Epizootic Spread of Schmallenberg Virus among Wild Cervids, Belgium, Fall 2011

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Schmallenberg virus was detected in cattle and sheep in northwestern Europe in 2011. To determine whether wild ruminants are also susceptible, we measured antibody seroprevalence in cervids (roe deer and red deer) in Belgium in 2010 and 2011. Findings indicated rapid spread among these deer since virus emergence ≈250 km away.

During summer and fall of 2011, a nonspecific febrile syndrome among adult dairy cows in northwestern Europe was reported. During November 2011, an enzootic outbreak causing fetal death or neurologic signs in newborn lambs, kids, and calves emerged throughout several countries in Europe. Both syndromes were associated with the genome of a new Shamonda/Sathuperi-like orbivirus named Schmallenberg virus (SBV) in the blood (adults) or central nervous system (newborns) (1,2). Susceptibility of wild ruminants can be expected on the basis of the behavior of related viruses of the Simbu serogroup. Therefore, we measured seroprevalence of antibodies against SBV in wild red deer (Cervus elaphus) and roe deer (Capreolus capreolus) and looked for the viral genome in fetuses from pregnant deer found dead.

The Study

Blood samples were collected during postmortem examination of 313 red deer and 211 roe deer shot during the 2010 and 2011 hunting seasons. The 524 samples were randomly collected during October–December from 35 hunting estates in 4 of the 5 provinces in southern Belgium (Figure 1). The animals’ sex; age; body condition; and macroscopic aspects of hooves, mucosae, and internal organs were recorded. IgG against the recombinant nucleoprotein of the emerging SBV was detected by using an ELISA kit (ID Screen Schmallenberg Virus Indirect, version 1; ID.vet Innovative Diagnostics, Montpellier, France). Results are expressed as percentages of the reference signal yielded by the positive control serum; serologic status is defined as negative (<60%), doubtful (60%–70%), or positive (>70%). Neutralizing antibodies against SBV were sought as described (3) in subsets of roe deer serum (IgG-negative and IgG-positive according to ELISA), and a linear relationship between percentages and reciprocal neutralizing titers was found. In addition, necropsies were performed on 22 fetuses and 5 newborn red deer fawns; brain samples were tested for SBV genomic RNA and cellular β-actin transcripts by reverse transcription quantitative PCR (3). Contingency tables were analyzed by using χ² analysis to detect associations between seroconversion and species, sex, age, sampling location, and sampling date. Significance level was p<0.05.

No gross lesions compatible with any disease were found in any deer. All 299 serum samples collected during the fall of 2010 were negative for IgG against SBV. However, among the 225 samples from deer killed in 2011, seroprevalence was 43.1% (95% CI 36.6%–49.6%). No significant association was found between species and seroconversion: 40.5% (95% CI 31.6%–49.5%) among red deer and 45.9% (95% CI 36.5%–55.2%) among roe deer (p = 0.42). Acquired immunity against SBV was thus already high, suggesting that SBV had quickly spread since its emergence ≈250 km northeast during late summer 2011.

A significant association between month of sampling and seroconversion was detected for both deer species

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Almost impossible to collect in the wild (quickly eaten by scavengers), and the absence of SBV-specific genetic material or morphologic alterations at necropsy are not evidence of noninfection. Therefore, no objective facts confirm or refute transplacental transfer.

Because the virus can infect the fetus only after the first placentome has developed and because roe deer embryos remain in diapause until January (7), it is unlikely that SBV has contaminated many roe deer fetuses. Because 90% of roe deer were already SBV positive in mid-December and because circulating antibodies prevent transplacental passage of the closest phylogenetic relatives of the virus (8), we suggest that roe deer fetuses were probably not infected. On the contrary, red deer mate in September, and the first functional placentome is established by the end of October (9); thus, 80% of pregnant red deer were exposed to the emerging virus when placental transfer was possible. Furthermore, 35% of pregnant red deer were infected in November and December, i.e., after establishment of the first placentome and before the fetus was immunocompetent. By extrapolating the rate of transplacental infection among cattle (6), we determined that 28% of these pregnancies resulted in contamination of the fetus, i.e., 10%, of expected pregnancies. Because unrestricted replication of Simbu-like viruses occurs in the central nervous system of immunologically incompetent ruminant fetuses (1), which can lead to a typical arthrogryposis/hydranencephaly syndrome, a 10% loss among fawns can be expected in 2012.

In the same geographic area, 5 years apart, 2 arboviruses have emerged: bluetongue virus serotype 8 (BTV-8) during the summer of 2006 and SBV during the summer of 2011. For each virus, Culicoides spp. midges function
as vectors and infect sheep, goats, cattle, and red deer. Although most (>50%) red deer seroconverted against BTV-8, only a few (<3%) roe deer sampled in the same places and at the same time were BTV-8-positive (10), which sharply contrasts with the SBV seroconversion rates reported here. This finding invalidates the assumption that less exposure of roe deer to infected midge bites explains the almost complete absence of seroconversion against BTV-8 in this species. The emergence of SBV thus reveals the existence of roe deer–specific anti–BTV-8 host factors, posing a fascinating question.

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