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## **Neglected vector-borne zoonoses in Europe: Into the wild**

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1 **Neglected vector-borne zoonoses in Europe: into the wild**  
2

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29 **Abstract**

30 Wild vertebrates are involved in the transmission cycles of numerous pathogens. Additionally, they  
31 can affect the abundance of arthropod vectors. Urbanization, landscape and climate changes, and  
32 the adaptation of vectors and wildlife to human habitats represent complex and evolving scenarios,  
33 which affect the interface of vector, wildlife and human populations, frequently with a consequent  
34 increase in zoonotic risk. While considerable attention has focused on these interrelations with  
35 regard to certain major vector-borne pathogens such as *Borrelia burgdorferi* s.l. and tick-borne  
36 encephalitis virus, information regarding many other zoonotic pathogens is more dispersed. In this  
37 review, we discuss the possible role of wildlife in the maintenance and spread of some of these  
38 neglected zoonoses in Europe. We present case studies on the role of rodents in the cycles of  
39 *Bartonella* spp., of wild ungulates in the cycle of *Babesia* spp., and of various wildlife species in the  
40 life cycle of *Leishmania infantum*, *Anaplasma phagocytophilum* and *Rickettsia* spp.

41 These examples highlight the usefulness of surveillance strategies focused on neglected zoonotic  
42 agents in wildlife as a source of valuable information for health professionals, nature managers and  
43 (local) decision-makers. These benefits could be further enhanced by increased collaboration  
44 between researchers and stakeholders across Europe and a more harmonised and coordinated  
45 approach for data collection.

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49 **Key words:** wildlife, zoonoses, arthropod vectors, surveillance

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## 52 **1. Introduction**

53 Wildlife has long been recognized to have a major role in the transmission and maintenance of  
54 zoonotic agents, as most emerging infectious diseases are of wildlife origin (Jones et al., 2008).  
55 However, knowledge of the pathogens that naturally occur in wild animals and their potential to  
56 spread to humans and domestic animals is still scarce (Thompson, 2013). This is particularly true  
57 for microorganisms transmitted by vectors, which have multi-component transmission cycles  
58 affected by the ecology as well as the dynamics and life cycles of both vectors and pathogens  
59 (Hollingsworth et al., 2015). Such transmission systems often include diverse wild vertebrate hosts,  
60 which can serve as reservoirs or amplification hosts for pathogens, as well as a food source for the  
61 hematophagous arthropods themselves.

62 Wildlife disease monitoring for emerging as well as for certain autochthonous, but neglected vector-  
63 borne diseases (VBD), is an essential component of surveillance systems, not only for public health,  
64 but also for veterinary and ecological health (Evensen, 2008; Braks et al., 2014). While the need for  
65 such wildlife disease monitoring programmes is internationally recognised (<http://www.glews.net>)  
66 as the emergence of infectious diseases of wildlife origin is frequently of global concern (Jones et  
67 al., 2008; Keesing et al., 2010; Olival et al., 2017), their surveillance, control and prevention chiefly  
68 require local actions. Work on the ground and allocation of resources is usually focused on local  
69 priorities and interests and subject to short-term planning. Any potential international surveillance  
70 programmes are further hampered by inconsistencies in case acquisition (capture and handling of  
71 animals), sampling strategies, diagnostics and data interpretation, and inadequate wildlife  
72 surveillance infrastructures (Stallknecht, 2007). Here we present the current state of knowledge of  
73 the role of wildlife in the emergence and ecology of a number of neglected vector-borne zoonoses  
74 in Europe. We hope that this critical review will help to promote future international collaborations  
75 focused on the detection, prevention and control of VBD in wildlife.

76

## 77 **2. Wildlife population dynamics and their effects on vector abundance**

78 The restoration of natural habitats under the European Union programme for protected areas  
79 (Natura 2000;<http://ec.europa.eu/environment/nature/>), aimed at establishing a connected network  
80 of natural habitats, as well as the legal protection and reintroduction of many wildlife species, and  
81 certain land use changes (Milner et al., 2006), are expected not only to benefit various wildlife  
82 species, but may also boost vector populations and allow them to spread and establish in new areas.

83 A much-cited example of how wildlife populations affect the abundance of vectors, is that of deer  
84 and *Ixodes ricinus* ticks. The last couple of decades have seen a dramatic increase in the abundance  
85 and geographic distribution of various deer species, particularly roe, red and fallow deer (Milner et  
86 al., 2006; Burbaitė and Csányi, 2009, 2010). While immature stages may feed on a variety of  
87 wildlife hosts, including small rodents, insectivores and birds (Hofmeester et al., 2016), several  
88 studies have documented high infestation levels of deer with all developmental stages of ticks  
89 (Kiffner et al., 2010; Qviller et al., 2013). Deer species are thus considered the main determinants of  
90 tick abundance (Mihalca and Sandor, 2013). In fact, some investigations have reported a direct  
91 correlation between deer and tick numbers (Gilbert et al., 2012; Qviller et al., 2013). Others have  
92 found that once a deer population has reached a threshold level, the number of deer does not  
93 significantly affect tick density, indicating that spread rather than abundance is the main driver for  
94 boosting tick populations (Hofmeester et al., 2017). It is important, however, to differentiate  
95 between the overall abundance of ticks in a habitat and the expected number of ticks questing at any  
96 point in time. According to Dobson and Randolph (2011), the former is boosted by greater host  
97 abundance (particularly in areas recently colonized by deer), while the latter is expected to decline  
98 in areas with high deer densities, as unfed ticks quickly find a new host and spend less time  
99 questing. It is also necessary to stress that the specific level of tick infestation on hosts is dependent  
100 on the host's feeding and roaming behaviour. For example, moose (*Alces alces*), which mainly feed  
101 from branches on trees, harbour fewer ticks on their heads and ears than red and roe deer

102 (Handeland et al., 2013); the latter two species are mainly ground feeders with ample opportunity to  
103 encounter all life stages of *I. ricinus*. In addition, deer represent important vehicles for tick  
104 distribution over long distances (Vor et al., 2010).

105 For other vectors populations, the effects of wildlife host dynamics have been less intensively  
106 studied. For example, sand flies are vectors of *Leishmania infantum*, phleboviruses and other  
107 pathogens in southern Europe, but there is a limited understanding of their spatial distribution of  
108 and relationship with wild host abundance. In particular, information is lacking on sandfly breeding  
109 sites, and trapping is mainly aimed at the adult stages (Felicangeli, 2004). Adult female sand flies,  
110 which are the only developmental stage able to transmit pathogens as they require a blood meal to  
111 develop the eggs, may feed on a wide variety of wild mammal and bird species (Bongiorno et al.,  
112 2003; Millán et al., 2014). In rural areas, sand flies congregate in buildings housing domestic  
113 animals, such as sheep sheds, bird houses and dog kennels (Dantas-Torres et al., 2014; Risueño et  
114 al., 2017). However, they are also found in natural and abandoned habitats such as rabbit burrows,  
115 caves and old ruined buildings, where they rely on wildlife for food. The ability of wildlife to  
116 increase and sustain enormous sand fly populations is demonstrated in the ongoing outbreak of  
117 human leishmaniosis in Fuenlabrada, Madrid, which is associated with an explosion in the  
118 population of hares (*Lepus granatensis*) in green areas integrated into a new housing development  
119 built on agricultural land (Molina et al., 2012; Carrillo et al., 2013).

120 While there are several studies on the role of wildlife on flea dynamics in North America, mostly  
121 focused on the ecology of the plague, such research is scarce in Europe. One survey reported that  
122 70% of all flea species are found on rodents (Medvedev, 2002). For example, fleas in the  
123 Palaearctic region preferentially parasitize voles, gerbils and hamsters (Medvedev and Krasnov,  
124 2006), and to a lesser extent other wildlife hosts such as hares and carnivores (mainly foxes) (Foley  
125 et al., 2017). Moreover, flea abundance positively correlates with host density in many flea-host  
126 associations (reviewed by Krasnov, 2008). On the other hand, more diverse host communities could

127 lead to a decrease in flea prevalence. Krasnov (2008) divided fleas into three main categories: (i)  
128 fleas of poultry, livestock and pets; (ii) fleas of commensal birds and mammals (sparrows, pigeons,  
129 house martins, rats and mice); and (iii) fleas of wild birds and mammals. While the first two  
130 groupings show a relatively uniform flea species composition, species in the third category have a  
131 much more diverse pattern, depending on the specific wildlife composition in the region and its flea  
132 fauna.

133 All of these examples show that a rise in the number of certain wildlife hosts can increase the  
134 abundance and distribution of vectors. In some cases this situation results directly in an increase in  
135 VBD, as shown in the example of sand flies and hares in Spain. Many wildlife host/vector/pathogen  
136 relationships, however, are more complex, particularly if the wildlife host is not a competent  
137 pathogen reservoir and the vector is a generalist. In this case, a boost in wildlife host population can  
138 have a ‘dilution’ effect, i.e. it can reduce the pathogen prevalence in the vectors (Dudek, 2014). It  
139 has also been postulated that reduced biodiversity may favour transmission of vector-borne  
140 pathogens because many severely degraded environments of low biodiversity still abound in rodents  
141 (Dudek, 2014), many of which are competent reservoirs for a multitude of disease agents.  
142 Consequently, the declining biodiversity currently experienced in many habitats all over the world  
143 may be advantageous to certain pathogens and their vectors, potentially increasing the risk of  
144 pathogen exposure (Daszak et al., 2007). However, it should also be borne in mind that there are  
145 natural habitats of low biodiversity, such as bogland or tundra, which do not necessarily represent  
146 high risk VBD areas. Care must be taken therefore when extrapolating from the wildlife  
147 transmission dynamics of one pathogen to another.

148

### 149 **3. Urbanization of wildlife and vectors**

150 Green spaces and corridors in cities and (sub)urban areas not only improve human well-being  
151 (Hansen and Pauleit, 2014) but can also help to mitigate the negative effects of heat waves, air



152 pollution, flooding and possible other health risks (IPCC, 2013). In addition, they can contribute to  
153 conservation strategies for wildlife and biodiversity. For example, forty-eight different mammal  
154 species, from bats to wild boars, have been recorded in Budapest (Tóth-Ronkay et al., 2015). Some  
155 mammal species, such as hedgehogs and squirrels, can reach higher densities in (sub)urban habitats  
156 than rural environments (Reeve, 1994; Tóth-Ronkay et al., 2015).

157 On the other hand, the trend in increasing urban green spaces and spatial expansion of urbanized  
158 areas into agricultural and nature habitats also increases the dispersal and abundance of vectors into  
159 urban areas and their contact with humans (Maetzel et al., 2005; Gassner et al., 2016; Paul et al.,  
160 2016; Vourc'h et al., 2016). As a matter of fact, *I. ricinus* (and to a lesser extent other tick species)  
161 are found in city parks, urban forests, private gardens and other green spaces in and around cities  
162 across Europe (Schorn et al., 2011; Buczek et al., 2014; Hornok et al., 2014; Mancini et al., 2014;  
163 Venclíková et al., 2014; Nelson et al., 2015; Starostzik, 2015; Szekeres et al., 2016). Although tick  
164 densities in these areas are generally low, the risk of acquiring a tick bite can be substantial, because  
165 of the relatively high exposure rates of humans. In fact, a Dutch survey found that approximately  
166 30% of tick bites were acquired in gardens (Mulder et al., 2013). While the ecological and  
167 environmental requirements for the establishment and maintenance of *I. ricinus* in its natural  
168 habitats are well known (Randolph, 2004; Medlock et al., 2013), our understanding of the tick's  
169 enzootic cycle in urban green spaces is very limited. For example, it is conceivable that shade from  
170 buildings, ornamental trees, shrubs and hedges provide protection from desiccation in urban  
171 settings. This could be critical for *I. ricinus* which does not survive for long at humidities below  
172 80% (Randolph and Storey, 1999). On the other hand, where the density of feeding or propagation  
173 hosts is too low to sustain a complete enzootic cycle, tick presence may depend on the continuous  
174 introduction from forest areas, particularly via birds (Hasle, 2013). It is possible that in urban  
175 settings medium-sized mammals, such as hedgehogs, squirrels, and (stone) martens replace deer as  
176 the main 'propagation hosts' (Gern et al., 1991, 1997; Labuda and Randolph, 1999).

177 With regard to sand flies, human residential environments can also provide suitable conditions for  
178 all stages of their life cycle, which takes around 30-45 days depending on environmental  
179 temperature and sand fly species (Killick-Kendrick, 1999; Alexander, 2000; Volf and Volfova,  
180 2011). While most species are predominantly exophagic and exophilic (i.e. feeding and egg  
181 development occur outdoors), adults take cover in houses, cellars and animal buildings, when they  
182 are inactive during the day. Moreover, gardens and other periurban habitats provide ideal breeding  
183 sites with organic matter for the larvae to feed on, and shelter from sunlight and desiccation. Foxes,  
184 which are one of the most common urbanised mammal species in Europe and highly susceptible to  
185 *L. infantum* infection, have long been considered a potential source of introducing *Leishmania* from  
186 sylvatic to domestic environments (Ashford and Bettini, 1987). However, more recent PCR-based  
187 studies have shown that *L. infantum* is also endemic in many other wildlife species such as  
188 mustelids, felines, rodents and lagomorphs that live in close proximity to humans and dogs (Del Río  
189 et al., 2014; Millán et al., 2014). Moreover, domestic and sylvatic *L. infantum* cycles are  
190 bidirectional. For example, the ITS-LOMBARDI *L. infantum* strain -recently isolated from hares in  
191 the Fuenlabrada outbreak-was first identified in a human cutaneous leishmaniosis case in 1987, and  
192 has probably been circulating in this area for some time (Chicharro, et al., 2013; Martín-Martín et  
193 al., 2015).

194 Urbanization has also important effects on mosquito vectors, with some species particularly being  
195 favoured by anthropogenic environmental changes. For example, several *Anopheles*, *Culex* and  
196 *Aedes* spp. easily find suitable habitats in urban areas, not least because of numerous artificial  
197 breeding sites created by humans (e.g. water deposits, swimming pools, gardens) (Ferraguti et al.,  
198 2016) and milder temperatures in winter (LaDeau et al., 2015). In fact, studies on the invasive  
199 mosquito species *Aedes albopictus* have shown that this parasite survives better in anthropized  
200 environments than in its natural habitats (Li et al., 2014; Roche et al., 2015). In anthropically altered  
201 areas, frequency of human bites may thus be increased, with higher rates of transmission of

202 mosquito-borne pathogens (LaDeau et al., 2015). More suitable climatic conditions for the vectors,  
203 and the presence of competent wild birds, also in urban areas, are implicated in the expanding  
204 incidence of West Nile Virus infections in Europe (Semenza et al., 2016).

205 While fleas usually gain access to human habitations via pets and periurban rodents, modern living  
206 conditions, particularly central heating, may help to create microclimate conditions suitable for the  
207 development of pre-imago stages throughout the year (Krasnov, 2008).

#### 209 **4. Examples of neglected zoonotic pathogens and their wildlife reservoirs**

210 ‘Neglected’ pathogens are pathogens characterized by a low level of public awareness, and research  
211 focus and/or funding. Some pathogens may be neglected only in certain geographical areas or  
212 certain hosts. Our review focuses on zoonotic pathogen-wildlife host systems that have received  
213 limited attention in the published literature and for which important knowledge gaps remain. Table  
214 1 provides an overview of these pathogens, their known and suspected vectors and wildlife  
215 reservoirs.

216

##### 217 **4.1. Rodents and flea-transmitted *Bartonella* spp.**

218 The genus *Bartonella* comprises several species that infect a large number of vertebrates,  
219 parasitizing erythrocytes and causing a persistent bacteraemia (Maggi et al., 2012). The main  
220 transmission route is via the faeces of ectoparasites (such as fleas and other hematophagous  
221 arthropods), which can enter the body through superficial scratches on the skin (Buffet et al.,  
222 2013). As a result of improved diagnostic techniques, the reported incidence of zoonotic *Bartonella*  
223 infections has been increasing over the last number of years (Edouard et al., 2015), particularly in  
224 people living under poor hygienic conditions and/or suffering from immunodeficiency (Mosepele et  
225 al., 2012).

226 Contact with wild rodents is likely to be a risk factor for infection, since these animals are the  
227 preferential reservoir hosts of several *Bartonella* species in nature. *B. elizabethae*, associated with  
228 the black rat and Oriental rat fleas (*Xenopsylla cheopis*), and *B. grahamii*, associated with wild mice  
229 and voles and transmitted by rodent fleas, are recognized as zoonotic pathogens (Chomel and  
230 Kasten, 2010). Moreover, the pathogenic *B. quintana* and *B. koehlerae* have been detected in rodent  
231 fleas (Mariè et al., 2006), and several studies have reported rodent infections with as yet unknown

232 genotypes (Silaghi et al., 2016).

233 While fleas are suspected to be the main vectors of *Bartonella* in wild rodent populations (Billeter  
234 et al., 2008), few flea species have been unequivocally shown to be competent vectors of *Bartonella*  
235 spp. A notable exception is *Ctenophthalmus nobilis*, a common parasite of small mammals in  
236 Western Europe and competent vector of *B.grahamii* and *B.taylorii* (Bown et al., 2004). The  
237 situation is further complicated by the fact that fleas only show host preference but no clear host  
238 specificity (Silaghi et al., 2016). The role of other arthropods as potential *Bartonella* vectors and  
239 reservoirs also remains to be elucidated. Reis et al. (2011) experimentally demonstrated the vector  
240 competence of *I. ricinus* for *B. birtlesii*. Moreover, certain *Bartonella* spp., including the rodent-  
241 associated *B. doshiae* and *B. tribocorum*, have recently been isolated from blood samples of human  
242 patients with nonspecific chronic symptoms and history of tick-bite. However, so far it has not been  
243 possible to establish a causal link between *Bartonella* spp., clinical signs, and tick bite (Vayssier-  
244 Taussat et al., 2016).

245 In Europe, *Bartonella* prevalence rates of between 14 and 85% have been reported from various  
246 species of rats, squirrels, voles and mice (Ellis et al., 1999; Bown et al., 2002, 2004; Telfer et al.,  
247 2007a, 2007b; Buffet et al., 2013; Kraljik et al., 2015; Silaghi et al., 2016). According to these  
248 studies, infection prevalence was affected by the level of infestation with the relevant vector, the  
249 rodents' resistance to infection, their population density, contact rates, and certain behaviours that  
250 could facilitate transmission by non-vectorial routes. Moreover, seasonal fluctuations in prevalence  
251 rates may be linked to seasonal activity patterns of various flea vectors, while the length of infection  
252 is dependent on the specific *Bartonella* species present (Telfer et al., 2007a, 2007b).

253 As this brief overview shows, many aspects of the complex interactions between zoonotic  
254 *Bartonella* spp., their wildlife hosts and arthropod vectors are yet to be determined. *Bartonella*  
255 diversity in rodents is particularly challenging, since co-infections with different species or variants

256 in rodent hosts and vectors (particularly in fleas) are very common (Gutiérrez et al., 2015), which  
257 may also have consequences on the transmission dynamics and clinical disease in humans.

258  
259 **4.2. Rodents and other wildlife as reservoirs of *Leishmania infantum***

260 *Leishmania infantum* is a protozoan transmitted by *Phlebotomus* spp. sand flies, causing life  
261 threatening zoonotic visceral Leishmaniosis (VL). In southern Europe, VL affects hundreds of  
262 people every year and is considered the most important disease of dogs (Moreno and Alvar, 2004;  
263 Ready, 2010; Gradoni, 2013). Many wildlife species can be infected by *L. infantum* but, in contrast  
264 to dogs and humans, disease is rarely reported and parasite burdens are often comparatively  
265 low (reviewed by Ashford and Bettini, 1987; Ashford, 1996; Quinnell and Courtenay, 2009;  
266 Antoniou et al., 2013; Del Río et al., 2014; Millán et al., 2014; Roque and Jansen, 2014). Given the  
267 large number of potential reservoirs that share habitats with *Leishmania* spp., it is difficult to  
268 determine which of them can serve as the primary reservoir of infection, capable of maintaining  
269 parasite endemicity indefinitely in the absence of a human or canine host. Moreover, even within  
270 specific host species, there are likely individual differences with regard to susceptibility to infection  
271 and infectiousness, depending on the parasite strain and host intrinsic and external factors (Roque  
272 and Jansen, 2014). The best approach to assess the reservoir status is to demonstrate host  
273 susceptibility to infection and ability to transmit the parasite to the vector by performing  
274 xenodiagnostic experiments. However, hosts able to meet these criteria may still not have primary  
275 reservoir capacity, in which case they are considered secondary reservoir hosts (Quinnell and  
276 Courtenay, 2009).

277 With the exception of hares (*Lepus granatensis*), no wildlife species has so far been associated with  
278 leishmaniosis outbreaks in Europe. Xenodiagnostic experiments have confirmed that hares, rabbits  
279 (*Oryctolagus cuniculus*), black rats (*Rattus rattus*) and the American crab-eating fox (*Cerdocyon*  
280 *thous*) can transmit *L. infantum* to sand flies (Quinnell and Courtenay, 2009; Jiménez et al., 2014).

281 Moreover, it is likely that infected red foxes (*Vulpes vulpes*) and other canids in Europe are also  
282 able to transmit these parasites to sand flies (Ashford and Bettini, 1987). The reservoir role of other  
283 sylvatic species known to be susceptible to *L. infantum* infection, including felines, mustelids,  
284 insectivores, and chiroptera, remains to be determined (Millán et al., 2014).

285 Murine (*Mus musculus*) and hamster (*Mesocricetus auratus*) laboratory models have been  
286 extensively used to investigate the clinical and immunological features of *Leishmania* infections,  
287 and the latter species is highly susceptible to visceralising *L. infantum* infection (Loría-Cervera and  
288 Andrade-Narváez, 2014; Moreira et al., 2016). Similarly in the wild, the potential epidemiological  
289 role of rodents in the *L. infantum* transmission cycle has attracted attention for a long time. In early  
290 experiments in France, Rioux et al. (1968), using non-molecular methods, failed to detect *L.*  
291 *infantum* in over 250 wild rodents including mice (*Apodemus* spp.), dormice (*Glis glis* and *Elyonis*  
292 *quercinus*) and rats (*Rattus* spp.). They were, however, able to infect these species with the parasite  
293 experimentally, noting differences in susceptibility. At the time, *L. infantum* strains had been  
294 isolated from black rats in Italy, showing identical isoenzymatic patterns to those from humans.  
295 Sand flies became infected with the strain after feeding on rats only when the rats were inoculated  
296 with high parasite doses or when immunosuppressed with a hydrocortisone treatment, leading to the  
297 conclusion that black rats are naturally resistant to *L. infantum* infection (reviewed by Ashford and  
298 Bettini, 1987). Several epidemiological studies have since demonstrated the presence of *L. infantum*  
299 DNA and specific antibodies in naturally infected rodents in Europe (Quinnell and Courtenay,  
300 2009; Millán et al., 2014). Infected rodents in these studies originated from areas where dogs and  
301 other wild carnivore primary hosts were also present, so that the rodents' epidemiological role in *L.*  
302 *infantum* transmission could not be confirmed. More recently, *L. infantum* DNA was detected in  
303 spleen samples from 11 out of 71 black rats from the Mediterranean island of Montecristo, a natural  
304 reserve where dogs are absent, suggesting that they may act as an alternative primary reservoir host  
305 (Zanet et al., 2014).

306 In summary, *L. infantum* has a remarkable ability to infect domestic and wild mammals, though  
307 clinical cases in wild animals are rarely observed. Little is known about the role of wildlife species  
308 as potential reservoirs or the degree of interaction between domestic and sylvatic *L. infantum* life  
309 cycles. However, there is strong evidence that anthropogenic disturbance of the vector and wildlife  
310 natural environment can lead to infection build-up and spill-over leading to epidemics in susceptible  
311 humans.

312

### 313 **4.3. Wildlife species with a role in zoonotic anaplasmosis**

314 The zoonotic obligate intracellular bacterium *Anaplasma phagocytophilum* occurs worldwide in the  
315 Northern Hemisphere and is transmitted between different species of vertebrates by the bite of ticks.  
316 In Europe the only known vector is *I. ricinus* (Jahfari et al., 2014; Stuen et al., 2013a). *Anaplasma*  
317 *phagocytophilum* causes granulocytic anaplasmosis in domestic ruminants, horses, dogs, cats, other  
318 mammalian species as well as in humans (Stuen et al., 2013a). Even though granulocytic  
319 anaplasmosis is generally seen as a mild and self-limiting disease, hospitalization and need for  
320 intensive care has been reported from isolated human cases (Dumler, 2012).

321 Although the pathogen has been detected in many vertebrates, including birds, deer, rodents and  
322 insectivores, it is unknown which species actually contribute as reservoir hosts in a significant way  
323 to the complex transmission cycle.

324 For example, few studies to date have focused on the role of ornithophilic hard-ticks in the eco-  
325 epidemiology of *A. phagocytophilum*. There is some evidence that blackbirds (*Turdus merula*) may  
326 be a reservoir host. For one, it is a common avian host of immature *I. ricinus*, mainly due to its  
327 ground-feeding behaviour (Hasle, 2013). For another, it is the most frequently reported bird in  
328 Europe to be infected with *A. phagocytophilum* and the most common species to carry infected  
329 ticks, including larval stages. A potential reservoir role for other bird species was also suggested by  
330 the higher prevalence in ticks collected from avian hosts, compared to those questing in the same



331 habitat in Switzerland (Lommano et al., 2014). However, Jahfari et al. (2014) showed that the *A.*  
332 *phagocytophilum* ecotype which was associated with avian hosts was absent in samples from all  
333 other hosts, indicating that it may be restricted to bird-ornithophilic tick systems. With regard to  
334 ecotypes that circulate in wild mammals and humans, it appears that birds are not an important  
335 source of infection and that their epidemiological role in zoonotic infections may be marginal.

336 Among mammalian wildlife species, roe deer and red deer are well-documented hosts of *A.*  
337 *phagocytophilum* with high prevalence rates reported (Petrovec et al., 2002; Michalik et al., 2009;  
338 Silaghi et al., 2011; Mysterud et al., 2013; Overzier et al., 2013; Stuen et al., 2013a,2013b). They  
339 are thought to significantly contribute to the spread of the organism by providing a persistent  
340 pathogen reservoir, in addition to serving as vehicles for infected and uninfected ticks. However,  
341 whether deer are significant contributors to human granulocytic anaplasmosis in Europe is doubtful,  
342 because clinical cases are only rarely reported (Stuen et al., 2013a).

343 While rodents are often suspected to serve as reservoir for *A. phagocytophilum*, their infection rates  
344 are actually quite low in Europe (Liz et al., 2000; Bown et al., 2003; Hulínská et al., 2004;  
345 Blaňarová et al., 2014; Kallio et al., 2014). In fact, several studies reported a complete absence of  
346 infection in all rodent species screened (Silaghi et al., 2012a; Blaňarová et al., 2014; Svitálková et  
347 al., 2015). Moreover, a xenodiagnostic study concluded that *Apodemus* spp. and *M. glareolus* were  
348 not competent reservoirs for *A. phagocytophilum* (Burri et al., 2014). Some authors consider them  
349 accidental hosts (Obiegala et al., 2014), while others suggest that they may act as hosts only for  
350 certain *A. phagocytophilum* variants (Blaňarová et al., 2014). On the other hand, certain insectivore  
351 species showed significantly higher prevalence rates (Liz et al., 2000; Barandika et al., 2007; Bown  
352 et al., 2011; Silaghi et al., 2012a; Földvári et al., 2014).

353 Several so-called 'niche cycle' have been suggested for *A. phagocytophilum*. These are defined by a  
354 competent reservoir host and at least two tick species, with at least two developmental stages each.  
355 One of these two tick species must be endophilic and specific for the reservoir host in question, and

356 the other one exophilic, with a broad host range. It is hypothesised that established niche cycles are  
357 sufficient to maintain a stable and constant endemic cycle of certain genetic variants of *A.*  
358 *phagocytophilum* in a given geographic area. Such a niche cycle has been proposed for hedgehogs,  
359 which are frequently infested with all three life stages of *I. ricinus* as well as the hedgehog tick,  
360 *Ixodes hexagonus* (Földvári et al., 2011; Pfäffle et al., 2011; Dumitrache et al., 2013; Dziemian et  
361 al., 2015). Other insectivore species, such as the common shrew, also fulfil the criteria for a niche  
362 cycle, as they are frequently infected with *A. phagocytophilum* in addition to numerous larvae and  
363 nymphs of *I. ricinus* and *I. trianguliceps* (nidicolous tick) (Bown et al., 2011). Moreover, niche  
364 cycles have been proposed for certain rodent species (*M. glareolus*, *A. flavicollis*, and *A. agrarius*)  
365 and *I. ricinus*/*I. trianguliceps*, chiefly involving *A. phagocytophilum* genotypes that do not have  
366 zoonotic significance (Blaňarová et al., 2014).

367 It is clear that significant knowledge gaps remain regarding the specific host associations of  
368 zoonotic *A. phagocytophilum* genotypes and the vector competence of various tick species. Without  
369 this information, endemic cycles of zoonotic anaplasmosis in nature will remain obscure.

370

#### 371 **4.4 Wildlife species and *Rickettsia* spp.**

372 *Rickettsiae* are obligate intracellular bacteria, which can be separated in two main groups: the  
373 typhus (TG) and spotted fever group (SFG). Although effective treatments exist for many *Rickettsia*  
374 species, some are still associated with severe, sometimes fatal disease. It is the case of *R. conorii*,  
375 the most pathogenic tick-borne rickettsia in Europe and causative agent of Mediterranean spotted  
376 fever (MSF)(Portillo et al., 2015). Two strains, *R. conorii* Malish and Israeli tick typhus strain, are  
377 associated with human and canine disease (De Sousa et al., 2008; Alexandre et al., 2011; Solano-  
378 Gallego et al., 2015). The ‘kennel tick’ or ‘brown dog tick’, *Rhipicephalus sanguineus*, transmits  
379 both strains. Although this tick has a close evolutionary relationship with domestic dogs and feeds

380 primarily on them, it can survive in a wide range of ecological niches and parasitize many wild and  
381 domestic species (Gray et al., 2013).

382 Serosurveys indicate that wild carnivores are frequently exposed to rickettsiae (Marquez and  
383 Millán, 2009; Lledó et al., 2016; Millán et al., 2016). However, as rickettsial DNA has never been  
384 detected in blood or tissue samples collected from wild carnivores (including genets, red foxes,  
385 martens and badgers) (Márquez and Millán, 2009; Torina et al., 2013; Millán et al., 2016), it is  
386 thought that these animals have a negligible role in the transmission (Millán et al., 2016).

387 In contrast, there is evidence that wild rabbits (*O. cuniculus*) and hares (*Lepus europaeus* and *L.*  
388 *granatensis*) have a role as reservoir hosts for *R. conorii* and also in the circulation of other  
389 *Rickettsia* species (Le Gac, 1966; Ruiz-Beltrán et al., 1992; Rovero et al., 2008), such as *R. slovaca*  
390 (Reháček et al., 1978). In Italy, antibodies against *R. conorii* and *R. slovaca* were detected in wild  
391 rabbits, and inoculation of guinea pigs with homogenates of *Rhipicephalus pusillus* ticks isolated  
392 from these rabbits resulted in a seroconversion to *R. conorii* (Ciceroni et al., 1988). However, due to  
393 cross-reactivity of antibodies within the SFG, these experiments do not unequivocally confirm the  
394 role of *Rh. pusillus* as a vector of *R. conorii*. To date, there is no further evidence that this tick  
395 species, which is a common ectoparasite of wild rabbits in the Mediterranean region, serves as a  
396 vector for *R. conorii*, although it is known to transmit *R. sibirica mongolotimonae* in Portugal,  
397 Spain, and France (De Sousa et al., 2006; Toledo et al., 2009; Parola et al., 2013).

398 Small mammals, particularly mice and voles, have also been considered important in the natural  
399 transmission cycle of certain SFG *Rickettsia* species including *R. slovaca*, *R. felis* and *R. helvetica*  
400 (Reháček et al., 1976, 1992; Schex et al., 2011; Martello et al., 2013). In contrast, other authors failed  
401 to identify *R. monacensis* and *R. helvetica* in the blood of *Apodemus* spp., *M. glareolus* and  
402 xenodiagnostic ticks, although attached *I. ricinus* tested positive, and suggested that these rodents  
403 are not reservoirs for SFG species (Burri et al., 2014; Biernat et al., 2016).

404 More recently, studies have shown that lizards may have a more prominent role as reservoirs than  
405 previously thought. In fact, systemic infection by *R. helvetica* was detected in lizards captured in  
406 Portugal and Italy (De Sousa et al., 2012; Tomassone et al., 2017).

407 In conclusion, a better understanding of the eco-epidemiology of rickettsial disease in specific  
408 geographic regions may help to reduce and even prevent outbreaks. For example, an imbalance in  
409 the rodent population in Porto Santo Island, Madeira, Portugal, lead to increased incidence of  
410 human cases of murine typhus, a disease caused by *R. typhi* and transmitted by the rat flea; weather  
411 conditions and human interventions were shown to cause such imbalance (Bacellar et al., 1998).  
412 The factors that affect the abundance, distribution and density of wild reservoir hosts and vectors,  
413 which can conspire to reduce or increase rickettsial infection rates, deserve detailed investigations.

414

#### 415 **4.4. Wild ungulates and *Babesia* spp.**

416 It is thought that the main - if not the only - vector for human babesiosis in Europe is *I. ricinus*  
417 (Gray et al., 2010). In contrast to the relatively high incidence of human babesiosis in the USA, the  
418 number of cases in Europe has remained extremely low. To date, less than 50 cases have been  
419 reported (Hildebrandt et al., 2013; González et al., 2015; Mørch et al., 2015). These include about  
420 40 cases attributed to the cattle parasite *Babesia divergens*, three to *B. venatorum* (formerly 'EU1-  
421 3') and just two autochthonous case attributed to *B. microti* (Hildebrandt et al., 2007; Arsuaga et al.,  
422 2016). Infections with *B. divergens* are mostly confined to asplenic patients, where they are  
423 characterized by septic fever, severe anaemia, haemoglobinuria and jaundice due to widespread  
424 haemolysis. By comparison, infections with the other two *Babesia* spp. appear to be less severe  
425 although all four reported cases also occurred in asplenic or immunocompromised patients  
426 (Herwaldt et al., 2003; Häselbarth et al., 2007; Gray et al., 2010). In contrast to this very low  
427 clinical incidence, significant seroprevalence rates have been recorded in many parts of Europe,  
428 particularly among people with a high risk of occupational exposure such as foresters, hunters,

429 farmers and veterinarians, or those with a history of tick-bite and/or tick-borne disease (Gorenflot et  
430 al., 1998; Foppa et al., 2002; Hunfeld et al., 2002; Gabrielli et al., 2014; Żukiewicz-Sobczak et al.,  
431 2014; Lempereur et al., 2015), indicating that human infection with *Babesia* spp. is not such a rare  
432 event, but that immunocompetent individuals may be largely resistant to disease. Alternatively, it is  
433 also conceivable that many of the *Babesia* species/subspecies carried by *I. ricinus* in Europe are not  
434 infectious to humans although they may cause seroconversion.

435 A large number of epidemiological surveys have screened deer blood or spleen for the presence of  
436 *Babesia* spp. using PCR protocols targeting the 18S rRNA gene. Sequence analysis revealed a  
437 bewildering array of strains and/or species many of which were described as ‘*B. divergens*-like’.  
438 However, detailed investigation of several human, bovine and deer isolates indicated that only  
439 isolates that were over 99.9% identical with the *B. divergens* reference sequence (U16370, a cattle  
440 isolate) shared the biological characteristics of this species, i.e. they were infective to gerbils *in vivo*  
441 and could be cultured in cattle, human and sheep red blood cells *in vitro* (Malandrin et al., 2010). In  
442 fact, all European *B. divergens* human isolates were homologous with U16370 by at least 99.94%.  
443 In contrast, other isolates, although 99.77% identical with U16370 and morphologically and  
444 serologically indistinguishable from *B. divergens*, were not infective to gerbils and could only be  
445 maintained in roe and fallow deer red blood cells. On the basis of these biological characteristics  
446 they were identified as *B. capreoli* (reference sequence AY726009), a species that is not considered  
447 to be zoonotic due to its inability to develop in human red blood cells *in vitro* (Malandrin et al.,  
448 2010). Only a very small number of deer isolates in the database, all from red deer, are over 99.9%  
449 homologous with *B. divergens* (Zintl et al., 2011) indicating that the role of red deer as a potential  
450 reservoir host for *B. divergens*, warrants further investigation. With regard to *B. venatorum*, isolates  
451 100% identical to the reference sequence AY046575 have only ever been identified from roe deer,  
452 the accepted reservoir host for this species.

453 Considering the high degree of genetic homology between *B. divergens* and *B. capreoli* in spite of  
454 their marked biological differences, it is possible that at least some of the numerous isolates that  
455 have been described in deer represent species in their own right. Unless they are assessed for their  
456 ability to infect human red blood cells, we cannot evaluate their potential public health risk.  
457 Furthermore the lack of sequence data for zoonotic babesias (currently there are only six human  
458 18S rRNA isolates in the database), should be addressed in order to determine their relationship  
459 with *Babesia* strains or species harboured by deer.

460

## 461 **5. Directions for future research and conclusion**

462 There is a tendency in the published literature to extrapolate from knowledge gained from  
463 intensively researched VBD to other less well-known pathogens. However, as this review shows,  
464 the relationship between pathogens, vector and wildlife hosts are often highly specific requiring a  
465 much more focused approach.

466 Moreover, effective and timely action in response to endemic and emerging zoonotic wildlife  
467 pathogens is only possible if potential hosts are routinely monitored (Mörner et al., 2002). This is  
468 not the case for the vast majority of VBD, particularly those that are not considered of major public  
469 health importance. Mannelli et al. (2012), reviewing surveillance activities by competent authorities  
470 across Europe, concluded that the only vector-borne zoonotic pathogens for which sufficient data  
471 was being collected were *Francisella tularensis* and West Nile virus, although some others  
472 (including *Borrelia burgdorferi*, *L. infantum*, tick-borne encephalitis virus, Crimean-Congo  
473 haemorrhagic fever virus) were also being recorded.

474 We propose that this conflict between the considerable research effort needed to monitor rare VBD  
475 and the lack of resources generally available for ‘neglected pathogens’ may be addressed, at least in  
476 part, by developing standardised guidelines for data collection and analysis, and a pan-European  
477 repository where up-to-date surveillance data is made available to all stakeholders. The logistics and

478 organisation of this network could replicate those of other similar European collaborations such as  
479 the EFSA/ECDC funded VectorNet initiative (<https://vectornet.ecdc.europa.eu/>), created to monitor  
480 the geographic distribution of arthropod disease vectors. This network could also be used to identify  
481 surveillance gaps and overlaps and further improve cost effectiveness by establishing links between  
482 researchers and organisations that routinely capture and/or cull wildlife (e.g. bird ringing, wildlife  
483 vaccination or rehabilitation centres, hunting management centres). This will no doubt result in  
484 significant cost savings as well as a better understanding of the current status of VBD.

485 From a technical point of view, most studies have relied chiefly on the molecular screening of  
486 potential vectors and wildlife hosts for the presence of certain pathogens. Recent years have seen  
487 the development of various novel blood meal analysis tools, such as stable isotope analysis, high-  
488 throughput sequencing, MALDI-TOF MS and high-resolution melting analysis (Schmidt et al.,  
489 2011; Campana et al., 2016; Collini et al., 2016; Niare et al., 2017). These methods, in addition to  
490 more traditional PCR-based assays, can be used to identify the last host a vector has fed on.

491 Inclusion of these novel tools in standard VBD surveys would thus contribute vital information on  
492 vector feeding habits and transmission cycles.

493 Finally, it is incumbent on the scientific community to demonstrate to policy makers and funding  
494 bodies the inherent value of disease surveillance and research into wildlife, as it shares many living  
495 spaces, pathogens and arthropod vectors with us.

496

497

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507



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1041

1042 **Tables.**

1043 Table 1. List of the zoonotic pathogens that are discussed in this review and their known

1044 (suspected) vectors and reservoir hosts

1045