Prevalence and genetic diversity of coronaviruses in wild birds, Finland

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ABSTRACT

Introduction: Migratory birds act as hosts for a number of zoonotic viruses, and have the ability to disperse these viruses to distant geographic locations. Coronaviruses (CoVs) represent a family of zoonotic viruses with wide variety of animal hosts, including birds and humans. The infections caused by coronaviruses vary from mild to severe, depending on the viral species and the host. Since the coronaviruses exhibit extraordinary large RNA genome, also the rate of homologous recombination is high, which in turn contributes to the genetic diversity and interspecies host-switches of CoVs. The emergence of novel CoVs has been rich during the last decades, and wild birds seem to serve as reservoirs for a variety of CoV strains. We examined the CoVs circulating among wild birds in Finland.

Materials and methods: Samples (cloacal swab, tracheal swab, oropharyngeal swab, or tissue) representing 61 bird species were collected during 2010-2013, and examined by RT-PCR targeting the RdRp gene for the presence of CoV RNA.

Results: Altogether 51/939 (5.4%) of the examined birds were found positive by RT-PCR. Diverse gamma- and deltacoronavirus sequences were detected.

Discussion: Gamma- and deltacoronaviruses circulate among wild birds in Finland. The number of CoV-positive birds detected each year varies greatly.

Introduction

Coronaviruses (CoVs) comprise a family under the order Nidovirales (family Coronavirusae) and infect a wide variety of mammals and birds. The course of infection varies greatly from asymptomatic to severe disease, depending on the host and virus species in question. The genome of CoVs is one of the largest (25–32 kb) viral RNA-genomes [1]. Based on phylogenetic analysis, the CoVs are divided into four different genera: Alpha-, Beta-, Gamma-, and Deltacoronavirus. The alpha- and betacoronaviruses are carried by mammals, whereas the gamma- and deltacoronaviruses mainly infect birds, with few exceptions [2,3]. The large genomes, infidelity of the RNA-dependent RNA polymerase, and high frequency of homologous RNA recombination are the main factors contributing to the high genetic diversity of CoVs [4–6]. The first CoV, Infectious bronchitis virus (IBV), was identified in 1937 [7]. IBV mainly infects chickens, but may infect other bird species as well. IBV is highly contagious and affects the respiratory tract, gut, kidney, and reproductive systems, causing substantial economic losses in the poultry industry [8]. Despite the global distribution of IBV, poultry in Finland remained free of clinical cases until April 2011 [9] after which outbreaks involving several CoV genotypes have occurred in Southern Finland.

The first human CoVs were identified in 1960s [10–12]. The human CoVs cause generally mild to moderate upper respiratory tract infections [13–15]. In 2003, a novel highly pathogenic betacoronavirus emerged in China, causing severe disease characterized by acute respiratory distress and it became known as severe acute respiratory syndrome (SARS)-CoV. The emergence of SARS-CoV [16] inspired virologists to more explore the highly divergent group of coronaviruses and their hosts, leading to the identification of a rapidly growing number of CoV species particularly in bats [2]. More recently, another highly pathogenic betacoronavirus infecting humans, the Middle East Respiratory Syndrome (MERS) CoV emerged in 2012 with a case fatality rate of over 40% [17,18].

Migratory birds have the ability to facilitate the dispersion of microorganisms with zoonotic potential. Wild birds have been associated with the ecology and dispersal of at least West Nile virus, tick-borne encephalitis virus, influenza A virus (IAV) and Newcastle disease virus (NDV) [19–21]. Since the discovery of IBV in 1937, it remained the only
known Gammacoronavirus for over 50 years, but the number has increased dramatically during the last 10 years [5]. Thereafter, representatives of the genera Gamma- and Deltacoronavirus have been isolated from both wild and domestic birds including species from the order Anseriformes, Pelecaniformes, Ciconiiformes, Galliformes, Columbiformes, and Charadriiformes [22-24]. In this report we provide a description of CoV species circulating in wild birds in Finland. Altogether 939 samples representing 61 different bird species were collected during 2010–2013 and examined for the presence of CoV RNA.

Materials and methods

Ethical statement

All active and passive surveillance samples tested in this study, including samples provided by hunters, and injured or diseased birds were initially sent to the Finnish Food Safety Authority Evira (Helsinki, Finland) for avian influenza testing as part of a national surveillance program. All hunters had appropriate permits, and no birds were killed for research purposes. The hunted birds were shot during the annual duck-hunting season, from non-endangered species, and thereby do not require ethical approval.

Sampling

The samples were collected as part of a national active and passive surveillance program, coordinated by the Finnish Food Safety Authority Evira and University of Helsinki. The sample panels included (1) hunted clinically healthy birds (active surveillance), (2) hunted clinically diseased birds (active surveillance), and (3) birds found dead (passive surveillance) or (4) clinically diseased (passive surveillance) (Table S1). The sample material used for RNA detection was cloacal swab, tracheal swab, oropharyngeal swab, or tissue. The swab samples were collected using nylon swabs and stored and transported in Universal Transport Medium (both by Copan International).

Prior to RNA isolation from the swab specimen (stored at −80°C), the samples were centrifuged to remove solid particles and supplemented with additional antibiotics (streptomycin-penicillin). The tissue samples were homogenized and separated by centrifugation and supplied with antibiotics (streptomycin-penicillin). The sample panel from 2010 consisted of 343 samples, the 2011 panel of 171 samples, the 2012 panel of 287 samples, and the 2013 panel of 138 samples. The samples were also screened for influenza A virus RNA [25].

Detection of coronavirus RNA

QIAamp Viral RNA Mini Kit was used to extract the RNA from the samples. The CoV screening was performed using degenerate primers [26] to amplify a 179 nt region in the polymerase (RdRp) gene (Orf1b). All positive samples were further characterized by amplifying a longer (608–610 bp) region in the RdRp gene using previously described primers (forward primer 5’TGGGWTGGGYTAYCCWAARTGYGA-3’ and reverse primer 5’GCATWGTRTGYTGNGARCARAATTCT-3’) [23]. All PCR reactions were performed on a BioRad PTC-100 Thermal cycler or Applied Biosystems Veriti Thermal cycler by a one-step RT-PCR protocol using QIAGEN One-Step RT-PCR Kit (Qiagen, Germany). The reaction volume for PCR was 25 µl containing 5 µl 5× buffer, 1 µl dNTP mix, 1 µM both primers, 0.2 µl RNAse inhibitor (Applied Biosystems, 20 U/µl), 2 µl RNA, and 1 µl Qiagen One-Step RT-PCR Enzyme Mix with thermal conditions as follows: reverse transcription at 50°C for 30 min, initial PCR activation at 95°C for 15 min, followed by denaturation step at for 30 s, annealing at 50°C for 30 s, and extension at 72°C for 30 s. The PCR products were analyzed on 2% agarose gels and gel purified using QIAquick gel extraction kit (Qiagen).

Sequencing and phylogenetic analysis

The RT-PCR amplicons were sequenced in both directions at the Sequencing Unit of Institute for Molecular Medicine Finland, (Helsinki), using BigDye v3.1 chemistry and run by ABI3730xl DNA Analyzer. The sequences were assembled in BioEdit and aligned using ClustalW [27-29]. The reference sequences were obtained through BLAST search [30], and from GenBank [31]. Phylogenetic analyses were based on sequence alignments of 464 nucleotides (position 100–547 of the RdRp gene, by IBV (AY392086) count) within Orf1ab. Sequence data from 46 of our samples was included in the analysis and the rest were left out because of inadequate sequence quality. The phylogenetic tree was constructed from Maximum-likelihood phylogenies and bootstrap values calculated by 1000 replicates. Gaps were handled by partial deletion. The evolutionary model GTR + G + I was used after evaluation of the best-fit model according to Bayesian Information Criterion. The alignments, selection of evolutionary model and phylogenetic analyses were performed within Mega5 [29].

Results

Coronavirus RNA in wild birds

We screened in total 939 samples from 61 bird species (Table S1) during a time period of four years
(2010–2013) using a traditional, conserved RT-PCR targeting a 179 fragment of the RdRp gene (Orf1ab) of all coronavirus lineages. Altogether 51/939 (5.4%) of the examined birds were found to be CoV RNA positive (Tables 1 and 2), of which 27 were found healthy, and 24 dead or diseased. CoV RNA was detected in eight species (Anas platyrhynchos, Anas crecca, Clangula hyemalis, Cygnus cygnus, Larus argentatus, Chroicocephalus ridibundus, Larus fuscus, and Columba sp.). CoV RNA was most abundant in the samples from 2010 (11%) and 2013 (7.2%), whereas during 2011 (0%) and 2012 (0.7%) only few of the tested samples were found positive. Influenza A virus (IAV) was also detected in four of the CoV positive samples (Table 1) [25,32]. The sampling locations are illustrated in Figure 1.

Phylogenetic analyses of the detected avian coronaviruses

The samples positive in the initial RT-PCR screen were consequently amplified for phylogenetic purposes by the method described by [23]. The number of reference strains for the alignments was unfortunately limited as GenBank deposited sequences vary in both location and length as a result of the wide array of approaches that have been employed for sequencing of coronaviruses. Table 2 shows the GenBank accession numbers of the viruses illustrated in the phylogenetic tree (Figure 2).

The viruses detected from wild ducks shared up to 99% nucleotide identity with other known CoVs of the Gammacoronavirus genus. The phylogenetically most closely related viruses are isolates from ducks in Russia and USA, China [33], Hong Kong [24] and South Korea [33,34]. The gammacoronaviruses that were detected in gulls in this study shared up to 99% sequence identity and clustered with previous isolates from gulls in Siberia and Alaska [23]. The viruses isolated from pigeons (Columba sp) (Fin6709 and Fin11782) and an isolate from a long-tailed duck (Clangula hyemalis) shared only up to 89% nucleotide identity with the currently published strains, with the highest nucleotide identity with duck gammacoronaviruses. Two isolates, AvCoV/Larus fuscus/Finland/10059/2013 and AvCoV/Chroicocephalus ridibundus/Finland/10083/2013, share 83% and 85% nucleotide identity with published avian and mammalian deltacoronaviruses, respectively. They clustered in the Deltacoronavirus clade in the ML trees, most closely to the White-eye CoV isolated in Hong Kong (JQ065044) (Figure 2). None of the viruses from wild birds in this study showed close relatedness to circulating IBV-strains.

Discussion

Several viruses, including zoonotic and economically significant pathogens, are known to circulate among wild birds. CoVs are a large group of viruses infecting mammals and birds, including infectious bronchitis virus (IBV), which is a highly contagious coronavirus infecting chickens. A variety of coronaviruses are known to circulate among wild birds. We set up this study to determine the occurrence of coronavirus infections in wild birds. Altogether 939 samples collected between 2010 and 2013 from wild birds were included in the study. The prevalence was fairly high (up to 11%) although with great annual variation. The sample panels differed to some extent in terms of time and geographic distribution (Figure 2) and sampled species, which might partly explain the annual variation. For example, majority of the samples of 2010 were from duck species (Anas sp.) sampled for active surveillance purposes. In 2011–2013 on the contrary, most samples were derived through passive surveillance and included a smaller proportion of ducks. However, no firm conclusions can be drawn whether the observed difference in prevalence is due to some regular pattern, or due to the sampling methods (heterogeneity of the sample material).

Some of the CoV positive samples were also positive for H3N8 and H9N2 IAV RNA. Similar double

| Species               | Clinical status (no.)                                                                 | % (no. positive/negative) | Gamma | Delta | Co-infections (no.) |
|-----------------------|---------------------------------------------------------------------------------------|--------------------------|-------|-------|---------------------|
| Anas platyrhynchos    | Hunted clinically healthy (27) and hunted clinically diseased (1)                     | 21.7% (28/129)           | x     |       | H9N2 (1) + H3N8 (2) |
| Anas crecca           | Hunted clinically healthy                                                              | 16.4% (9/55)             | x     |       | H3N8 (1)            |
| Larus argentatus      | Found dead                                                                             | 9.6% (5/52)              | x     |       |                     |
| Chroicocephalus       | Found dead                                                                             | 4.2% (1/24)              | x     |       |                     |
| ridibundus            |                                                                                       |                          |       |       |                     |
| Larus fuscus          | Found dead                                                                             | 8.3% (1/12)              | x     |       |                     |
| Columba sp.           | Hunted clinically diseased (1) and found clinically diseased (1)                      | 3.6% (2/56)              | x     |       |                     |
| Cygnus cygnus         | Found dead                                                                             | 3.6% (1/78)              | x     |       |                     |
| Clangula hyemalis     | Found dead                                                                             | 1.3% (4/8)               | x     |       |                     |
| Total                 |                                                                                       | 51/412                  |       |       |                     |
### Table 2. Avian coronaviruses described in this study. The strain names, species, clinical status of the bird, date of sampling, location of the sample indicated in the codes of sub-regional units of Finland, and Genbank accession numbers.

| Taxonomic Family of the host | Strain | Species | Clinical status* | Date of sampling | Location | GenBank accession number |
|-----------------------------|--------|---------|------------------|------------------|----------|-------------------------|
| Anatidae                    |        |         |                  |                  |          |                         |
| Avian Coronavirus/Anas platyrhynchos/Finland/8585/2010 | Mallard (Anas platyrhynchos) | 1 | 20 August 2010 | 112 | KX588654 |
| Avian Coronavirus/Anas crecca/Finland/8589/2010 | Eurasian Teal (Anas crecca) | 1 | 20 August 2010 | 112 | KX588655 |
| Avian Coronavirus/Anas platyrhynchos/Finland/8597/2010 | Mallard (Anas platyrhynchos) | 1 | 20 August 2010 | 112 | KX588657 |
| Avian Coronavirus/Anas crecca/Finland/8627/2010 | Eurasian Teal (Anas crecca) | 1 | 20 August 2010 | 112 | KX588658 |
| Avian Coronavirus/Anas platyrhynchos/Finland/8629/2010 | Mallard (Anas platyrhynchos) | 1 | 20 August 2010 | 112 | KX588659 |
| Avian Coronavirus/Anas platyrhynchos/Finland/8764/2010 | Eurasian Teal (Anas crecca) | 1 | 20 August 2010 | 193 | KX588660 |
| Avian Coronavirus/Anas crecca/Finland/8770/2010 | Eurasian Teal (Anas crecca) | 1 | 20 August 2010 | 193 | KX588661 |
| Avian Coronavirus/Anas platyrhynchos/Finland/8782/2010 | Mallard (Anas platyrhynchos) | 1 | 20 August 2010 | 193 | KX588662 |
| Avian Coronavirus/Anas platyrhynchos/Finland/8788/2010 | Eurasian Teal (Anas crecca) | 1 | 21 August 2010 | 193 | KX588663 |
| Avian Coronavirus/Anas crecca/Finland/8790/2010 | Mallard (Anas platyrhynchos) | 1 | 21 August 2010 | 112 | KX588664 |
| Avian Coronavirus/Anas platyrhynchos/Finland/8802/2010 | Mallard (Anas platyrhynchos) | 1 | 20 August 2010 | 112 | KX588665 |
| Avian Coronavirus/Anas platyrhynchos/Finland/8812/2010 | Mallard (Anas platyrhynchos) | 1 | 20 August 2010 | 112 | KX588666 |
| Avian Coronavirus/Anas crecca/Finland/9195/2010 | Eurasian Teal (Anas crecca) | 1 | 28 August 2010 | 152 | NA |
| Avian Coronavirus/Anas crecca/Finland/9203/2010 | Eurasian Teal (Anas crecca) | 1 | 28 August 2010 | 152 | KX588667 |
| Avian Coronavirus/Anas crecca/Finland/9209/2010 | Eurasian Teal (Anas crecca) | 1 | 28 August 2010 | 152 | KX588668 |
| Avian Coronavirus/Lates argentatus/Finnland/9211/2010 | European Herring Gull (Larus argentatus) | 3 | 1 September 2010 | 171 | KX588669 |
| Avian Coronavirus/Anas platyrhynchos/Finland/9362/2010 | Mallard (Anas platyrhynchos) | 1 | 24 August 2010 | 171 | KX588670 |
| Avian Coronavirus/Anas crecca/Finland/9368/2010 | Eurasian Teal (Anas crecca) | 1 | 28 August 2010 | 171 | KX588671 |
| Avian Coronavirus/Anas platyrhynchos/Finland/10485/2010 | Mallard (Anas platyrhynchos) | 1 | 20 August 2010 | 171 | KX588672 |
| Avian Coronavirus/Anas crecca/Finland/10539/2010 | Eurasian Teal (Anas crecca) | 1 | 28 August 2010 | 154 | KX588673 |
| Avian Coronavirus/Anas crecca/Finland/13345/2010 | Mallard (Anas platyrhynchos) | 1 | 2 September 2010 | 71 | KX588674 |
| Avian Coronavirus/Anas crecca/Finland/13347/2010 | Mallard (Anas platyrhynchos) | 1 | 2 September 2010 | 71 | KX588675 |
| Avian Coronavirus/Anas crecca/Finland/13349/2010 | Mallard (Anas platyrhynchos) | 1 | 2 September 2010 | 71 | KX588676 |
| Avian Coronavirus/Anas crecca/Finland/13367/2010 | Mallard (Anas platyrhynchos) | 1 | 2 September 2010 | 71 | KX588677 |
| Avian Coronavirus/Anas crecca/Finland/13369/2010 | Mallard (Anas platyrhynchos) | 1 | 2 September 2010 | 71 | KX588678 |
| Avian Coronavirus/Anas crecca/Finland/13371/2010 | Mallard (Anas platyrhynchos) | 1 | 2 September 2010 | 71 | KX588679 |
| Avian Coronavirus/Anas crecca/Finland/13377/2010 | Mallard (Anas platyrhynchos) | 1 | 2 September 2010 | 71 | KX588680 |
| Avian Coronavirus/Anas crecca/Finland/13385/2010 | Mallard (Anas platyrhynchos) | 1 | 2 September 2010 | 71 | KX588681 |
| Avian Coronavirus/Anas crecca/Finland/13389/2010 | Mallard (Anas platyrhynchos) | 1 | 2 September 2010 | 71 | KX588682 |
| Avian Coronavirus/Anas crecca/Finland/13395/2010 | Mallard (Anas platyrhynchos) | 1 | 2 September 2010 | 71 | KX588683 |
| Avian Coronavirus/Anas crecca/Finland/13399/2010 | Mallard (Anas platyrhynchos) | 1 | 2 September 2010 | 71 | KX588684 |
| Avian Coronavirus/Anas crecca/Finland/13401/2010 | Mallard (Anas platyrhynchos) | 1 | 2 September 2010 | 71 | KX588685 |
| Avian Coronavirus/Cygnus cygnus/Finland/9483/2013 | Whooper Swan (Cygnus cygnus) | 3 | 28 July 2013 | 71 | KX588672 |
| Avian Coronavirus/Cygnus cygnus/Finland/14383/2013 | Long-tailed Duck (Clangula hyemalis) | 3 | 21 October 2013 | 11 | NA |
| Avian Coronavirus/Cygnus cygnus/Finland/14385/2013 | Long-tailed Duck (Clangula hyemalis) | 3 | 21 October 2013 | 11 | NA |
| Avian Coronavirus/Cygnus cygnus/Finland/14387/2013 | Long-tailed Duck (Clangula hyemalis) | 3 | 21 October 2013 | 11 | NA |
| Avian Coronavirus/Cygnus cygnus/Finland/14395/2013 | Long-tailed Duck (Clangula hyemalis) | 3 | 6 September 2012 | 11 | KX588651 |
| Avian Coronavirus/Anas platyrhynchos/Finland/11136/2013 | Mallard (Anas platyrhynchos) | 2 | 2012 | 138 | KX588657 |
| Avian Coronavirus/Lates argentatus/Finnland/9211/2010 | European Herring Gull (Larus argentatus) | 3 | 1 September 2010 | 171 | NA |
| Avian Coronavirus/Lates fuscus/Finnland/10059/2013 | Lesser Black-backed Gull (Larus fuscus) | 3 | 29 July 2013 | 43 | KX588673 |
| Avian Coronavirus/Chroicocephalus ridibundus/Finland/10083/2013 | Black-headed Gull (Chroicocephalus ridibundus) | 3 | 14 August 2013 | 191 | KX588674 |
| Avian Coronavirus/Lates argentatus/Finland/10087/2013 | Black-headed Gull (Chroicocephalus ridibundus) | 3 | 14 August 2013 | 171 | KX588670 |

(Continued)
infections have been reported by other researchers [35], and low pathogenic avian influenza viruses (LPAI) together with type-1 avian paramyxoviruses (APMV) are also commonly detected [36]. The co-infections observed here were in four apparently healthy ducks, three mallards and a teal, suggesting that infections by a number of low virulence viruses are well tolerated by migratory ducks.

CoV infections in mallard ducks (Anas platyrhynchos) have been documented in Sweden [26,35], and one of the studies reported the prevalence of CoV infections to be 6.9% albeit with seasonal variation [35]. In a more recent study, Wille et al. documented the prevalence of CoVs among wild waterbirds in Sweden to be even 18.7% [37]. A study from Norway reported the prevalence of CoV RNA among graylag goose (Anser anser) in Northern Europe to be as high as 38% in 2004, but they also documented great annual and geographical variation; in 2003 the prevalence was only 18% [38]. Also in England, wildfowl (Anseriformes) and waders (Charadriiformes) have been reported to carry CoVs [39]. A report from South Korea documented a prevalence of CoV RNA to be 0.95% in wildfowl (Anseriformes), and they also observed annual variation in the detection of CoV RNA

![Figure 1. A map illustrating the sampling locations. The sampling locations (sub-regional units, year 2013) are indicated in the Table 2, and are here illustrated by numbers on a map of Finland.](image-url)
Based on the studies from South Korea [34] and Northern Europe [38], and the results we obtained, it seems that large annual variation in the circulating RNA seems to be characteristic to avian coronaviruses. A study in China reported Gammacoronavirus and Deltacoronavirus RNA in 12% of the screened, apparently healthy, wild aquatic birds (Anseriformes, Pelecaniformes, and Ciconiiformes) [24], which is in concordance with our results. Another study, conducted in the Beringia area (encompassing areas of Alaska and Siberia), reported Gammacoronavirus RNA in 6.4% of the examined birds (Anseriformes, Pelecaniformes, and Charadriiformes) [23]. Interestingly, some of the sequences found in the Chinese study [24] were identical to those found from the Beringia area [23]. Also the Finnish Duck CoVs reported in this study cluster very closely with duck CoVs of the Gammacoronavirus genus from Siberia and China, which are connected by migratory routes. The close genetic relatedness of these strains (as well as those found in South Korea and England) indicates that migratory birds do have a specific role in dispersing CoV to distinct geographical locations. However, the sequence analyses might be somewhat biased due

Figure 2. Maximum likelihood phylogeny of avian coronaviruses detected in Finland in wild birds. A 464 nucleotide long sequence region of the RdRp gene within Orf1ab was sequenced and aligned with reference sequences: the analysis includes alignments of 46 viruses detected in this study and a diverse set of reference strains obtained through BLAST search. Maximum-likelihood analysis using 1000 bootstrap replicates was used to infer tree topology. The evolutionary model GTR+G + I was used after evaluation of the best-fit model according to Bayesian Information Criterion. Bootstrap support values exceeding 70 are shown next to the nodes. Strains described in this study are marked by dots. Branches have been collapsed where our viruses share high sequence identity, and the branch wideness is relative to the number of viruses included.
to the limited availability of reference strains. This and previous studies have mainly been focused on duck and shorebird species of the Northern hemisphere, hence the involvement of other avian species and a wider global distribution of avian CoVs remains unclear.

Interestingly enough our results show that the genus *Larus* (order *Charadriiformes*) can serve as a host for both genera, the *Deltacoronavirus* (species: *Larus fuscus*) and *Gammacoronavirus* (species: *Larus argentatus*). Also, the fact that we detected a *Deltacoronavirus* in *Chroicocephalus ridibundus* is interesting, since in another study [23] *Gammacoronavirus* RNA was found from the same bird species. It was previously reported, that the bird species from the order *Anseriformes* can be infected with both delta- and gammacoronaviruses [24], and we now demonstrate that the same applies for the order *Charadriiformes*. The abundance of the detected CoV sequences originated from ducks supports the hypothesis that the Duck CoVs would persist as endemic in duck populations causing subclinical infections similarly to LPAI and APMV.

To conclude, we show that there is a wide range of gammacoronaviruses and also some deltacoronaviruses circulating in wild birds in Finland. In addition, we show that the number of CoV-positive birds detected each year varies greatly. Due to the high recombination rate of CoVs, new species constantly arise that are able to adapt to new hosts and ecological niches [5]. Since the number of CoV sequences detected from birds was so abundant, it would be interesting to study what types of CoVs are circulating in small Finnish mammals, especially those in close contact with birds.

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**Disclosure statement**

No potential conflict of interest was reported by the authors.

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