Common midwife toad ranaviruses replicate first in the oral cavity of smooth newts (*Lissotriton vulgaris*) and show distinct strain-associated pathogenicity

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*Ranavirus* is the second most common infectious cause of amphibian mortality. These viruses affect caudates, an order in which information regarding *Ranavirus* pathogenesis is scarce. In the Netherlands, two strains (CMTV-NL I and III) were suspected to possess distinct pathogenicity based on field data. To investigate susceptibility and disease progression in urodeles and determine differences in pathogenicity between strains, 45 adult smooth newts (*Lissotriton vulgaris*) were challenged via bath exposure with these ranaviruses and their detection in organs and feces followed over time by PCR, immunohistochemistry and *in situ* hybridization. *Ranavirus* was first detected at 3 days post infection (p.i.) in the oral cavity and upper respiratory mucosa. At 6 days p.i., virus was found in connective tissues and vasculature of the gastrointestinal tract. Finally, from 9 days p.i onwards there was widespread *Ranavirus* disease in various organs including skin, kidneys and gonads. Higher pathogenicity of the CMTV-NL I strain was confirmed by higher correlation coefficient of experimental group and mortality of challenged animals. *Ranavirus*-exposed smooth newts shed virus in feces intermittently and infection was seen in the absence of lesions or clinical signs, indicating that this species can harbor subclinical infections and potentially serve as disease reservoirs.

The common midwife toad virus (CMTV) is a *Ranavirus* (family *Iridoviridae*), initially reported in Spain in 2008. *Ranaviruses* clustering genetically within the common midwife toad clade were subsequently recorded as the cause of mortality events involving various amphibian and reptile species throughout Europe and Asia.

In the north of the Netherlands (NL), CMTV was the cause of an amphibian mass mortality event in 2010 at Dwingelderveld National Park (DNP). Since then, over half of the amphibian species native to the Netherlands have been affected by CMTV ranaviruses; particularly water frogs (*Pelophylax kl. esculentus*) and smooth newts (*Lissotriton vulgaris*). Macroscopic lesions consistently observed across affected amphibian host species included skin hemorrhages and edema. Histology revealed widespread necrosis and the presence of intracytoplasmic inclusion bodies.

Molecular tests revealed that in addition to the northern CMTV *Ranavirus* strain, two other phylogenetically distinct CMTV *Ranavirus* strains occurred in the east and south of the Netherlands. Dutch *Ranavirus* strains were designated as CMTV-NL I, II and III. Based on gene truncations, field mortality patterns and changes in animal abundance after mortality events, the strain from the north (CMTV-NL I) appeared to be more pathogenic phenotype than the southern strain (CMTV-NL III). From this, it was previously concluded that CMTV
NL-I, the more virulent strain, was recently introduced into the Netherlands and that CMTV NL-III was potentially endemic with attenuated virulence10,11.

The other Dutch strain (CMTV-NL II) was obtained from a pond within a private property in the centre-east of the country. No field data was available for this strain. CMTV NL-II was not included in this experiment.

The smooth newt is one of the amphibians commonly affected by CMTV-NL ranaviruses. Juvenile smooth newts can quickly colonize new habitats and disperse hundreds of meters from breeding to overwintering sites within a year12. They are often found in the same waterbodies as water frogs (Pelophylax spp.) which have suffered population declines due to Ranavirus infection10. All of these characteristics make them potential Ranavirus hosts, and if so, potential spreading vectors. Subclinical Ranavirus infections have been recorded in other caudate species: adult tiger salamanders introduce Ambystoma tigrinum virus (ATV) into waterbodies when returning to breeding sites after hibernation or when they are used as bait13,14.

Recent research on Ranavirus pathogenesis, tissue tropism, and dose-dependent susceptibility has focused on anurans15. In contrast, most of the experimental work regarding caudates is limited primarily to distinct aspects of Ranavirus strain pathogenicity or the influence of environmental factors on the development of disease16.

In order to study the susceptibility of smooth newts to CMTV-NL Ranavirus disease, to determine whether CMTV NL Ranavirus strains differ in pathogenicity, and to find out whether this species can harbor subclinical infections and excrete virus, smooth newts were challenged with Dutch CMTV isolates. The smooth newt was chosen as an experimental model due to field data on its susceptibility to CMTV-NL10. Through this experiment, the pathogenesis and fecal shedding of CMTV-NL ranaviruses were studied by tracing the virus within the tissues and feces of the hosts over time. The pathogenicity of CMTV-NL strains I and III was compared by characterizing lesion severity through semi-quantitative scoring.

**Results**

**Smooth newts are susceptible to infection with both CMTV-NL Ranavirus strains.** Adults newts captured in the wild were challenged with two CMTV-NL strains or medium and euthanized at 5 post-infection (p.i.) time points to study disease progression. The experimental set up is schematically illustrated in Fig. 1.

Smooth newts in all experimental groups experienced a minor loss of weight by the end of the trial, but weight loss did not vary by treatment (Supplementary Table S1.; ANOVA; p = 0.6). All animals from the control group remained healthy until the end of the experiment.

Three of the CMTV-NL I-exposed animals had mild erythema and/or hemorrhages in the mouth and gular region, lesions were evident from day 9 p.i. and persisted until the end of the experiment (Fig. 2), while only one CMTV-NL III-exposed animal exhibited mouth erythema at day 7 p.i. Sudden death without clinical signs...
occurred in three CMTV-NL I-exposed newts, at days 7, 9 and 10 p.i., while none of the CMTV-NL III-exposed animals died or reached humane end-points before the end of the experiment.

**Ranavirus first replicates in the upper alimentary tract of smooth newts.** All smooth newt tissues sampled at 6 hours p.i. tested negative for infection using *in situ* hybridization ISH. However, viral RNA was first detected at 3 days p.i. by ISH staining in the upper respiratory tract mucosa, oral mucosa, and stomach contents of a CMTV-NL I-infected animal. (Fig. 3). ISH showed evidence of infection in multiple tissues at 6 and 9 days p.i., (Table 1).
Lesions are more severe in newts infected with CMTV-NL I. Pathogenicity of the CMTV-NL Ranavirus strains were evaluated through semi-quantitative scoring based on a previously established system\(^\text{16}\) which is detailed further in the Materials and Methods section.

Results showed that lesions were more severe in animals from Group 1 infected with CMTV-NL I Ranavirus, than in Group 2 animals infected with the CMTV-NL III Ranavirus (Fig. 5). In animals infected with CMTV-NL I Ranavirus, inclusions, necrosis and vascular damage were consistently present at days 9 and 12. The only consistent finding in animals infected with CMTV-NL III Ranavirus on those days was vascular damage. In the liver of CMTV-NL I Ranavirus-infected animals, IHC positive signal increased over time along with the evidence of necrosis and inclusions bodies. These changes were observed in the cytoplasm of hepatocytes and in the sinusoids. In contrast the IHC positive signal in CMTV-NL III Ranavirus infected animals remained scarce, mostly

| CMTV-NL I | 6 h p.i. | 3 d p.i. | 6 d p.i. | 9 d p.i. | 12 d p.i. |
|-----------|---------|---------|---------|--------|---------|
| Mouth     | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |
| Esophagus | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |
| Stomach   | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |
| Spleen    | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |
| Kidney    | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |
| Pancreas  | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |
| Heart     | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |
| Liver     | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |
| Intestine | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |
| Skin      | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |

| CMTV-NL III | 6 h p.i. | 3 d p.i. | 6 d p.i. | 9 d p.i. | 12 d p.i. |
|-------------|---------|---------|---------|--------|---------|
| Mouth       | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |
| Esophagus   | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |
| Stomach     | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |
| Spleen      | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |
| Kidney      | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |
| Pancreas    | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |
| Heart       | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |
| Liver       | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |
| Intestine   | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |
| Skin        | ISH 1ND,2– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND | ISH 1ND,1– 3– 3ND |

Table 1. ISH, IHC, and PCR results of Ranavirus-infected animals. Animals from the 6 hours p.i., 3 days p.i. and the control group tested negative for all tests in all organs. Positive (+), negative (−), organ not done (ND) * signal present only in the lumen.

Positive immunohistochemistry (IHC) staining indicative of viral protein production was not observed until 6 days p.i., when mild single cell staining was detected in the liver, esophagus, oral mucosa and upper respiratory tract mucosa of both Ranavirus-exposed experimental groups. Positive staining also occurred in splenic macrophages, blood vessels of the heart and connective tissues of the stomach and gonads in CMTV-NL I-exposed animals. By day 9 p.i., an IHC signal was still present in the previously positive connective tissues of the reproductive tract and liver, and had spread to the skin, intestine, pancreas, and kidney in affected animals from both exposed groups. At day 12 p.i. viral proteins were still present in most tissues (upper respiratory and mouth mucosa, stomach, intestine, liver, connective tissues of the gonads and in the skin) of animals from both Ranavirus-infected groups (Fig. 3). However, viral proteins could no longer be detected in the kidney, spleen, heart and pancreas (Table 1). Virus protein production was not observed at any time point in the peripheral or central nervous system of CMTV-NL infected smooth newts.

None of the control animals tested Ranavirus positive at any of the time points using PCR and they all tested PCR-positive for the control gene β-actin (Supplementary Fig. S1). Ranavirus DNA was detected by PCR at the earliest time point (6 hours p.i.) in intestine samples of both Ranavirus-infected groups and in the skin of one CMTV-NL III-infected animal. At day 3 p.i., DNA was detected in the liver of animals from both Ranavirus-infected groups, while 6, 9 and 12 days p.i., DNA was present in the liver and intestine of both Ranavirus-infected groups, but only the skin of CMTV-NL I-exposed animals tested positive 9 and 12 days p.i. (Fig. 4, Table 1/Supplementary Fig. S2)
limited to sinusoids. Histopathology revealed an increase in the number of melano-macrophages, but necrosis and inclusion bodies were rarely observed (Fig. 6).

The Spearman rank correlation method was used to evaluate the lesion severity scores. It revealed high correlation coefficients among the lesion scores in CMTV-NL I-infected animals (all $>0.70$; Supplementary Table S2); however, due to multiple ties, it was not possible to calculate reliable p-values (Supplementary Table S2).

No inclusion bodies were observed in animals from the control group. The results of the semi-quantitative scoring are available in Supplementary Table S3.

Smooth newts infected with CMTV-NL-I Ranavirus shed virus intermittently through the feces. Ranavirus DNA was found in the feces of both CMTV-NL Ranavirus-infected animals at day 2 p.i. At day 3 p.i., none of the feces tested positive for Ranavirus DNA. Feces of CMTV-NL I-exposed animals tested positive intermittently and in a few animals repeatedly from day 4 to 10 p.i. In contrast, feces from CMTV-NL III-exposed animals tested positive only once from day 5 to 11 p.i. (Fig. 7). Control animals always tested PCR negative for Ranavirus.

Discussion

The aims of this study were to determine the susceptibility of smooth newts to CMTV-NL Ranavirus infection, the pathogenesis of Ranavirus disease in these hosts, the difference in pathogenicity between two closely related Dutch CMTV Ranavirus strains and the ability of smooth newts to excrete Ranavirus and harbor subclinical infections.

Pathology, PCR, ISH and IHC, all illustrate how smooth newts are susceptible to CMTV-NL Ranavirus across tissues and that the virus first targets the oral cavity and upper respiratory mucosa. Subsequently, gastrointestinal invasion likely occurs through the vasculature as indicated by the sites of immunolabelling (connective tissues and endothelial cells). Finally, there is wide-spread involvement of numerous organs including kidneys, skin and gonads.

The ISH staining of the gastric contents shows that smooth newts in this study likely became infected by ingestion. This mode of transmission has long been recognized as highly efficient for ranaviruses$^{15,17,18}$. The initial virus replication in the mouth region of these newts and the subsequent dissemination through connective tissues and blood vessels probably explains how the virus could avoid highly damaging physiological barriers like low gastric pH$^{19}$.

Experimental studies with a caudate-specific Ranavirus (Ambystoma tigrinum virus) have previously shown that skin to skin contact is also a plausible route of infection in tiger salamanders (Ambystoma tigrinum)$^{17}$. However, in this study, Ranavirus skin involvement in most Ranavirus-exposed animals was only evident at the latest stages of viral infection.

The efficiency of the skin barrier against Ranavirus transmission has been shown to depend on several factors, including the amount of antimicrobial peptides produced$^{20}$ and the diversity of the host skin microbiome$^{21}$. Alterations to the previous factors likely result in disruption of skin integrity and probably facilitate invasion of the pathogen and the development of disease.

Severity of disease was highest in animals infected with CMTV-NL I as seen by the type of lesions and death of some animals prior to euthanasia. These results are consistent with the previous hypothesis that the CMTV-NL
IRanavirus is the most pathogenic of the Dutch Ranavirus strains. This likely explains the association of this strain to the on-going high mortality involving over half of the endemic amphibian species in the Netherlands, and the local declines in populations of water frogs (Pelophylax spp) in natural and artificial water bodies. In contrast, CMTV-NL III Ranavirus exposed animals did not experience mortality prior to the designated euthanasia time point and even though infection was confirmed in most animals, the overall lesion severity was low. This indicates attenuated virulence of the strain, which had previously been suggested by low Ranavirus-associated mortality in areas where the strain occurs and by molecular data showing truncations of virulence-associated genes.

Regarding Ranavirus-excretion by infected smooth newts, evidence of viral DNA was observed in the feces of both infected animal groups. It is possible that the initial excretion at day 2 p.i. and DNA intestinal tissue detection at 6 hours p.i., corresponded to virus inoculated during the challenge and did not necessarily reflect replication.
Figure 6. Comparison of lesion severity in the liver of smooth newts infected with CMTV-NL *Ranavirus* strains over time. (A, B) CMTV-NL I *Ranavirus*-infected smooth newt with a small area of necrosis (square) Hematoxylin and Eosine, and positive ISH immunolabelling of sinusoids (green arrow and circle) and hepatocytes (black arrow and circle). C and D) CMTV-NL III *Ranavirus*-infected smooth with no changes in Hematoxylin and Eosine, ISH immunolabelling restricted to sinusoids. (E, F) CMTV-NL I *Ranavirus*-infected smooth newt with numerous intracytoplasmic inclusion bodies (arrows) in hepatocytes, Hematoxylin and Eosine. Marked IHC immunolabelling of hepatocytes (black arrow and circle) and in sinusoids (green arrow and circle) (G, H) CMTV-NL III *Ranavirus*-infected smooth newt at 9 days p.i. with mild increase in the number of melano-macrophages, Hematoxylin and Eosine. Positive IHC immunolabelling is restricted to the cytoplasm of one hepatocyte (black arrow and circle). (I, J) CMTV-NL I *Ranavirus*-infected smooth newt with marked necrosis of the parenchyma (square) and rare presence of inclusion bodies (arrow) Hematoxylin and Eosine. IHC immunolabelling is observed within hepatocytes (black circles). K and L) CMTV-NL III *Ranavirus*-infected smooth newt, No microscopic abnormalities are seen with Hematoxylin and Eosine, IHC immunolabelling is scant and limited to a few sinusoids (green circle) IHC Anti-ECV polyclonal antibody, ISH FV3 probe.

Figure 7. PCR detection of *Ranavirus* in fecal samples of animals from CMTV-NL I and CMTV-NL III *Ranavirus*-infected animals: The X axis shows the p.i. time point of euthanasia, while the Y axis shows the days p.i. Positive *Ranavirus* detection is represented by + sign while negative fecal samples are represented by a − sign. Grey bands represent the day at which the animals were euthanized, while the black bands represent days at which infected animals died prior to reaching the set time point of euthanasia. The letter “N” stands for no feces found. Control animals always tested negative for *Ranavirus* in fecal samples.
This is supported by the positive ISH staining limited to the gastric contents at 3 days p.i. in CMTV-NL I-infected animals and the absence of IHC signal in the intestine of these animals until 9 days p.i. The number of animals with fecal viral DNA was comparable in both exposed groups. However, only CMTV-NL I Ranavirus-infected animals shed virus constantly until the day of euthanasia.

The results also show that Ranavirus-exposed smooth newts intermittently excrete virus in feces and that the virus replicates within tissues in the absence of lesions or clinical signs. This confirms that smooth newts, like other species of caudates\(^{13}\) may harbor subclinical infections and serve as natural reservoir hosts. These findings highlight the importance of increasing the scope of CMTV-NL Ranavirus monitoring efforts, to focus not only on sampling highly susceptible amphibian species or life stages but also amphibian accompanying species which do not show evident Ranavirus-associated clinical signs.

To conclude, this study signals the oral route as an effective and likely common transmission pathway of ranaviruses in smooth newts; showcases distinct pathogenicity of two CMTV-NL Ranavirus isolates and confirms the ability of these animals to harbor subclinical infections, excrete virus in feces and potentially serve as disease reservoirs.

Materials and Methods

Collection and husbandry of animals. All experimental procedures were undertaken at the Institute of Zoology (IoZ), London, UK. The husbandry methods were first approved by the Zoological Society of London Ethics Committee (project's license number: P8897246A) and carried out in accordance to regulations.

A total of 45 adult smooth newts were collected at several natural sites in Cambridge, UK (Longitude: 52.24, Latitude:0.04) and Essex, UK (Longitude:51.58, Latitude 0.50) without a previous history of amphibian mortality or Ranavirus detection during the spring of 2017. Upon arrival to the animal facility at the IoZ, newts were kept in two groups housed in plastic tanks (80 × 40 × 25 cm) filled with soil, reptile hides (Exo Terra, UK), cork bark and Petri dishes with aged tap water. In order to exclude the presence of subclinical infections, a two-month long quarantine period was implemented. Animals were monitored daily and fed a diet consisting primarily of live crikets (Gryllodes sigillatus) dusted with the calcium supplement (Nutrobal, US), every two days.

In vivo infection experiment. After the quarantine period, animals were randomized into three different groups of 15 individuals each in accordance to the three assigned treatments: Group 1: CMTV-NL I (10\(^8\) TCID50), Group 2: CMTV-NL III (10\(^8\) TCID50) and finally the control group: cell medium (L-15). These groups were further split into five subgroups of three animals each based on established post-infection (p.i.) experimental time points: 6 hours, 3, 6, 9 and 12 days p.i. Each animal was transferred to an individual plastic box using a different set of powder-free vinyl gloves. Before inoculation with the treatments, the animals were measured (snout to vent), weighed, and swabbed on the dorsal and ventral skin to exclude subclinical Ranavirus infections acquired in the wild. The swabs were labelled with the identification number of the animal and stored in a −80°C freezer.

The newts were exposed percutaneously to 3 ml of Ranavirus suspended in cell medium (10\(^8\) TCID\(_{50}\)) for 6 hours. The virus was pipetted directly on individual plastic trays and the newts were placed on the virus puddle to ensure full palmo-plantar and abdominal contact. These bath exposure procedures were performed under the UK Home Office license (license P8897246A), and by a trained and licensed technician. Procedures were always performed first on the control animals to avoid cross-contamination. All animals were fed every two days and their water changed daily during the experiment. Feces were collected daily when present and stored in 70% alcohol for PCR analysis.

Once each of the assigned time points was reached, the animals were humanely euthanized by trained staff using a schedule 1 method for amphibians, consisting of an overdose of MS-222 (tricaine methanesulfonate) and confirmation of death through pitting of the brain. Samples of liver, intestine and a hind-foot were collected for Ranavirus PCR analyses and stored in 70% alcohol. The rest of the animal was fixed in 10% formalin for 24 hours and then post-fixed in 70% alcohol.

Histopathology and semi-quantitative scoring. All organ samples fixed in formalin were processed for routine histopathological analysis and stained with hematoxylin/eosin. To compare the pathogenicity of the two virus strains, three hallmark lesions of Ranavirus infection were separately and blindly evaluated in 10 organs of each individual by two veterinary pathologists using an established semi-quantitative scoring system\(^{10}\). These lesions consisted of viral inclusion bodies (scores: 0 none; 1 if <5 per high power field; 2 if 5–10 per high power field, etc).
field; 3 if >10 per high power field), extent of necrosis of epithelial and mesenchymal cells (scores: 0 none; 1 < 5% of organ; 2 5–50% of organ; 3 > 50% of organ), and the level of vascular damage in the form of hemorrhage, vasculitis or thrombosis (scores: 0 none; 1 mild; 2 moderate; 3 if severe). The small size of two organs (spleen and kidney) resulted in loss of the tissue from several animals during histopathology processing. These organs were scored but excluded in the final statistical analyses for lesion severity to avoid over or underestimation of the final average score per animal.

**Statistical analysis.** For each treatment (a combination of both CMTV-NL Ranavirus strain and euthanasia day), there were three animals. Only the 10 organs available for all animals were considered for the analysis. To examine the correlation among lesions within groups, the lesion scores were analyzed with a Spearman rank correlation test, given the small sample size for each treatment at each time point (n = 3).

**In situ hybridization.** In order to evaluate virus tissue tropism several animals from both infected groups were processed for in situ hybridization. Given the presumably higher sensitivity of in situ hybridization over immunohistochemistry, animals belonging to the first two time points (6 hours and 3 days p.i.) were selected along with animals belonging to the 9 days p.i. time point which served as a positive control. The control group was excluded based on negative PCR and immunohistochemistry results. Slides were stained with an in situ hybridization probe (ACBio RNAscope® Probe- V-FV3-orf90R, Cat No. 439991) which, although labeled as a general manufacturer (Advanced Cell Diagnostics, 7707 Gateway Blvd. Newark, CA, 94560, USA) specifications.

**Immunohistochemistry.** For immunohistochemistry, a polyclonal antibody against European catfish virus provided by Anna Toffan (Istituto Zooprofilattico Sperimentalle delle Venezie, Italy) was used. The specificity of the antibody was tested by western blot analysis. Tissues from a PCR Ranavirus-positive and Ranavirus-negative amphibians were used as positive and negative controls respectively. Details of the immunohistochemistry protocol are available elsewhere.

**Ranavirus PCR.** To detect viral DNA in the skin swabs and in intestine and liver, the tissues were processed for conventional PCR using diagnostic primers for the major capsid protein: Forward primer (5′-GACTTGGCCACTTATGAC-3′), reverse primer (5′-GTCTCTCAGAGAGAAA-3′). Primers targeting salamander β-actin: Forward (5′-TGAAACCCAAAGGGAACAGGAAAGAT-3′) and reverse (5′-TACGACGGAGGATACAAGGACAGGAC-3′) were used as an internal control. DNA extraction of the skin swabs was performed using an established protocol. The skin swab from a Ranavirus-infected smooth newt carcass from a separate unrelated experiment conducted at the IoZ was used as a positive control. Fecal samples of each newt were processed for DNA extraction as previously described.

**Data Availability** The majority of the data generated from this work is included in this paper. The rest is readily available from the corresponding author on reasonable request.

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Author Contributions
This study was funded by Utrecht University, Dutch Wildlife Health Center, Zoological Society of London, Cornell University and Consejo Nacional de Ciencia y Tecnologia. B.S. developed the concept of the project and the experimental design, performed the in vivo experiment, performed PCR and immunohistochemical staining, analyzed all the data and wrote the paper. T.W.J. provided assistance with the experimental design and gave open access to the laboratory facilities during the experiment, N.K. provided technical assistance during the experiment and aided in animal husbandry, S.J.R.A., M.J.G. and R.J.C. collected the specimens from the wild, C.S. provided assistance in animal husbandry during the experiment, D.A.V. analyzed the data and performed immunohistochemical staining, M.J.F. performed the in situ hybridization, provided assistance with the experimental design and analyzed the histopathology data, M.J.L.K. provided assistance with the semi-quantitative scoring and histopathological analyses, S.J.v. B. and A.G. participated in developing the concept of the project and provided assistance with the experimental design. All authors reviewed and approved the paper.

Additional Information
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