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Two Different Macaviruses, *ovine herpesvirus-2* and *caprine herpesvirus-2*, Behave Differently in Water Buffaloes than in Cattle or in Their Respective Reservoir Species

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**Abstract**

The ongoing global spread of “exotic” farm animals, such as water buffaloes, which carry their native set of viruses, may bear unknown risks for the animals, into whose ecological niches the former are introduced and vice versa. Here, we report on the occurrence of malignant catarrhal fever (MCF) on Swiss farms, where “exotic” water buffaloes were kept together with “native” animals, i.e. cattle, sheep, and goats. In the first farm with 56 water buffaloes, eight cases of MCF due to *ovine herpesvirus-2* (OvHV-2) were noted, whereas additional ten water buffaloes were subclinically infected with either OvHV-2 or *caprine herpesvirus-2* (CpHV-2). On the second farm, 13 water buffaloes were infected with CpHV-2 and two of those succumbed to MCF. In neither farm, any of the two viruses were detected in cattle, but the Macaviruses were present at high prevalence among their original host species, sheep and goats, respectively. On the third farm, sheep were kept well separated from water buffaloes and OvHV-2 was not transmitted to the buffaloes, despite of high prevalence of the virus among the sheep. Macavirus DNA was frequently detected in the nasal secretions of virus-positive animals and in one instance OvHV-2 was transmitted vertically to an unborn water buffalo calf. Thus, water buffaloes seem to be more susceptible than cattle to infection with either Macavirus; however, MCF did not develop as frequently. Therefore, water buffaloes seem to represent an interesting intermediate-type host for Macaviruses. Consequently, water buffaloes in their native, tropic environments may be vulnerable and endangered to viruses that originate from seemingly healthy, imported sheep and goats.

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**Introduction**

The water buffalo was introduced into Switzerland in 1996, as 5 farmers from a western-central region imported 14 pregnant water buffalo cows and one water buffalo bull from Romania. For many years, artificial insemination with semen imported from Italy was performed, in order to maintain a broad genetic basis among the water buffalo population [1]. Meanwhile, water buffaloes are held in 75 farms all over Switzerland [personal communication, Tierverkehrsdatenbank, 12-23-2011] and further breeding is possible without importing new animals.

The Mediterranean River Buffalo, antecedent of the Romanian and Italian buffalo breeds, originates from the Domestic Asian Water Buffalo (*Bubalus bubalis*) [2,3]. Although water buffaloes seem to be susceptible to most diseases that affect cattle, showing some variety in sensitivity or resistance, they are generally known to be in good health and well adapted to the hot and humid climates of tropical countries with the corresponding pathogens. However, reactions to some diseases may vary depending on the region, environment and genetic basis of the buffalo breed [4].

Foot-and-mouth disease, Rinderpest, Infectious Bovine Rhinotracheitis, Bluetongue, Bovine Viral Diarrhea, Buffalo pox, Rabies, Ephemeral Fever and Malignant Catarrhal fever are some of the viral infections detected in water buffaloes in Asia and Europe [5].

Malignant catarrhal fever (MCF) is an often-fatal lymphoproliferative disease of mainly ruminant species including domestic cattle, water buffalo, American bison, various species of cervids and other wild living ruminants, caused by closely related *Gammaherpesvirus* of the genus *Mavrus* [6]. Two viruses are primarily responsible for the disease; the *ovine herpesvirus-2* (OvHV-2) as seen worldwide in sheep-associated-MCF (SA-MCF), with sheep as the main reservoir host [7–9], and the *alcelaphine herpesvirus-1* (AHSV-1), endemic in wildebeest, inducing wildebeest-associated MCF (WA-MCF) in Africa and zoological gardens [8–10]. In recent years, further MCF agents have been recognized; the MCF-causing virus in white-tailed deer [11,12], cattle [13] and red brocket deer [14] with unknown reservoir host, although a potential connection with goats has been suggested [14,15]; the ibex MCF virus with outbreaks of MCF in bongo
anteelopes [16,17]; the AHV-2-like virus introducing MCF in Barbary red deer [18]; and the caprine herpesvirus-2 (CpHV-2) identified in healthy domestic goats [19–21], leading to clinical cases in different kinds of cervids [22–27] and probably in domestic cattle, banteng [13,27] as well as in water buffaloes [28].

Clinical signs of MCF in water buffaloes are characterized by depression, anorexia, high fever, lymphadenopathy, conjunctivitis and corneal opacity, inflammation, ulceration and exudation of the upper digestive- and respiratory tract, diarrhea and neurological deficiencies leading to death [29–33].

In Switzerland, SA-MCF sporadically occurs in domestic cattle [34]; one case in a farmed sika deer has been mentioned [35]. OxHV-2 has been detected in pigs [36]. In water buffaloes, only one case of probable CpHV-2-associated MCF-like disease has been described [28].

Over the last few years the popularity of housing water buffaloes for milk and meat production has risen. Up to now approximately 12002 water buffaloes are kept in Switzerland, in smaller herds and herds of up to more than 50 animals (personal communication, Tierverkehrsdatenbank, 12-23-2011), distributed over all geographical regions. The economic loss in case of death is quite high. Preceding our survey, a Swiss farm reported increased fatalities following MCF-like symptoms in water buffaloes. For the current case report we therefore decided to investigate 2 Swiss water buffalo farms with history of disease and one farm with no known MCF-like case, by means of real-time PCR analysis, in order to detect OxHV-2 and/or CpHV-2 as probable triggers of disease and gain more insights concerning epidemiological issues.

Materials and Methods

2.1. Ethics statement

This study was carried out in strict accordance with the Swiss regulations for animal experimentation. The protocol for this study was approved by the Cantonal Veterinary Office Zurich, ZH, Switzerland (Permit Number: 102/2012).

Moreover, the owners of the water buffaloes gave permission for their animals to be used in this study.

2.2. Animals

In the course of our study we investigated 3 Swiss farms housing water buffaloes (Bubalus bubalis) as well as additional ruminants. Switzerland is essentially free of bovine herpesvirus-1 (BoHV-1) and keeps this state on the basis of serological surveys. Before granting import permits, it is mandatory to test susceptible cloven-hoofed animals, including water buffaloes, for antibodies against BoHV-1. Only seronegative animals will be issued with a permit for importation. Due to serological cross-reaction, antibodies against bubaline herpesvirus-1 (BuHV-1) can be detected in the same test (Engels and Ackermann, personal observation). Moreover, for differential diagnostic reasons, several individuals with MCF (detected in our study) were tested with negative result for BoHV-1 antibodies. Additionally, a total of 31 buffaloes, including individuals from all 3 farms, were specifically tested for BoHV-1 and BuHV-1 shedding by real-time PCR (Ackermann and co-workers, unpublished data); all with negative result. Therefore, we assume that neither BoHV-1 nor BuHV-1 had any effect on the present study.

Farm 1. The first herd, located in the lowlands, consisted of 56 water buffaloes, including 53 cows and calves (not individually discriminated) and three adult bulls. During summertime the water buffaloes lived in a free-stall barn with periodical pasturing, in direct contact with 42 Holstein-Friesian cattle. Moreover, 60 Dorper sheep and 6 Boer goats were kept in a separate barn, approximately 30 m off the buffaloes’ and cattle’s barn. Some water buffaloes spent the summer on an external pasture belonging to a remote farm, where also sheep were kept [relevance for case No. 8]. During autumn and wintertime the water buffaloes and cattle were housed indoors under the same roof together with the small ruminants. As Dorper sheep do not keep to a restricted lambing period, birth giving occurs year round. Milk lambs were allowed to roam freely among buffaloes and cattle, and occasionally fed from the same manger. The sheep and goats were removed from the premises in August 2011; 3–4 months after a first water buffalo came down with Malignant Catarrhal fever (MCF).

Farms 2. The second herd of 21 water buffaloes was held in the alpine regions of Switzerland. Throughout the year, including birth-giving times, the water buffaloes were housed in the same barn together with 4 cattle and 7 goats. They shared the summer pasture with the goats and additional cattle. Remaining feed of the goats as well as goat milk was occasionally fed to the water buffaloes. No sheep were kept on the farm, and no contact with external sheep is known.

Farm 3. The third herd of 43 water buffaloes was situated in the central region of Switzerland. Apart from the water buffaloes living in a free-stall barn with periodic access to the pasture, fluctuating numbers of 20 to 50 sheep of a Swiss breed (Schweizer Alpenschaal) were housed in a separate stable and separate pasture throughout the year. The sheep gave birth from January to March; no free roaming of offspring occurred on the premises. There was no known direct contact between sheep and water buffaloes. Neither cattle nor goats were kept on the same farm.

2.3. Sample collection

5–10 ml EDTA-treated blood was collected from 118 water buffaloes and 13 goats and from a selection of 17 cattle and 12 sheep. Additionally to the 160 blood samples nasal swabs were taken from 30 water buffaloes and six goats. From one water buffalo and 3 water buffalo fetuses only fresh organ samples could be collected. Two water buffalo calves from farm 1 were monitored beginning by time of birth by analyzing a monthly blood sample over a period of 8 months. The blood of 28 water buffaloes from farm 1 was re-tested 8 to 13 months after the first blood sampling.

2.4. Sample preparation and DNA extraction

Buffy-coat cells from each sample were gained through the addition of 40 ml lysis buffer (0.13 M NH₄Cl, 10 m M CH₃CO₂, 0.1 m M EDTA [pH 7.2]) to the EDTA-treated blood, followed by centrifugation for 10 min at 868 × g and disposal of the supernatant. This step was repeated 1–3 times. The final pellet was re-suspended in 40 ml phosphate-buffered saline, and after a another centrifugation step and disposal of the supernatant, was stored at −20°C until further use. Dry nasal swabs were used directly for DNA extraction. Fresh organ samples were crushed before lysis and DNA extraction. DNA extraction was carried out using the QiAamp DNA Mini Kit (Qagen, Hombrechtkon, Switzerland) according to the manufacturer’s recommendations for each of the collected sample materials. The extracted DNA was used directly for PCR typing or stored at −20°C.

2.5. PCR

OxHV-2. The genomic DNA sequence of the ORF 63 tumour protein from OxHV-2 provided the basis for the design of the real-time PCR. The sequence described by Taus et al. [37] (GenBank accession no. DQ159083.1), as well as the sequence of Hart et al. [38] (GenBank accession no. AY839756.1), were
considered. Primers and probe were designed using the Perkin-
Elmer Primer Express software (version 1.0, Perkin-Elmer, Foster
City, California). The following primers and probe were selected:
forward primer 5′-GAG AAC AAG CGC TCC CTA CTG A-3′
[Life Technologies Europe BV, Zug, Switzerland], reverse primer
5′-CGT CAA GCA TCT TCA TCT CGA G-3′ [Life Technolo-
gies], probe 5′-FAM-AGT GAC TCA GAC GAT ACA CGA
CGC GAC A-TAMRA-3′ [Micsysn, Balgach, Switzerland].
Possible cross-reactions with other Hepatoviridae were evaluated
with the National Center for Biotechnology Information Basic Local
Alignment Search Tool (NCBI BLAST, http://blast.ncbi.nlm.nih.
gov/Blast.cgi). PCR reaction mixture and cycling conditions
were carried out as described previously by Hussy et al. [39] consider-
ing the above-mentioned modifications of primers and probe and run
on an 7900HT Fast Real-Time PCR System [Life Technologies]
with 9600 emulsion ramping. 10 nl of diluted and undiluted
samples were tested in duplicates.

**CpHV-2.** For the detection of CpHV-2 primers and probes
described by Cunha et al. [40] were used. Real-time PCR
was performed using the same conditions as for the OvHV-2 PCR.
However only 1 nl of diluted or undiluted template DNA was
added in duplicates and the ramping of the cycler was set to
the standard rate.

**12S rDNA.** The 12S rDNA gene was used as an internal
control in order to confirm positive DNA extraction. The PCR
was based on the finding of a consensus sequence between
previously published [41] 12S rDNA sequences of various bovid
taxa. 5′-GGG GTG CTT TAT AYC CTA CTA GAG-3′,
600 nM, served as forward primer, 5′-TCA GCA AGT ATT
GGT GAT TAT C-3′, 600 nM, [Micsysn, Balgach,
Switzerland] as the reverse primer. The probe 5′-VIC-AGC CTG
TTC TAT AAY CGA T-MGBNFQ-3′, 160 nM, [Life Technolo-
gies] was used. Samples from water buffaloes, cattle, sheep
and goats were positively tested for the described house-keeping gene,
confirming their sensitivity to the reaction. For real-time PCR the
same conditions as mentioned above were used. 10 nl of the
diluted sample in question applied in dual wells served as template.

**Results**

3.1. Incidence of clinical MCF
A total of 4 severe and 4 relatively mild cases of MCF were
noticed over a period of 17 months, from April 2011 to August
2012, among the water buffaloes of farm 1 (Table 1). In each single
case OvHV-2 DNA was detected in the diseased animals. In farm
2, two cases of clinical MCF were reported, one in October 2008
and one in November 2010 but testing for OvHV-2 DNA
remained negative. No cases of clinical MCF were reported from
farm 3.

Thus, despite the presence of small ruminants on all three
farms, three different types of outcomes were noticed: (1) relatively
frequent (almost monthly) cases of OvHV-2-associated MCF in
water buffaloes of farm 1; (2) relatively low frequency of clinical
MCF in water buffaloes without detection of OvHV-2 DNA on
farm 2; (3) no evidence of clinical MCF on farm 3. Based on these
observations, questions arose concerning the prevalence of OvHV-
2 and other potential MCF agents among the small ruminants
living on the same farms as well as about the possibility of subclini-
cal circulation of MCF agents among the water buffaloes.

3.2. Search for OvHV-2 and CpHV-2 among cattle and
small ruminants on the same farms

13 of 42 cattle on farm 1 were available for testing and all
turned out to be negative for both viruses, OvHV-2 and CpHV-2.
Since OvHV-2 is known to be highly prevalent in sheep, only 6
out of 60 sheep on farm 1 were tested. Indeed, 5 of those 6 sheep
carried OvHV-2, suggesting a high prevalence of this MCF agent
among the sheep of farm 1. In contrast, CpHV-2 DNA was not
detected in the samples of the same sheep, suggesting no or a low
prevalence of CpHV-2 among the sheep. Among the 6 goats on
the same farm, one was identified as positive for CpHV-2 and
another as positive for OvHV-2 DNA in blood and nasal swabs.
No viral DNA was detected in a nasal swab sample from a third
goose, which however tested positive for both OvHV-2 and CpHV-
2 in the blood. Thus, at least two different MCF agents were
present and apparently circulating in their primary hosts on farm
1.

In farm 2, the 4 cattle tested negatively for both OvHV-2 and
CpHV-2, whereas CpHV-2 DNA was detected in the blood
samples from 6 of the 7 co-housed goats. This was consistent with
a high prevalence of CpHV-2 in its native host species. In contrast,
OvHV-2 was not detected in the same goats.

In farm 3, again 6 individual sheep were randomly selected for
testing against OvHV-2 and CpHV-2. All of them were shown to
be infected with OvHV-2, but none of them provided evidence of
CpHV-2 infection.

Thus, OvHV-2 circulated among sheep in farm 1 and farm 3,
whereas CpHV-2 showed a high prevalence among the goats in
farm 2 and a lesser prevalence among the goats in farm 1.

3.3. Prevalence of OvHV-2 and CpHV-2 among water
buffaloes on the same farms

Fig. 1 shows that both MCF agents present on farm 1, OvHV-2
and CpHV-2, had taken an opportunity to infect water buffaloes.
However, CpHV-2 caused only one subclinical infection among
the 56 water buffaloes, whereas OvHV-2 was associated with 8
cases of clinical MCF and 9 incidences of subclinical infections on
the same farm. Thus, OvHV-2 infections among water buffaloes
in this farm amounted to a prevalence of about 30%, but only
50% of the infected individuals succumbed to MCF. These
observations indicate that, with equal frequency, OvHV-2
infections of water buffaloes may either take a subclinical or a
manifest course with respect to MCF. Interestingly, one of the
eight water buffaloes with clinical MCF (case No. 8, Table 1)
tested OvHV-2- negative in December 2011. However, five
months later and at a total of 9 months after the OvHV-2-positive
sheep had been removed from the farm, the same animal fell ill
with OvHV-2-associated MCF. In this case, the source of OvHV-
2 remains obscure.

Analysis of nasal swab samples from farm 1 revealed that in 3
out of 3 water buffaloes with clinical MCF and in 5 of 10
subclinically OvHV-2-infected water buffaloes the agent could
simultaneously be detected in white blood cells and in nasal
secretions. In contrast, excretion of CpHV-2 could not be detected
in the nasal sample of the one CpHV-2-positive water buffalo.

Organs of 2 fetuses from seriously diseased, OvHV-2-positive
water buffaloes of farm 1 were also analyzed; one fetus was proven
to be positive for OvHV-2 in the spleen. Thus, vertical
transmission of OvHV-2 in water buffaloes seemed to be possible.
In an organ pool of fresh brain, kidney, heart and intestine of the
second fetus, no virus was detected. Unfortunately no spleen
sample was available from this second fetus. The only CpHV-2-
positive water buffalo of farm 1 had an abortion without showing
further clinical symptoms. Fresh fetal brain and lymph node
samples were tested negative for both viruses; no obvious reason of
prenatal death could be found. Thus, vertical transmission of
CpHV-2 among water buffaloes could not be substantiated.
Table 1. Time scale of events and OvHV-2 real-time PCR results from water buffaloes with clinical MCF of farm 1.

| Buffalo No. | Clinical MCF | Apr-11 | May-11 | Jul-11 | Aug-11* | Sep-11 | Oct-11 | Nov-11 | Dec-11 | Jan-12 | May-12 |
|-------------|--------------|--------|--------|--------|---------|--------|--------|--------|--------|--------|--------|
| severe      | OvHV-2+<sup>3</sup> |        |        |        |         |        |        |        |        |        |        |
| 1           |              |        |        |        | OvHV-2+<sup>3</sup> |        |        |        |        |        |        |
| 2           |              | nd     | OvHV-2+ | nd     | OvHV-2+<sup>3</sup> |        |        |        |        |        |        |
| 3           |              | nd     | nd     | OvHV-2+ | nd     | OvHV-2+<sup>3</sup> |        |        |        |        |        |
| 4           |              | nd     | nd     | nd     | nd     | OvHV-2+<sup>3</sup> |        |        |        |        |        |
| mild        |              |        |        |        |         |        |        |        |        |        |        |
| 5           |              | nd     | nd     | OvHV-2+ | nd     | nd     | OvHV-2+<sup>3</sup> |        |        |        |        |
| 6           |              | nd     | OvHV-2+ | nd     | nd     | OvHV-2+<sup>3</sup> |        |        |        |        |        |
| 7           |              | nd     | nd     | OvHV-2+ | nd     | nd     | OvHV-2+<sup>3</sup> |        |        |        |        |
| 8           |              | nd     | nd     | nd     | nd     | OvHV-2+<sup>3</sup> |        |        |        |        |        |

OvHV-2+ = positive for OvHV-2;
OvHV-2- = negative for OvHV-2;
nd = not determined;
<sup>3</sup> = clinical signs of MCF;
<sup>4</sup> = euthanasia or slaughter due to clinical MCF;
<sup>5</sup> = sheep removed from farm.
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Fig. 2 shows that water buffaloes may be very susceptible to CpHV-2. With 13 positive individuals among 20 animals tested on farm 2, the prevalence amounted to 65%. However, only two out of those 13 animals did succumb to an MCF-like disease. In the absence of OvHV-2, these observations strongly suggest that CpHV-2 was the most likely agent for the disease. However, the subclinical course with CpHV-2 was much more frequent than with OvHV-2 in farm 1. Interestingly, CpHV-2 DNA was only detected in the nasal swab from one out of 11 CpHV-2-positive animals. Therefore, independent circulation of CpHV-2 among water buffaloes was not strongly supported by our data.

Remarkably, despite of its apparently high prevalence among the sheep in farm 3, transmission of OvHV-2 from the reservoir host to water buffaloes was not detected and clinical cases did not occur. Thus, strict separation of sheep and water buffaloes on such farms may be effective to prevent MCF.

Discussion

Considering the ongoing global spread of “exotic” farm animals from their native to novel environments, it is imperative to keep in mind that they all carry their own sets of viruses, which may be well adapted to their original hosts, but may bear unknown risks for the animals, into whose ecological niches they are introduced. In contrast, “exotic” farm animals may be especially sensitive for viruses circulating in foreign countries and various MCF-causing agents seem to be an interesting topic in this context.

For the present communication, we started off with clinical cases of malignant catarrhal fever (MCF) in water buffaloes (considered “exotic”) from two Swiss farms and, consecutively, analyzed the genoprevalence of two different MCF agents, OvHV-2 and CpHV-2, in various susceptible ruminants on the same farms. A third farm, from which no cases of MCF had been reported, served as comparison. The following insights were gained:

![Figure 1. Proportion of Macavirus-affected water buffaloes on farm 1.](https://example.com/figure1.png)

Samples from water buffaloes (n = 56) were analyzed by real-time PCR for the detection of OvHV-2 DNA and CpHV-2 DNA, respectively. The figure plots the percentage of healthy animals (“No virus detected”: white; “infected and healthy”: grey) versus the proportion of animals with MCF due to OvHV2 infection (red). In the secondary pie, the proportion of infected but healthy animals is further subdivided into animals with OvHV2 (green) and CpHV2 (blue), respectively.
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Our report confirms earlier work mentioning that water buffaloes were more susceptible to OvHV-2 infection than cattle [8,31,33,42]. As expected, sheep and goats, the reservoir hosts of OvHV-2 and CpHV-2, respectively, were shown to be Macavirus-carriers in both, farm 1 and farm 2. Yet, despite similar exposure, cases of MCF as well as subclinical genoprevalence of the MCF agents were noted only in the water buffaloes but not in cattle. It is well known that susceptibility to infection by Macaviruses may depend on animal species. Certain species of deer (i.e., Père David’s deer [43]), Bali cattle [42,44] as well as American bison [45-49] are considered highly susceptible to OvHV-2.

(2) Despite this increased susceptibility, the mere presence of Macavirus-carriers on the farms did not appear to be sufficient for the transmission of the two MCF agents from the reservoir hosts to the water buffaloes as indicator hosts. For example, transmission of CpHV-2 from the infected goats to water buffaloes was only observed in one case on farm 1, whereas the genoprevalence of CpHV-2 amounted to 65% among the water buffaloes on farm 2. The true reason for this difference could not be determined under the present conditions. However, possible explanations for the high CpHV-2 transmission rate in farm 2 may be the joint keeping of goats and water buffaloes on pastures and in the stable throughout the entire year. Moreover, the water buffaloes were occasionally fed with remaining feed and milk of the goats. Alternatively, one may consider independent circulation of CpHV-2 among water buffaloes. However, the virus was detected only in one out of 11 nasal swabs taken from CpHV-2-positive water buffaloes of farm 2, which did not heavily support this possibility.

OvHV-2 was highly prevalent in sheep on farm 1 as well as on farm 3. Whereas, on farm 3, the virus was not transmitted to the water buffaloes, it caused subclinical infection and mild to severe MCF outbreaks among the water buffaloes on farm 1. The obvious explanation for this may be attributed to the different types of management on the two farms: whereas the water buffaloes and the sheep had been co-stabled at winter time in farm 1, they were housed and grazed separately in farm 3, which may be considered a successful preventive measure. Although long distance spread of OvHV-2 from sheep to susceptible animals, such as bison, has been reported by others [49], close contact among sheep and other susceptible species, especially keeping them under the same roof, is considered as a more efficient means to transmit OvHV-2 [30,51]. Also, the temporary free ranging of hand-raised lambs in the water buffalo stable of farm 1 seems to favor OvHV-2 transmission. The role of the OvHV-2 infected goats of farm 1 is not clear. OvHV-2 DNA in blood samples and nasal swabs of goats housed together with sheep has been detected previously [15,19,21,27,44,52]. In cattle and bison without clinical signs of MCF, DNA of OvHV-2 has been extracted from blood, milk, nasal secretions and conjunctiva [13,46-48,50,51,53,54].

It is unclear whether the OvHV-2 and CpHV-2-positive results from healthy water buffaloes in farms 1 and 2 were merely due to ongoing prolonged incubation periods prior to an MCF outbreak or else were attributable to water buffaloes as potential reservoir hosts.

Although the detection of OvHV-2 DNA in nasal secretions does not provide direct proof for virus excretion, the relatively high frequency of detection of OvHV-2 DNA in nasal swabs of infected water buffaloes on farm 1, might support the notion that OvHV-2 may circulate by horizontal transmission among water buffaloes in a sheep-independent manner. However, re-testing of blood samples of originally virus-free water buffaloes in farm 1, at eight to 13 months after removal of the sheep from the premises, did not reveal any newly infected carriers of OvHV-2. Similarly, two water buffalo calves, born after the removal of the sheep, were monitored at monthly intervals for OvHV-2. Neither calf turned OvHV-2 positive until the monitoring was ceased seven months later. The source of infection for water buffalo No. 8 from farm 1 remains unclear. Since the water buffalo spent the summer months of 2011 on an external farm, which also housed sheep, an infection on those premises must be considered. However, the summering farm housed various water buffaloes and to
our knowledge no further case of MCF had been registered. Indirect transmission, as well as direct horizontal transmission from a subclinically infected water buffalo as reservoir host, cannot be ruled out. A further possible explanation is that the virus load of the sample taken in December 2011 was under the detection limit of realtime PCR analysis; temporary variation of OvHV-2 DNA detection in peripheral blood lymphocytes of susceptible hosts has previously been demonstrated in cattle [51,53–55] and bison [56,57].

Interestingly, OvHV-2 DNA was detected in the spleen of a fetus from a deceased, OvHV-2-positive dam of farm 1, which confirms potential trans-placental transmission of OvHV-2 among water buffaloes. To our knowledge, this might represent the first report on vertical Macavirus transfer among MCF indicator hosts. Yet, as no spleen samples were available in the cases of the other two examined fetuses of a further OvHV-2 and of a CphHV-2-positive dam, no definitive conclusion can be drawn concerning the frequency of such a transmission. Vertical transmission seems to play an important role in wildebeest, the reservoir host of AIHV-1 [10,58,59]. In contrast, in sheep, vertical transmission of OvHV-2 may occur only rarely [60], whereas trans-placental transmission from CphHV-2 positive goats to their kids seems not to play a major role [21]. It will be important to address these issues more deeply, especially with regard to water buffaloes.

(3) Despite the considerations mentioned above, the water buffaloes seemed to be generally more susceptible to CphHV-2 (two-thirds of exposed animals positive) than to OvHV-2 (one-third positive). However, with 50% of infected water buffaloes of farm 1 succumbing to MCF due to OvHV-2, compared to only 15% of the CphHV-2-positive animals developing MCF on farm 2, the penetration rate of MCF appeared to be higher with OvHV-2 than with CphHV-2. Compared to cattle, the best studied indicator host, water buffaloes seem to be more susceptible to Macavirus infections but less prone to succumb to MCF. However, in comparison to typical reservoir hosts, water buffaloes seem to be less susceptible to Macavirus infections but more prone to subsequently succumb to MCF. Thus, they may represent an interesting intermediate-type host for the Macaviruses.

Therefore, it may be important to extend the research on MCF not only to relatively poorly characterized viruses, such as CphHV-2, but also to thus far neglected indicator host species, such as water buffaloes. Indeed, MCF is of great concern among different species of buffaloes, particularly in Asia, but also in Brazil and Italy [29,30,42]; clinical MCF in free-ranging African buffaloes is considered an emerging disease [61]. In association with shared housing of water buffaloes and sheep and the frequency of MCF, OvHV-2 seems to be of high relevance [29–31]. However, our investigation confirms that water buffaloes can also succumb to MCF due to CphHV-2 [28]. Hence, shared housing of water buffaloes and goats has to be considered as a risk factor for MCF as well.

In the present case report, the question whether or not Macaviruses actively circulate among water buffaloes could not be unambiguously solved. The notion that Macavirus DNAs were detected in nasal secretions of virus-positive animals and in the spleen of one fetus seems to argue in favor of active virus transmission among water buffaloes. However, after the removal of the original reservoir hosts from one farm, newly infected individuals, with just one exception (case No. 8, Table 1), could not be identified, which rather speaks against efficient intra-species transmission.

In conclusion, two different Macaviruses, OvHV-2 and CphHV-2, behaved differently in the “exotic” host, the water buffalo, than in “native” host species, such as cattle, sheep, and goats. On the one hand, the water buffaloes seemed to be more susceptible than cattle to infection with either virus; on the other hand, the case fatality rate among water buffaloes was lower than the perceived lethality among cattle [34]. However, recent reports from countries with a high concentration of farms, where OvHV-2 reservoir species are kept together with MCF-indicator species, suggested an increasing incidence of non-lethal outcomes of MCF, which also involved either subclinical cases among cattle, and more chronic, cutaneous forms of the disease [50,62–64]. These observations may signify that either OvHV-2 has a capacity to adapt to new hosts upon serial passaging or else that the cattle in those countries are selected to become resistant to MCF. Both possibilities would be worthwhile to consider further.

Finally, it may be valuable to also look into the water buffalo’s immune responses against the different types of Macaviruses in order to better understand the reasons for the relatively high number of subclinical cases.

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Author Contributions

Conceived and designed the experiments: MA ABJS ME MF. Performed the experiments: ABJS RB MF. Analyzed the data: ABJS MA ME. Contributed reagents/materials/analysis tools: RB MF. Wrote the paper: MA ABJS ME. Revised the paper: MA ABJS ME RB MF.

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