units (pfu) per animal of MVA-NP diluted in endotoxin-free PBS. A total volume of 100 μl was delivered equally across 2 sites. Animals received a booster vaccination 14 d later. Control animals received 10⁷ pfu of non-recombinant MVA 1974 or an equivalent volume of saline. Animals were euthanised and tissues were collected 7 or 14 d after the final vaccination.

Interferon-gamma (IFN-γ) ELISpot assay

Splenocytes were assessed for antigen recall response via IFN-γ ELISpot (Mabtech, Sweden), performed as per the manufacturer’s instructions. Cells were seeded in PVDF microtitre plates at 2 × 10⁶ splenocytes per well and re-stimulated with peptide pools (Mimotopes, Australia). Overlapping peptides spanning the length of the CCHF virus nucleoprotein consisting of 20mers, offset by 8 residues, were applied at a final concentration of 25 μg/ml per peptide in pools of 28–32 peptides. Plates were developed after 18 hours at 37°C in a humidified incubator supplemented with 5% CO₂. Spots were counted visually on an automated ELISpot reader (Autoimmun Diagnostika GmbH, Germany). Background values from wells containing medium but no peptides were subtracted and pools were summed across the target protein. Results were expressed as spot forming units (SFU) per 10⁶ cells.

Antibody testing by Western Blot

SW13 cell monolayers were infected with CCHF virus strain IbAr10200 at a multiplicity of infection (MOI) of approximately 0.01, and incubated at 37°C in Leibovitz’s L-15 medium containing 2% foetal bovine serum. 48 hours post-infection, the medium was removed and the cells were treated with Laemmli buffer supplemented to contain 10% sodium dodecyl sulfate (SDS) (Sigma). The resultant mixture was collected into vials and heat treated at 90°C for 10 minutes before use in Western Blot analysis. Uninfected SW13 monolayers were treated similarly for use as a negative control.

Lysates from CCHFv-infected or uninfected SW13 cells were subjected to SDS-PAGE on a 4–12% Bis-Tris gel (Life Technologies) and transferred to a PVDF membrane. After blocking in 5% milk protein, membranes were incubated with mouse serum for 2 hours, washed 6 times with PBS containing 0.05% NP40, incubated for 1 hour with HRP-conjugated rabbit anti-mouse IgG (Sigma) or goat anti-mouse IgG/A/M (AbD Serotec) and washed as before. All antibody dilutions were made in PBS containing 0.05% NP40 and 5% milk protein. Bound antibody was detected with ECL-Prime WB detection reagent (GE Life Sciences, UK) according to the manufacturer’s directions and visualised on a ChemiDoc system (BioRad, UK). Molecular weights were calculated by comparison with markers of known molecular weight using QuantityOne software.

CCHF virus challenge in A129 mice

Fourteen days after the final vaccination with MVA-NP or a control substance, A129 mice received 200 TCID₅₀ CCHF virus strain IbAr10200 intradermally in the midline of the lumbar region in a volume of 100 μl divided equally across 2 sites. 50 μl is the maximum recommended volume for intradermal inoculation of mice.\(^{67}\) and confirmation of intradermal delivery was seen by a visible bleb formation under the skin. Post challenge, animals were weighed and body temperature measured daily by a subcutaneously located temperature chip. In addition, they were observed for clinical signs of disease twice daily (arching, ruffled fur, lethargy and immobility). Criteria for euthanasia on welfare grounds consisted of 20% weight loss or observation of 2, abnormal clinical signs. At 4 d post-challenge, randomly selected animals were euthanised and samples of blood, spleen and liver collected for viral load studies. Spleen and liver samples were also collected for histopathological examination.

Viral load analysis

Whole blood (100 μl) was collected into RNA Protect Animal Blood tubes (Qiagen) and stored at −80°C. Tubes were thawed, inverted and left for a further 2 hours at room temperature to ensure efficient cell lysis. Samples were treated with Red
Blood Cell Lysis Solution (Miltenyi Biotec) before purification of total RNA using an RNaseasy Mini kit (Qiagen). For viral load analysis, spleen and liver samples were collected into RNALater (Qiagen) and stored at −80°C. Thawed tissue was transferred to RLT buffer (Qiagen), homogenized by passing through a 70 μm sieve and then treated using an RNaseasy Mini kit (Qiagen) for extraction of total RNA.

CCHFv S segment was detected by RT-PCR on the ABI 7500 RT-PCR platform as described (Atkinson et al, 2012), with cycling conditions adjusted to those described in the Quantifast probe assay: 50°C for 20 min, 96°C for 5 min, followed by 45 cycles of 95°C for 15 sec and 60°C for 30 sec (with quantification analysis of fluorescence performed at the end of each 60°C step), and final cooling of 40°C for 30 sec. A synthetic S segment of known concentration was used to quantify S segment copy number in blood and tissue samples. All reactions were run in triplicate.

To normalize the CCHFv expression data, the hypoxanthine guanine phosphoribosyl transferase (HPRT) housekeeping gene was used. A one-step RT-PCR with singleplex detection was performed targeting an 89 bp product in the mouse HPRT gene (NCBI Reference sequence NM_013556) using the QuantiFast probe assay (Qiagen) and the ABI 7500 RT-PCR platform. CT values for CCHFv and HPRT were each inverted by subtracting the CT value from 45 (the total number of cycles), where CT is the number of cycles to reach the fluorescence threshold value. The mean value of CCHFv was then divided by the mean value of the HPRT reference gene for each sample.

**Histopathology studies**

Samples of spleen and liver were placed in 10% neutral buffered formalin for 7 days, processed routinely to paraffin wax, sections cut at 3–5 μm, stained with haema-toxylin and eosin (H&E) and examined microscopically. Lesions referable to infection with CCHF virus were scored subjectively using the following scale: normal, minimal, mild, moderate and marked.

For immunohistochemistry, formalin-fixed, paraffin-embedded sections of spleen and liver, cut between 3–5 μm, were mounted on positively charged X-tra Adhesive slides (Leica Biosystems, UK), deparaffinised and rehydrated. Immunohistochemical staining was achieved using a BOND-MAX Immunostainer (Leica Microsystems, UK) and a Novocastra Bond Intense R (Leica Biosystems) detection kit. A heat-induced epitope retrieval cycle with buffer ER1 (Leica Biosystems) was performed for 20 minutes. Slides were incubated with rabbit serum (4%) (Abcam, Cambridge, UK) for 20 minutes followed by an avidin/biotin blocking stage (15 minutes each) (Abcam). Polyclonal antibody raised in sheep immunised against recombinant CCHFv nucleoprotein (kindly provided by Dr John Barr, University of Leeds, UK) was incubated with the tissue for 30 minutes, followed by a biotinylated rabbit anti-sheep polyclonal antibody (Abcam) at a dilution of 1500, for 10 minutes. Haematoxylin was used as the counterstain. Positive and negative control slides were included. Immunolabelled slides were evaluated using light microscopy.

**Disclosure of Potential Conflicts of Interest**

No potential conflicts of interest were disclosed.

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