

Comparative survey of canine parvovirus, canine distemper virus and canine enteric coronavirus infection in free-ranging wolves of central Italy and south-eastern France

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Abstract Diseases likely affect large carnivore demography and can hinder conservation efforts. We considered three highly contagious viruses that infect a wide range of domestic and wild mammals: canine parvovirus type 2 (CPV-2), canine distemper virus (CDV) and canine enteric coronaviruses (CECoV). Infection by either one of these viruses can affect populations through increased mortality and/or decreased general health. We investigated infection in the wolf populations of Abruzzo, Lazio e Molise National Park (PNALM), Italy, and of Mercantour National Park (PNM), France. Faecal samples were collected during one winter, from October to March, from four packs in PNALM ($n=79$) and from four packs in PNM ($n=66$). We screened samples for specific sequences of viral nucleic acids. To our knowledge, our study is the first documented report of CECoV infection in wolves outside Alaska, and of the large-scale occurrence of CPV-2 in European wolf populations. The results suggest that CPV-2 is enzootic in the population of PNALM, but not in PNM and that CECoV is episodic in both areas. We did not detect CDV. Our findings suggest that density and spatial distribution of susceptible hosts, in particular free-ranging dogs, can be important factors influencing infections in wolves. This comparative study is an important step in evaluating the nature of

possible disease threats in the studied wolf populations. Recent emergence of new viral strains in Europe additionally strengthens the need for proactive monitoring of wolves and other susceptible sympatric species for viral threats and other impairing infections.

Keywords *Canis lupus* · Infectious disease · Host density · Emerging viral strain · Free-ranging dog

Introduction

Disease-induced population decline has been reported in several large carnivore species, and most epidemic events appear to be caused by viruses (Murray et al. 1999). Even when the effects of diseases are not epizootic and are apparently sublethal, pathogens may affect the size or resilience of infected host populations and increase the probability of decline caused by other factors (Cleaveland et al. 2002). When infections are endemic in reservoir hosts and transmitted horizontally among taxa, the threat of disease epidemics in large carnivores can be important, as demographic effects of disease may occur regardless of host population size or disease transmission rate (Murray et al. 1999). Spread of existing and emerging pathogens in free-ranging animals can cause rapid changes in the abundance and genetic diversity of susceptible populations (Altizer et al. 2003).

Canine parvovirus type 2 (CPV-2) and canine distemper virus (CDV) are well-known pathogens of canids and are reported to occur in several free-ranging wolf populations in Europe and around the world (Kreeger 2003; Zarnke et al. 2004; Frölich et al. 2005; Sobrino et al. 2008; Almberg et al. 2009; Santos et al. 2009; Di Sabatino et al. 2014). Infection by *Alphacoronaviruses* (ACVs) is poorly documented in free-ranging canid populations. The variants of ACVs infecting canids are canine enteric coronaviruses type I and II (CECoV).

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Canine enteric coronaviruses have been detected in wolves only in Alaska (Zarnke et al. 2001). Nevertheless, this rapidly evolving virus appears to be enzootic worldwide in dogs (Pratelli 2006) and has recently received increased attention in Europe (Benetka et al. 2006; Buonavoglia et al. 2006; Decaro and Buonavoglia 2008; Decaro et al. 2008). Transmission of numerous viruses is highly influenced by local carnivore densities (Murray et al. 1999). Dogs can transmit each of these three pathogens to other carnivores, through direct contact (e.g. saliva) or contact with contaminated material (e.g. faeces, vomitus; Kreeger 2003), and might therefore introduce and/or help maintain infection in susceptible wolf populations.

Canine parvovirus type 2, CDV and CECoV are highly contagious pathogens that can infect a wide range of domestic and free-ranging species (Deem et al. 2000; Buonavoglia et al. 2006; de Oliveira Hübner et al. 2010; Nandi and Kumar 2010). Infection by each of these viruses has been associated with high morbidity and mortality in domestic and free-ranging carnivores including wolves (Johnson et al. 1994; Pence 1995; Gese et al. 1997; Di Sabatino et al. 2014), with usually more severe symptoms reported in young individuals (Murray et al. 1999; Deem et al. 2000; Kreeger 2003; Pratelli 2006; Almberg et al. 2009; Nandi and Kumar 2010; Mech and Goyal 2011). Canine distemper infection typically causes pneumonia, encephalitis and/or diarrhoea (Murray et al. 1999; Deem et al. 2000; Kreeger 2003). Canine parvovirus type 2 and CECoV primarily affect the small intestine, causing sometimes severe enteritis and consequent dehydration (Kreeger 2003; Pratelli 2006; Decaro and Buonavoglia 2008). Additionally, foetal or neonatal infection by CPV-2 can trigger severe myocarditis (Kreeger 2003). Systemic infection by CDV or CECoV may also cause various neurological manifestations (Deem et al. 2000; Kreeger 2003; Buonavoglia et al. 2006). Co-infection by CPV-2 and CECoV is associated with more severe symptoms (Decaro et al. 2006; Pratelli 2006). New variants of each of these three viruses have recently been identified in Italy and in other countries of Western Europe, some of which show increased virulence and increased capacity of horizontal transmission (Buonavoglia et al. 2001, 2006; Evermann et al. 2005; Martella et al. 2005; Decaro and Buonavoglia 2008; Le Poder 2011; Monne et al. 2011; Origgi et al. 2012; Di Sabatino et al. 2014). These variants result from mutations and recombination events in local viruses and possibly also from importation of infected animals from other countries (Benetka et al. 2006; Buonavoglia et al. 2006; Demeter et al. 2007; Allison et al. 2012; Origgi et al. 2012). These shared characteristics of CPV-2, CDV and CECoV make them important conservation threats for susceptible host species. The impact of infection on recolonizing wolf populations might be exacerbated compared to large, well-established populations (Johnson et al. 1994).

Wolves typically live in family-based packs, consisting of a mated pair and their offspring of one or several generations,

born in early spring (Packard 2003). The studied wolf subspecies, *Canis lupus italicus* (Randi et al. 2000), is protected. It is only present in Italy and in recently recolonized areas of the Alps in the neighbouring countries (Valière et al. 2003). The wolf never disappeared from central Italy, where small groups of individuals survived the large-scale extermination of the species in Western Europe (Boitani 2003). Since the protection of the carnivore in the 1970s, the population has been recolonizing the alpine range from the Apennines (Lucchini et al. 2002; Fabbri et al. 2007; Ciucci et al. 2009). In France, the first wolf pack settled in Mercantour National Park (PNM) in 1993, after over 50 years of absence (Houard and Lequette 1993).

Most disease surveys on CPV-2, CDV and CECoV use serological investigations to detect specific antibodies. Antibodies indicate previous exposure to an infectious agent but do not provide information on current infection. The virus rapidly disappears from faecal material once active infection is over. In agreement with this, a previous study in Canada showed 100 % seroprevalence of antibodies against CPV-2 in sampled wolves ($n=18$), but absence of the virus in all faecal samples collected from the same population (Stronen et al. 2011). Search for CPV-2 DNA in tissue samples also led to negative results in a large-scale survey of free-ranging carnivores, even though detected antibodies proved previous exposure to the virus in some individuals (Frölich et al. 2005).

Serological investigation on four wolves captured in central Italy in 1993 and 1994 showed previous exposure of four and one individuals to CPV-2 and CDV respectively, whereas no exposure to CECoV has been detected (Fico et al. 1996). Similar results were obtained in captive and free-ranging bears in Abruzzo, Lazio e Molise National Park (PNALM) between 1991 and 1995 (Marsilio et al. 1997). An extended study conducted in Northern Italy in 1994 and 1995 reported the presence of CPV-2 in 3.5 % of the analysed wolf scats (Martinello et al. 1997). A severe CDV outbreak recently spread through part of Europe (Sekulin et al. 2011; Origgi et al. 2012). In Italy, the outbreak was first detected in the north of the country in 2006 and rapidly expanded southwards (Monne et al. 2011). In central Italy, including all around the area of PNALM, the death of 20 wolves was recently attributed to infection by CDV; five of these animals were also infected by CPV-2 (Di Sabatino et al. 2014). In PNM, a study investigated the presence of CPV-2 in the wolf population in 1996 and 1997, but was not able to give conclusive evidence of the presence of the virus (Rossi 2000). To our knowledge, no further large-scale viral disease survey has been conducted on this wolf subspecies. In Italy, high seroprevalence rates of CPV-2, CDV and CECoV were lately reported in domestic (Priestnall et al. 2007) and free-ranging (Corrain et al. 2007) dog populations. Spiss et al. (2012) recently showed high seroprevalence of CECoV in a large-scale study of domestic dogs in Austria.

The emergence of very contagious and highly virulent variants of these viruses in Western Europe highlights the crucial role of surveys in wolves in the concerned areas. Identification of potentially harmful pathogens in susceptible populations is an important first step to evaluate and mitigate their potential impact on population dynamics of free-ranging animals (Murray et al. 1999). Understanding which ecological factors shape the spread and severity of diseases can help control the impact of infections on host populations (Murray et al. 1999). The objectives of our study were (a) to investigate the occurrence and spatial distribution of CPV-2, CDV and CECoV infections in wolves in PNALM (Italy) and PNM (France) and (b) to search for environmental correlates of infection in order to help understand and mitigate the spread of the diseases. We expected that viruses would be less widespread in the wolf population of PNM due to its more recent origin, its lower density and/or lower density and spatial distribution of other susceptible sympatric hosts.

Material and methods

Study areas

We conducted this survey on wolves in Abruzzo, Lazio e Molise national park (PNALM), in central Italy, and in Mercantour National Park (PNM), in south-eastern France. The wolf populations of these two study areas are connected through a dispersal corridor (Fig. 1). Both study areas are mountainous, partly forested and at similar latitudes (Table 1). Livestock and different wild ungulate species are present year-round in both national parks. The main prey species of wolves vary with the abundance and accessibility of the ungulate species present on the territory of each pack.

Mercantour National Park is an area only recently recolonized by wolves. In PNM, the density of the wolf population is low compared to PNALM, from where the carnivore never disappeared. Besides this, the study areas differ in the presence, density and/or spatial distribution of other carnivores susceptible to infection by the investigated viruses: brown bears (*Ursus arctos*) are present in PNALM but absent from PNM, and dogs are widespread and very common in the Abruzzo region (Boitani and Ciucci 1995; Boitani et al. 2002; Ciucci pers. com.) while rare and mostly localized in restricted areas (around villages) in PNM. In PNALM, a free-ranging dog population is well established in and around the park. Sheepdogs and livestock protection dogs are present in both study areas, especially in summer. Most dogs are vaccinated in PNM (Luddeni pers. com.), but not in PNALM (Ciucci pers. com.).

Investigated packs

We examined spatial distribution of the viruses through sampling of different packs in each national park, considering \geq two wolves travelling together as a pack. When fieldwork was conducted, seven packs were known to have at least part of their home range within the buffer zone boundaries of both PNALM and PNM (MEEDEM and MAP 2008; Grottoli 2011). Based on the quality and quantity of collected faecal samples, we retained four packs in each national park for our survey (Table 2). The number of animals present in each studied pack was assessed by snow-tracking sessions conducted in winter, complemented in PNM by genetic analyses (Duchamp et al. 2012). In summer, wolf-howling sessions provided data on the reproductive success of these packs (Ciucci and Boitani 2006, 2007; Grottoli 2011; Duchamp et al. 2012). Because samples were collected in winter, they were from individuals of over 6 months of age.

Additionally, we analysed five samples from individuals that dispersed, died or that could not be assigned to one of the packs in PNM, as indicated by genetic analysis of the collected faecal samples (see Miquel et al. 2006; Duchamp et al. 2012 for details). Two of these samples are from the same individual.

Sample collection and identification

In both study areas, wolf scats were collected year round by scientists, local co-workers and rangers (Ciucci and Boitani 2006, 2007; Grottoli 2011; Duchamp et al. 2012) for the purpose of non-invasive molecular tracking or diet analysis. In the present survey, we considered samples collected between the 1st of October 2005 and the 31st of March 2006 in PNM, and between the 1st of October 2006 and the 31st of March 2007 in PNALM. In both study areas, most wolf scats were collected while snow-tracking the studied packs. In the absence of snow cover, samples were collected at known scent posts, at exploited carcasses or during opportunistic surveys along pathways (Grottoli 2011; Duchamp et al. 2012).

In PNALM, multiple criteria were used to conservatively discriminate wolf scats from those of other species, among which a diameter ≥ 2.5 cm and estimated volume ≥ 100 cc (Ciucci and Boitani 1998; Grottoli 2011). Based on mtDNA and nuclear markers (Boggiano et al. 2013), all fresh scats ($n=107$) collected on the snow along wolf trajectories from December 2005 to March 2006 in this study area were from wolves, except for two samples from foxes. This provides a direct validation (98 % accuracy) of the selection criteria adopted (Ciucci pers. com.).

In PNM, genetic data based on a set of seven microsatellite loci were available from previous pilot studies in France. These allowed the discrimination of wolf scats from those of

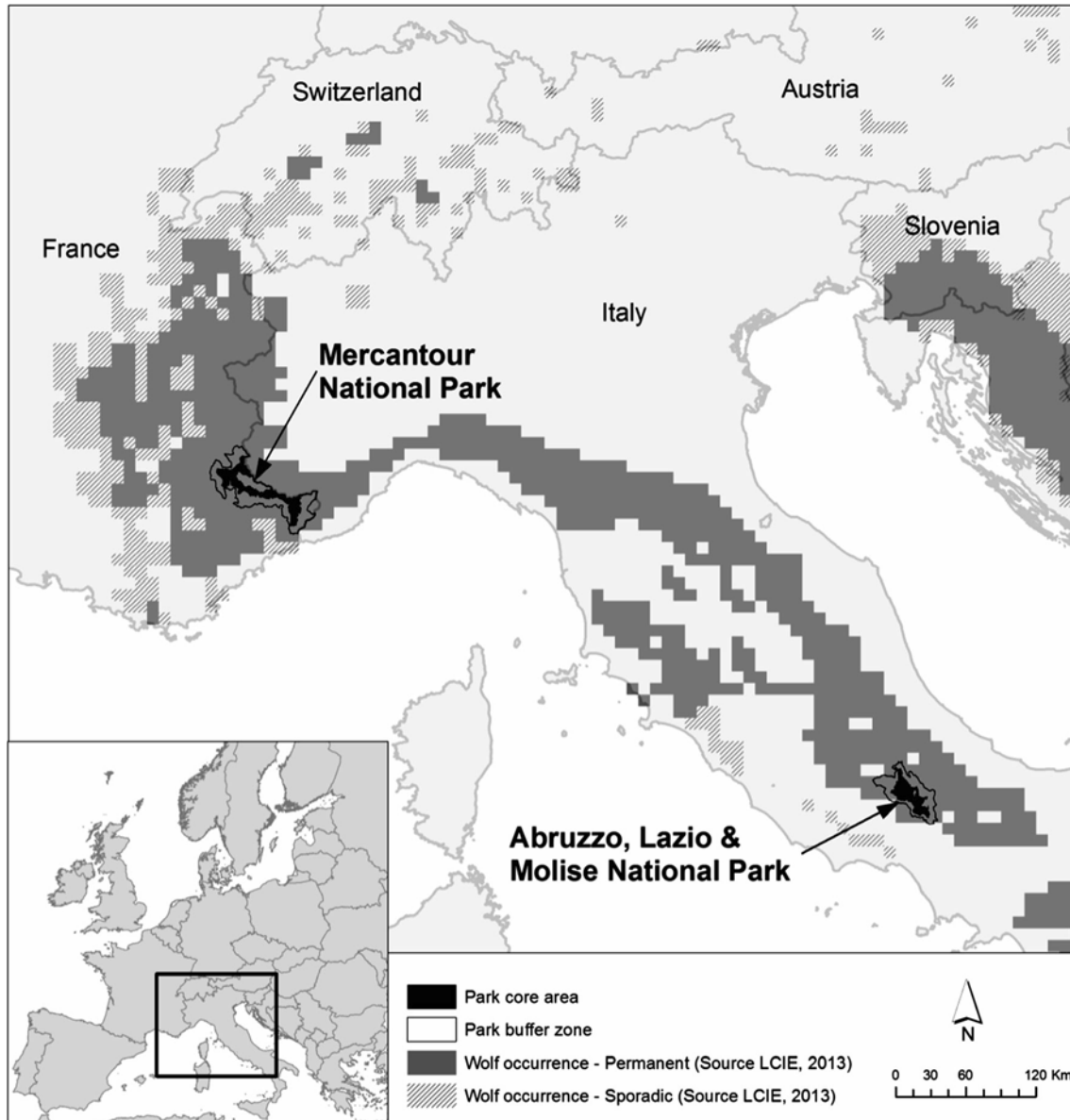


Fig. 1 Study areas and wolf distribution in 2012, showing the dispersal corridor between Italy and France. Redrawn from Kaczensky et al. (2013)

other species (Valière et al. 2003) and the identification of the sex and identity of most contributing animals by the detection

of individual genotypes from faecal samples (see Miquel et al. 2006; Duchamp et al. 2012 for details).

Table 1 Characteristics of the study areas

Study area (winter)	Park creation year	Location (coordinates)	Mountain range	Wolf presence/return ^a	Wolf density in studied winter (ind./1,000 km ²)
PNALM (2006–2007)	1923	Central Italy (41°76' N; 13°84' E)	Apennines	Always present	40–50 ^b
PNM (2005–2006)	1979	South-eastern France (44°18' N; 7°05' E)	Alps	Since 1992	11.5 ^c

PNALM Abruzzo, Lazio e Molise National Park, Italy, *PNM* Mercantour National Park, France, *ind.* individuals

^aHouard and Lequette 1993; Boitani 2003

^bMinimum estimated values (Ciucci and Boitani 2006, 2007)

^cWolf density was calculated as the mean number of wolves per pack divided by the mean estimated size of the territory of the packs in the park (estimated territory size: 260–350 km², ONCFS Réseau Loup/Lynx 2006; Duchamp et al. 2012)

Table 2 Ecological characteristics of investigated packs in PNALM (Abruzzo, Lazio e Molise National Park, Italy) and PNM (Mercantour National Park, France)

Study area and packs	Detected repr. in S_{t-1} ^a	Individuals/pack in W_t ^b	Detected repr. in S_t ^c	Individuals/pack in W_{t+1} ^d
PNALM (year)	(2006)	(2006–2007)	(2007)	(2007–2008)
Iorio	Yes	6	Yes	4
Mainarde	Yes	9	Yes	5
Orsara	Yes	3	Yes	6
Villavalelonga	Yes	7	Yes	6
Total		25		21
PNM (year)	(2005)	(2005–2006)	(2006)	(2006–2007)
Haute Tinée	Yes	3–4	No	2–4
Moyenne Tinée	No	2–3	No	2
Vésubie-Roya	No	3–5	Yes	4–5
Vésubie-Tinée	Yes	3–5	Yes	3–5
Total		11–17		11–14

Repr reproduction, W_t winter of sample collection, S_t summer following sample collection

^{a, c} Detected presence of pups during the summer preceding sample collection (S_{t-1}) and during the summer following sample collection (S_t ; ONCFS Réseau Loup/Lynx 2005, 2007; Grottoli 2011)

^{b, d} Pack size in winter of sample collection (W_t) and in winter following that of sample collection (W_{t+1}). Pack-size estimates are based on snow-tracking sessions in PNALM (Ciucci and Boitani 2007, 2008; Grottoli 2011), and on snow-tracking sessions and genetic analyses performed on faecal samples in PNM (ONCFS Réseau Loup/Lynx 2006, 2007; Duchamp et al. 2012)

We analysed only well preserved samples. We did not retain for analysis faecal samples that were partly consumed by birds, dried out, or exposed to rain or to temperatures obviously above freezing point. We excluded samples composed mostly of hair (estimated as >90 % of the scat volume), as well as scats over-marked with urine or lying less than 50 cm away from another scat. On the day of collection, all samples were stored at $-20\text{ }^{\circ}\text{C}$ in labelled plastic bags and kept frozen until analysis.

Nucleic acid screening and sequence analysis

To assess effective infection by the viruses, and not only exposure of the individuals, we screened collected scats for specific sequences of viral nucleic acids. Before extraction of nucleic acids, samples were vortexed for 1 min and centrifuged at $4,000\times g$ for 10 min; 140 μl served as template using a commercially available kit (QIAamp viral RNA Kit, Qiagen, Hilden, Germany, suitable to extract viral RNA as well as DNA) following the manufacturer's instructions. Extracts were stored at $-80\text{ }^{\circ}\text{C}$.

Detection of CDV specific RNA was carried out with the primers PP-I p1 and p2 described by Frisk et al. (1999). RT-PCR assays were run in a volume of 20 μl (18.4 μl reaction mixture, OneStep RT-PCR Kit, Qiagen; 1.6 μl template) and a primer concentration of 0.4 μM . The thermocycler scheme consisted of two pre-PCR steps of $50\text{ }^{\circ}\text{C}$, 30 min and $94\text{ }^{\circ}\text{C}$, 15 min followed by 40 cycles of denaturation ($94\text{ }^{\circ}\text{C}$, 30 s), annealing ($58\text{ }^{\circ}\text{C}$, 30 s) and extension ($72\text{ }^{\circ}\text{C}$, 1 min) and a final extension ($72\text{ }^{\circ}\text{C}$, 10 min). For the detection of CECoV specific

nucleic acids (member of genus *Alphacoronavirus*) realtime-PCR was performed using the primers and probe described by Gut et al. (1999), who indicate a high cross-reactivity among *Alphacoronaviruses*. Canine parvovirus type 2 specific nucleic acids were detected by realtime-PCR with primers and probe described by Decaro et al. (2005). Negative controls, consisting only of the components of the kit, were run together with the samples through all procedure steps. Equivocal results were not considered in the interpretation of data.

Sequencing was performed on 409 bp of the CCoV gene (primers CCoV1 and CCoV2 according to Pratelli et al. 2002) and 683 bp of the CPV-2 gene (primers F/CPV-2F: 5'-ATGG AGCAGTTCAACCAGAC-3' and F/CPV-2R: 5'-TGTTGG TGTGCCACTAGTTC-3'). Amplified DNA was extracted using a commercially available kit (QIAquick[®] PCR purification kit) following the manufacturer's instructions and served as template for sequencing PCR, which was carried out in a volume of 20 μl with a ready to use sequencing PCR mixture (DNA Sequencing Kit). Forward and reverse sequences of PCR products were analysed using ABI Prism 310 Genetic Analyser.

Prevalence and confidence interval

Prevalence refers to N_{Pos}/N , with N_{Pos} the number of wolf scats in which viral nucleic acids of CPV-2, CDV or CECoV were detected, and N the total number of samples analysed for the considered virus within each study area. As not all scats are statistically independent (i.e. multiple analysed faecal

samples are from the same individuals), calculated prevalence does not represent prevalence of the virus in the populations and cannot be interpreted as such. However, we used these values in a simple comparison of the study areas. We calculated confidence intervals (95 %) following a binomial distribution for large sample (Sokal and Rohlf 1995), using R 2.15 (R Core Team 2013).

Results

We analysed 79 wolf faecal samples from PNALM and 66 from PNM, collected from four packs in each study area (Table 3). In PNM, we also analysed five samples from individuals that could not be assigned to a pack, dispersed or died. In order to confirm specificity, we analysed randomly selected sequences of five CPV-2 and two CECoV positive samples. The homology between the five CPV-2 sequences was 99.5–100 % and 99 to 99.3 % to CPV-2 strain C-780916 (American Type Culture Collection ATCC VR-953). The sequences of the two CECoV positive samples were up to 98–99 % homolog to various CECoV sequences available in GenBank (EU856361.1, DQ112226.1, EU924791.1 and EU924790.1).

We identified CPV-2 in all four packs in PNALM ($n=12$, prevalence=15.2 %) and in two different packs in PNM ($n=8$, prevalence=12.1 %; Table 3). In PNM, all positive samples but one were from the Haute Tinée pack, in which the two females and the one male all shed CPV-2 DNA in their faeces. Another positive sample was from a male in the Vésubie-Tinée pack. We detected CECoV in two packs in PNALM ($n=7$, prevalence=8.9 %) and one pack as well as two other individuals in PNM ($n=4$, prevalence=6.1 %). In PNM, positive samples from identified individuals were from the Haute Tinée pack ($n=1$) and a dispersing male ($n=2$). We did not detect nucleic acids of CDV in the analysed samples.

We detected both CPV-2 and CECoV in two samples from the Italian Villavalelonga pack. Considering infections at the pack level, we detected both CPV-2 and CECoV in two packs in PNALM and in one pack in PNM (Table 3).

Discussion

The present study is the first large-scale multi-viral infection survey conducted on wolves in Italy or in France through non-invasive techniques and is among the rare investigations on wolves in Western Europe (Martinello et al. 1997; Sobrino et al. 2008; Santos et al. 2009; Di Sabatino et al. 2014). To our

Table 3 Detection of CPV-2, CDV, and CECoV nucleic acids in wolf scats from studied packs in PNALM (Abruzzo, Lazio e Molise National Park, Italy) and PNM (Mercantour National Park, France)

National parks and investigated packs/individuals	CPV-2			CDV	CECoV			N
	N _{Pos}	P	CI		N _{Pos}	N _{Pos}	P	
PNALM (2006–2007)	12	15.2	(7.0–23.4)	0	7	8.9	(2.6–15.2)	79
Iorio	3			0	0			13
Mainarde	4			0	3			27
Orsara	2			0	0			19
Villavalelonga	3			0	4			20
PNM (2005–2006)	8	12.1	(4.2–20.0)	0	4	6.1	(0.3–11.9)	66
Haute Tinée	7			0	1			18
Moyenne Tinée	0			0	0			12
Vésubie-Roya	0			0	0			10
Vésubie-Tinée	1			0	0			21
Dispersed/died/unidentified ^a	0			0	3 ^b			5
Total	20			0	11			145

Positive (N_{Pos}) results are illustrated, together with the prevalence (P) of each virus in each study area and the corresponding 95 % confidence intervals (CI). P and CI are expressed as percentages (%). N: total number of faecal samples analysed

CPV-2 canine parvovirus type 2, CDV canine distemper virus, CECoV canine enteric coronaviruses

^a Samples from individuals that dispersed, died or that were not assignable to one of the packs in PNM, as indicated by genetic analyses of microsatellite DNA

^b Two of the 3 positive samples are from a single individual, as indicated by genetic data obtained through the analysis of the collected faecal samples (see Miquel et al. 2006 and Duchamp et al. 2012 for details)

knowledge, our findings are the first reported CECoV infections in wolves outside Alaska. Although exposure of wolves to CPV-2 was previously reported in France (based on a few opportunistic necropsies of dead animals; Duchamp and Gauthier, unpublished data) and in Italy (Fico et al. 1996; Martinello et al. 1997; Di Sabatino et al. 2014), our study provides the first conclusive evidence of CPV-2 infection in several established wolf packs in these two countries. Previous opportunistic testing of animals found dead revealed no infection by CDV or CECoV in French wolves, whereas similar investigations recently detected CDV in wolves around PNALM (Di Sabatino et al. 2014). Given the possible negative impact of these viruses on canid populations, and because only little information is available on *C. lupus italicus*, the results from our study are important for conservation management and highlight the need for continued monitoring.

We used molecular investigations to detect viral nucleic acids in faecal samples from PNM and PNALM. Whereas this technique is unable to detect previous exposure as indicated by specific anti-viral antibodies, it detects recent infection of individuals (Martinello et al. 1997; Murray et al. 1999). The same molecular techniques were used for each virus and analyses were undertaken in a single laboratory, ensuring consistency of results and thus enabling direct comparison of the two studied populations. As most previous studies investigated exposure to viruses through serological surveys, derived findings cannot be directly compared with our results.

Close physical contact between group members is characteristic of social canids such as wolves and greatly enhances within-pack transmission of pathogens (Johnson et al. 1994). Pack members regularly use urine and faeces to mark their territory (Harrington and Asa 2003) and inspection of faecal markings is frequent along territory edges. Investigation of the ano-genital area of conspecifics is part of common social interactions (Harrington and Asa 2003). These behavioural characteristics of wolves enhance oro-faecal transmission of pathogens between individuals. Therefore, and given that these viruses are highly contagious, the detection of CPV-2 or CECoV in one or more samples from a pack suggests that several members of that pack were probably infected. Thus, we discuss our results mostly based on the infection at the pack-level.

The wolf populations of PNALM and PNM are connected through a dispersal corridor (Ciucci et al. 2009; Falcucci et al. 2013) since over 20 years, and the region separating the two study areas is home to several widely distributed alternative susceptible host species (e.g. the red fox—*Vulpes vulpes*). Therefore, similar infection rates of wolves by the highly contagious studied viruses could be expected in the two areas. However, differences between PNALM and PNM in the density of wolves as well as the presence and density of other susceptible host species may be important ecological factors

shaping the distribution of viruses in the environment, and consequently, the exposure of wolves to these pathogens. In particular, the spatial distribution of dogs, widespread in PNALM while more localized in PNM, may play a specific role in the transmission of diseases to wolves. In PNALM, an unvaccinated free-ranging dog population lives sympatrically with the studied wolf packs. In the Mercantour area, numerous farm dogs and hunting dogs are reported seropositive to CPV-2, as recorded by local veterinarians (Luddeni pers. com.). This can however be the consequence of vaccination or exposure to the virus in the environment. Among the four studied French packs, the territory of the Haute Tinée pack is the only one that contains a major village in its centre, which may enhance contact rates between wolves and contaminated faeces from domestic dogs. The territory of this pack also lies along one of the main roads crossing the Alps, and is used by many travellers and their pet dogs. Such important anthropogenic influences acting in this specific area may favour transmission of pathogens to wolves through contamination of the environment by domestic dogs, and could explain the detection of both CPV-2 and CECoV in the Haute Tinée pack.

Canine parvovirus type 2

Prevalence of CPV-2 in wolf faecal samples ranged from 12.1 % to 15.2 % in PNM and PNALM respectively. Identification of CPV-2 in all investigated packs of PNALM suggests that the virus is enzootic in that wolf population (Almberg et al. 2009; Mech and Goyal 2011). In PNM, however, CPV-2 was only detected in two packs out of four, indicating that the virus may not yet be established in the whole population.

Although the density of various species susceptible to infection by CPV-2 might be similar in both study areas, the population of free-ranging dogs only present in PNALM may play a significant role in the dissemination of CPV-2 in the environment. Brown bears are also absent from PNM, whereas individuals infected by CPV-2 have been reported in PNALM (Marsilio et al. 1997). Thus, even though CPV-2 is highly resistant (Steinel et al. 2001), contamination of the environment by the virus may be limited in PNM because of a lower density and/or distribution range of other susceptible hosts as compared to PNALM.

Canine enteric coronaviruses

As the *Alphacoronaviruses* (ACVs) group comprises porcine enteric coronaviruses (Decaro and Buonavoglia 2008), the detection, in our study, of ACVs from infected wild boars consumed by wolves cannot be excluded. Indeed, these ungulates are prey species of the carnivore in PNALM (Grottoli 2011) and in PNM. However, only very low prevalence of infection by ACVs is reported in wild boars in Europe

(Vengust et al. 2006; Ruiz-Fons et al. 2008; Sedlak et al. 2008; Kaden et al. 2009), and dilution of viral particles in wolf faeces would further decrease chances of detection. Additionally, our sequencing of randomly selected positive samples from PNALM and PNM indicated specificity for CECoV.

High density of susceptible host populations (Zarnke et al. 2001) and frequent social interactions with conspecifics (Priestnall et al. 2007) enhance transmission opportunities of CECoV. We identified this virus in two packs in PNALM and one pack in PNM. That CECoV was not detected in all packs suggests that the virus is not enzootic in these wolf populations. Coronaviruses are inactivated in a few days at 37 °C, but remain infective for up to several months at 4 °C and below (Pratelli 2008). In their serological survey of a wolf population, Zarnke et al. (2001) reported that CECoV might mainly be transmitted in wintertime. They also showed that immunity against CECoV seems to be short-lived and disappears rapidly without re-exposure to the virus. Therefore, our results suggest that CECoV is better maintained in the high-density multi-host populations in PNALM and/or that the virus is recurrently introduced into the wolf population by sympatric susceptible hosts. In Italy, exposure of dogs to CECoV is widespread and frequently reported (Pratelli et al. 2003; Priestnall et al. 2007; Decaro and Buonavoglia 2008; Decaro et al. 2008). Thus, the free-ranging dog population of PNALM may serve as a reservoir for the infection or reinfection of wolves by this virus.

Co-infection by CPV-2 and CECoV

Co-infection by CPV-2 and CECoV is known to enhance the severity of symptoms (Evermann et al. 2005; Decaro et al. 2006; Pratelli 2006), and fatal outcomes have been reported in dog pups (Decaro et al. 2006). In PNALM, we detected both CECoV and CPV-2 in two packs and found concurrent infection by both viruses in samples collected in one of them. In the Italian packs, young pup mortality typically caused by CPV-2 and CECoV co-infection did however not result in early loss of entire litters. Once CPV-2 becomes enzootic in a population, its negative impact on pup survival seems to decline (Mech and Goyal 2011). The development of long-lasting, possibly life-long, immunity following CPV-2 infection (Steinel et al. 2001; Mech and Goyal 2011) may have a protective effect on the wolf population of PNALM, and help explain the reproductive success in all four packs, despite the detection of CPV-2 and CECoV in two of them. In PNM, we found both CECoV and CPV-2 only in the Haute Tinée pack, with no sample containing both viruses. Possibly, fatal co-infection with CPV-2 and CECoV can be a determinant factor explaining the absence of surviving pups in the Haute Tinée pack in the summer following that of samples collection. We however also know that one of the mating partners disappeared from the pack during the winter of investigation.

Despite the subsequent detection of a new individual in this pack before the onset of the mating season, it is unclear whether this new pack member replaced the missing partner.

As our results suggest that CPV-2 is not enzootic in the wolf population of PNM, individuals may be more vulnerable to CPV-2 infection as well as to co-infection by CPV-2 and CECoV. This may have contributed to the low detected pup production in two consecutive summers in PNM (2005 and 2006) compared to PNALM (2006 and 2007).

Canine distemper virus

Serological surveys conducted 15 years ago reported exposure to CDV in three out of nine free-ranging brown bears (Marsilio et al. 1997) in PNALM, and in one out of four wolves in a neighbouring geographical area (Fico et al. 1996). Exposure of foxes and badgers to CDV was also documented in the same general area (Di Sabatino et al. 2014). As none of the samples that we analysed tested positive for CDV, our results suggest that the virus was absent from the investigated wolf populations at the time of sample collection. However, that infected individuals typically shed the virus only for 4 to 5 days in their faeces minimizes the chance to detect the pathogen, even from sick animals.

Broad implications

Negative impact of diseases on population dynamics is underestimated, as morbidity and mortality are difficult to evaluate in free-ranging populations (Zarnke et al. 2004). This applies even more to large carnivores, owing to their secretive behaviour (Murray et al. 1999). The extent of the impact depends on the proportions of additive and compensatory diseases-caused mortalities. In social species such as wolves, mortality caused by infections can, however, also affect other important biological parameters, as the social structure of groups.

Infection by either one of the viruses considered in the present study can have considerable effects on population dynamics of susceptible canids through increased mortality and/or decreased general health, and consequently impact dispersal in free-ranging populations (Johnson et al. 1994; Kreeger 2003; Pratelli 2006; Almberg et al. 2009; Nandi and Kumar 2010; Mech and Goyal 2011; Monne et al. 2011; Prager et al. 2012). Large-scale disease surveys are yet seldom undertaken in European wild carnivores. Disease-induced mortality and morbidity in the long-established and saturated wolf population of PNALM can have consequences on a larger scale, slowing down the dispersal dynamic of the species and thus directly affecting the connected and expanding populations.

Our results indicate that CECoV should be widely included in epidemiological surveys in free-ranging canids. Given that

CECoV remains infective for extended periods of time at cold temperatures (Pratelli 2008), this virus might represent a more significant conservation concern in boreal and/or mountain ecosystems experiencing yearly winters conditions. In Europe, this applies to most northern countries and to mountainous areas located at higher elevations, as the Alps and the Apennines.

In the future, the contact between large carnivores and domestic animals, and thus the risks of infectious disease transmission, will probably increase as the consequence of range overlap (Murray et al. 1999) resulting from habitat fragmentation (Cleaveland et al. 2002). Pathogens infecting multiple taxa and those that are highly contagious will be of highest conservation concern (Murray et al. 1999). Wild canids are at specific risk of exposure to diseases, as they share susceptibility to numerous pathogens with the dog, the most abundant carnivore (Randall et al. 2004). Among others, it has been suggested that dogs might be a reservoir for the infection of wild canids by CPV-2, CECoV and CDV (Corrain et al. 2007; Prager et al. 2012; Di Sabatino et al. 2014). This may also be the case for CPV-2 and CECoV infections in wolves in PNALM and PNM. Indeed, when comparing our two study areas, global prevalence of both CPV-2 and CECoV is similar, but the spatial distribution of infection in the packs differs: It is consistent with the presence of dogs, widely distributed in PNALM whereas more localized around villages in PNM. The large population of unvaccinated free-ranging dogs present in Italy (Verardi et al. 2006; Corrain et al. 2007) considerably increases the density of susceptible hosts, and may thus importantly impact the spread and maintenance of canid pathogens in the environment. A study of the feral dog population of Abruzzo, which includes PNALM (Boitani and Ciucci 1995), supports this hypothesis. It reports very low survival rate of pups (30 % and 7.5 % at 70 days and 4 months of age, respectively), and population demography mostly driven by stochastic mechanisms. Such observations could well be explained by infection or co-infection by the viruses considered in our study, and suggest that sympatric dogs may play a significant role in the infection of wolves. As foxes and dogs, jackals are susceptible to infection by canid pathogens including CECoV (Goller et al. 2012) and are even reported reservoirs hosts of CPV-2 and CVD (Aguirre 2009). As the golden jackal (*Canis aureus*) is extending its range in Europe and recently reached northern Italy (Arnold et al. 2012), this additional susceptible host might play an increasing role in the spread of the studied viruses in Europe.

The non-invasive sample collection used in this work is well adapted to our study of free-ranging wolf populations. Additionally, scats collected in winter provide the most representative data, as snow-tracking procedures potentially give access to samples from each individual, independent of the

marking behaviour characteristics of pack leaders (Verardi et al. 2006).

Conclusion

Highly contagious pathogens with important potential for horizontal transmission will be of increasing concern for the conservation of carnivores. Our findings suggest that CPV-2 is enzootic in the wolf population of PNALM but not in PNM, and that CECoV is episodic in both areas. In each study area, infection detected in packs was consistent with the spatial distribution of dogs, which may play an important role in the infection of wolves. We therefore strongly recommend the vaccination of domestic and working dogs, as well as of stray dogs whenever possible. The recently established wolf population of PNM may be more vulnerable to viral infections and less resilient to epizootic events than the long-established population of PNALM. On the other hand, the wolf population of PNALM is at increased risk of exposure to emerging strains of highly virulent viruses, as transmission is more likely in high-density susceptible multi-host populations. Infections in the source population of PNALM can have a direct negative impact on connected recolonizing populations, through decreased survival and dispersal. Future large-scale infectious disease surveys, both in wolves and in other susceptible sympatric species, would help understand the epidemiological and spatiotemporal patterns of infections.

To take potentially harmful diseases into account may also importantly refine demographic modelling of free-ranging large carnivore populations and sharpen our understanding of the population dynamics of these species. Our findings strongly suggest that prospective large-scale longitudinal surveys are essential to monitor and evaluate the spread of viruses, of both established and new strains, and their consequences on the studied populations. They underline the necessity for continued monitoring of viral and other infectious diseases, in conservation programs and in local as well as global management strategies of wolves and other carnivores, as an important first step in attempting to mitigate the impact of infections on wild populations.

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