



Opinion piece



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Special Feature: A contribution to our Special Feature on Disease Ecology organised by Joseph Hoyt and Kate Langwig.

How do infections impact social relationships?

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Infectious disease dynamics are both a cause and a consequence of variation in sociality. Social interaction rates can shape parasite transmission, and conversely, parasite infection can alter social interaction rates. At the core of this feedback is a trade-off: although many social interactions yield fitness benefits, parasites impose fitness costs on infected hosts and risks to uninfected partners. Both the benefits of social interaction and the costs of infection are context-dependent, dynamic and often asymmetric within dyads. We therefore hypothesize that variation in this cost-benefit trade-off explains how and why behavioural responses to parasitism differ between individuals across stages of relationship development and across different types of social relationships (parent-offspring, pair-bond, affiliative, dominance relationships). We use these hypothetical cost-benefit trade-offs to generate testable predictions about how parasites will impact the behaviour of infected and uninfected hosts across different social relationships. We also explore the potential for acute infections to have long-term social consequences by influencing the development of social relationships.

1. Introduction

Host contact networks shape the transmission patterns of disease-causing organisms (i.e. parasites). In most epidemiological models, every host individual has an equal probability of contacting others (e.g. [1]; figure 1A). Yet interaction rates are often non-random, forming structured social networks that can influence epidemic outcomes (figure 1B). For example, disease spread can be accelerated by highly connected ‘superspreaders’ [2] and slowed by network modularity [3]. Social network structure has important implications for both short-term epidemic dynamics and the long-term coevolution of hosts and parasites [4–7], and network-based models can guide disease control strategies in humans [8] and wildlife [9,10].

Parasites are also powerful agents that can influence contact rates through the behaviour of both infected and uninfected hosts [11,12], which can feed back on transmission dynamics [13–15]. While some host behavioural changes involve parasite manipulation of hosts [11,16], other responses are by-products or adaptive responses to infection. Infected individuals often reduce their movement or social interactions following infection [12,17–19]. Uninfected hosts can either avoid infected conspecifics [20–23] or provide care [24,25]. These parasite-driven changes in contact rates can scale up to affect broader social networks and epidemiological dynamics [26–29] (figure 1C).

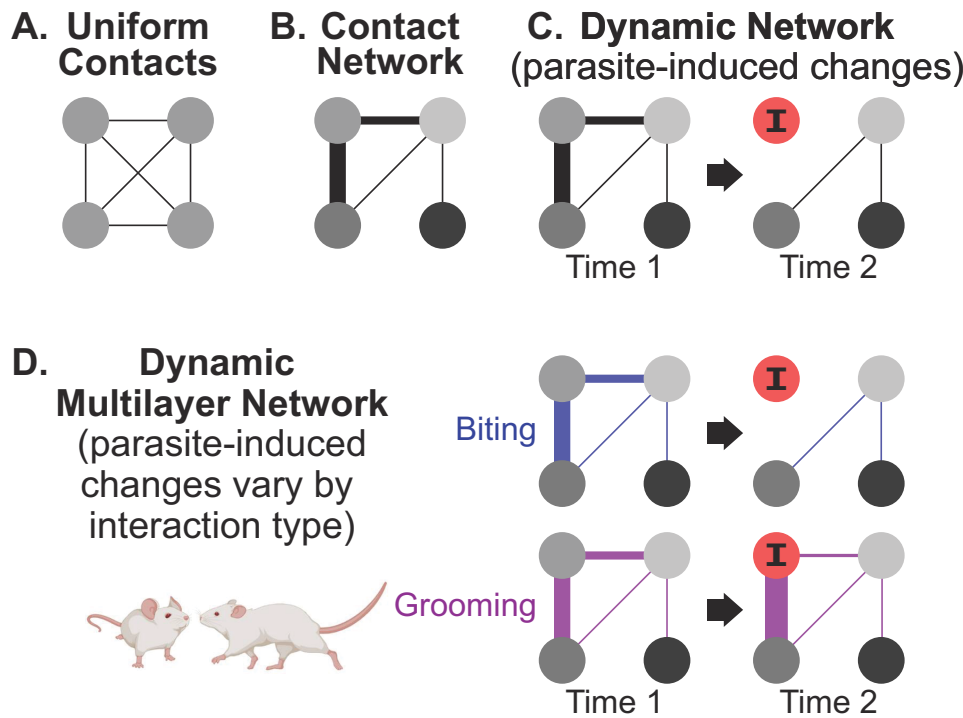


Figure 1. Increasingly complex representations of social structure in social networks can reflect how relationships change with infection. An infected individual (shown in red and marked with I) is embedded in a social environment that can be modelled using (A) uniform contacts, (B) variation in pairwise contact rates aggregated over time (i.e. a social network), (C) dynamic changes in those contact rates over time, including changes caused by infection and (D) parasite-induced behavioural changes that vary across interaction types and/or social relationships.

For example, decreases in human contact rates caused by influenza symptoms were associated with a near 75% reduction in transmission potential [30].

Shifts in contact rates in response to parasites emerge from decisions made within individual dyadic relationships. Within these pairs, uninfected and infected hosts face a trade-off between the benefits of social interactions and costs imposed by parasites, but these costs and benefits may differ between infected and uninfected partners. For uninfected hosts, social interactions can increase infection risk [31], and reducing social interactions can limit this risk [22,32,33]. In contrast, infected hosts already suffer the fitness costs of infection, arising from parasite damage and energetic costs of immunity [34]. Reducing social interactions may conserve energy for recovery [17], but infected hosts may also benefit from social interactions (e.g. caretaking) that facilitate host defences [35]. For both uninfected and infected hosts, however, foregoing social interactions also means losing any survival or reproduction-related benefits of the relationship [12,36–39].

The ratio of social benefits to infection costs is context-dependent, dynamic and asymmetric within a dyad, shaping how uninfected and infected hosts should navigate trade-offs. Many factors can influence how individuals balance social benefits and parasite costs, including perceived infection risk or severity [40,41], immunity [42], sex [43], the presence of potential mates [44,45] and predation risk [38]. Differences in which social interactions individuals prioritize or forego create variation in behavioural responses to infection [37,38,46] that scale up to context-dependent shifts in contact networks (figure 1D).

The costs and benefits of social interactions underlying trade-offs can also change as a pair of individuals forms differentiated *social relationships*—a term we use for unobserved social preferences that one animal has for affiliating with or avoiding another. Social relationships vary in strength, predictability and effect on fitness [47]. Behavioural ecologists often attempt to infer these latent social relationships from estimates of interaction or contact rates [48–51]. While disease ecologists often focus on contact rates as drivers of parasite exposure, social interactions and their underlying social relationships shape the social benefits hosts experience [47], influencing trade-offs and thereby potentially shaping how social behaviour may change in response to parasites (figure 1D).

Social relationships are also dynamic, and how relationships change over time may be both a cause and consequence of variation in how hosts respond to infection within dyads. Through repeated, mutually beneficial interactions, individuals can form stable associations that reduce conflict and increase cooperation, thereby developing a ‘stake’ in the survival of their partner due to the benefits of that partner’s continued existence, a factor called ‘fitness interdependence’ [52–54]. However, because social relationships are dynamic, the cost–benefit trade-offs between sociality and infection will likely change as social relationships form, strengthen, stabilize, weaken or dissolve. A strong social relationship may represent a significant social benefit justifying infection risk for uninfected partners or energy expenses by infected hosts, while a weak or dissolving relationship may favour avoidance or isolation. Therefore, we suggest that the current state of the relationship is a primary driver of immediate behavioural decisions in the face of disease.

Crucially, present social interactions and disruptions thereof can impact the long-term future trajectory of social relationships. Growing evidence, for example, suggests that animals preferentially associate and interact with partners who have provided greater benefits, which can change over time [55–58]. Several experiments also suggest that long-term social relationships can be impacted by relatively short-term social perturbations. For example, male prairie voles (*Microtus ochrogaster*)

separated from their pair-bonded female for four weeks lost their partner preference [59], and monk parakeets (*Myiopsitta monachus*) removed from their group for eight days lost and could not immediately regain their previous social rank [60], while female vampire bats (*Desmodus rotundus*) that were co-housed for one week were more likely to establish allogrooming relationships that persisted for months and could develop into life-saving food-sharing relationships [49,61]. These findings raise an interesting question about the potential for short-term social impacts of parasites to affect the long-term trajectory of social relationships.

Here, we address three questions: How do social relationships influence behavioural responses to parasites? Do parasites alter the formation and maintenance of relationships? And how might parasite-induced changes in social behaviour differ across relationship types? We hypothesize that variation in the parasite-cost-to-social-benefit trade-off explains how and why behavioural responses to parasitism differ across stages of relationship development and across different types of social relationships (e.g. parent-offspring, pair-bond, affiliative, dominance relationships). We then use these hypothesized cost-benefit trade-offs to generate testable predictions about how parasites impact the behaviour of infected and uninfected hosts across different social relationships. We also explore when acute infections might have long-term social consequences by affecting how social relationships develop.

(a) How do social relationships shape behavioural responses to parasites?

Parasites impose a trade-off between the benefits of social interaction and the costs of either parasite exposure [22,38,39,62] or excess energy expenditure during infection [17]. An optimal response, therefore, requires balancing the social benefits against the likelihood and costs of infection. Consequently, less beneficial social interactions and more costly parasites (e.g. those with higher transmissibility or severity) should favour greater avoidance by healthy hosts or greater social isolation by infected hosts (figure 2A,B). Conversely, hosts should maintain their investment in social relationships that are sufficiently beneficial, even when parasite costs are high. In some cases, individuals may increase their investment in relationships if there are unique benefits under infection, such as care or sanitary behaviours [63]. Similarly, growing evidence suggests that infected hosts may uniquely benefit from social interactions that improve access to resources or antipredator defence [64–66], potentially leading to asymmetries in infection costs and social benefits for sick versus healthy individuals (figure 2B).

The role of social relationships in this cost-benefit framework is well demonstrated by how responses to infection can differ by sex. Females often show a stronger aversion to infection cues than males [67]. In olive baboons (*Papio anubis*), for instance, females avoid mating with males that have visible yaws lesions, whereas males mate irrespective of infection cues [68]. While these results could represent female choice based on male immune quality [69], they can also be explained by sex-dependent cost-benefit trade-offs: for males, the additional mating opportunity may outweigh the costs of a possible infection, while for females, the same cost of possible infection may exceed the marginal reproductive benefits. Similarly, immune-challenged male zebra finches (*Taeniopygia guttata*) reduce activity in isolation but increase courtship when exposed to females [44], suggesting that the potential reproductive benefits outweigh the investment in immune defence. Sex-based differences in cost-benefit tradeoffs are also present in non-reproductive behaviours. For example, virus-infected male fruit flies (*Drosophila melanogaster*) socially distance while virus-infected females remain clumped [70]. Male social distancing likely reduces competition and energetically costly aggression, saving resources for immune defence [70]. In all these examples, the benefits of social interactions differ by sex, influencing how hosts balance social benefits and parasite costs. Next, we turn to the question of if and how parasites impact social relationships.

(b) Can parasites influence the development of long-term social relationships?

If social benefits increase in frequency and certainty over time as social relationships develop, then infections can be more disruptive to relationship formation than to relationship maintenance. Consider the simple case of two unrelated individuals developing a symmetrical, non-reproductive affiliative relationship, which is inferred from reciprocal allogrooming (or allopreening). Without an infection, allogrooming rates might increase over time during relationship formation (figure 2C) and stabilize once established (figure 2D). When the individuals first meet, there is no shared history to inform social interactions, the benefits of the social relationship are least certain, and fitness interdependence is absent. Therefore, at this stage, the cost of a potential infection should be most likely to exceed the social benefits (figure 2C,D). Yet the ability of an uninfected host to avoid a new infected partner requires detecting infection. When future social rewards are high or exclusion is costly, infected individuals may even conceal infection, as commonly observed in humans [71]. Additionally, uninfected hosts may be poor at detecting conspecifics' infections early in relationships if symptoms are progressive or if recognizing symptoms is easier with more familiar individuals [12].

In some cases, the impact of infection-induced behavioural changes on new relationships might delay or entirely prevent the establishment of a possible long-term relationship (figure 3A). This prediction is consistent with recent opportunistic observations of a *Staphylococcus* outbreak among female vampire bats captured from different wild populations and introduced in captivity [72]. Infected bats were lethargic and engaged in less allogrooming than did asymptomatic bats. The negative effects of infection on allogrooming relationships were stronger for unfamiliar pairs than for pairs with existing allogrooming relationships. This observation of different impacts of an acute infection on developing and established relationships is consistent with evidence that the immediate and anticipated benefits of social interactions early in vampire bat relationships are relatively low compared with later stages of relationship development [49,76,77]. Moreover, the effects of infection on newly introduced pairs lasted several weeks after antibiotic treatment and recovery [72], consistent with the hypothesis that infection can influence future relationship development.

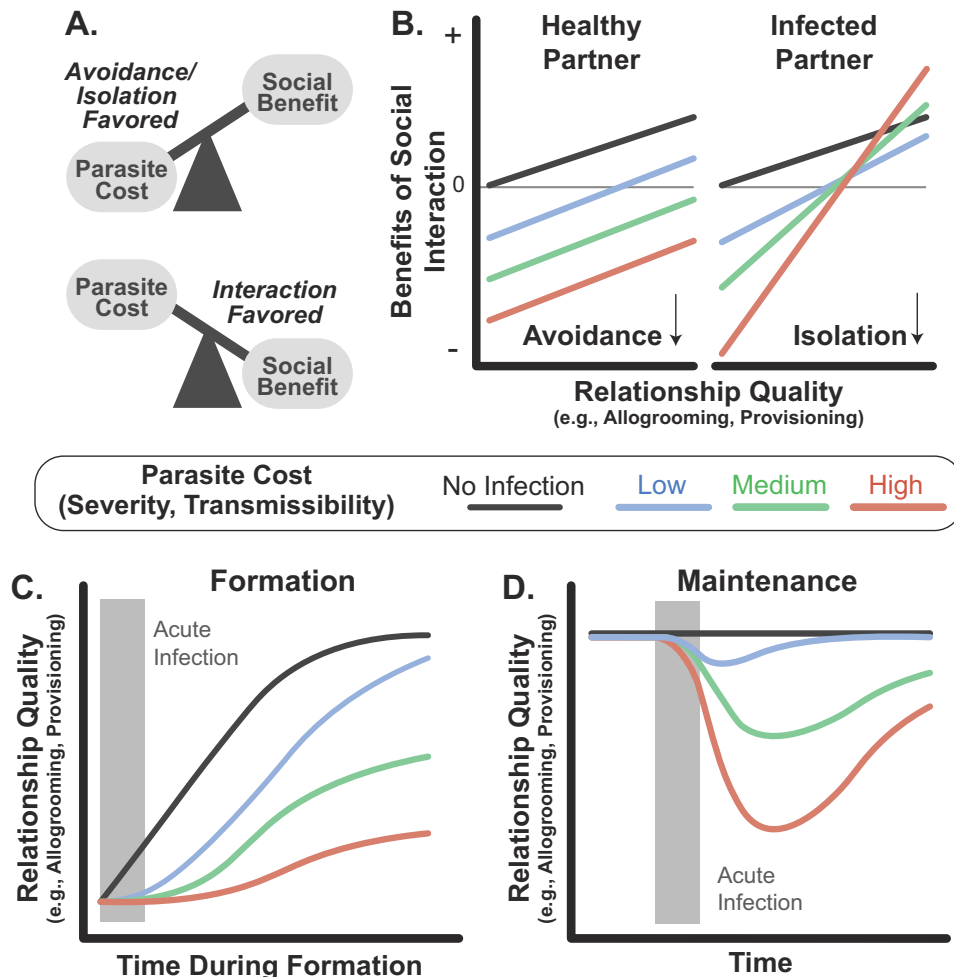


Figure 2. Trade-offs between social benefits and parasite costs shape investments in social relationships. (A) Individuals face a trade-off between the fitness benefits of social relationships and the fitness costs of infection. (B) This trade-off influences the optimal patterns of social interaction: more beneficial relationships (inferred from the rate of allogrooming, for instance) should increase the likelihood of interaction, and more severe or transmissible parasites should favour avoidance or isolation, but these effects may differ between uninfected and infected hosts. If this trade-off underlies changes in social interactions, then parasite severity and transmissibility could shape (C) relationship formation and (D) maintenance because delays in recovery will slow relationship formation or impair relationship maintenance (assuming recovery of the infected host).

Similar to affiliative relationships, infections should impact initial mate choice more than continued investment in pair bonds that enhance parental care. Parasites are known to shape mate choice [67,78]: female rock doves (*Columba livia*), for example, strongly avoided males infected with lice [79], and female mice (*Mus musculus*) exposed to parasite cues prefer familiar males over unfamiliar males, even though unfamiliar males are normally preferred [73]. However, when longer term reproductive relationships are established, particularly in pair bonds where the fitness of infected and healthy mates is strongly interdependent, infections should have weaker (or positive) effects on relationships (figure 3A). For instance, pair-bonded zebra finches remained in close proximity despite immune challenges [80], and pair-bonded prairie voles more strongly preferred their bonded mate than an unfamiliar conspecific after an immune challenge [81]. In mating systems where partnerships are continually assessed, the impact of infections can change as the balance between parasite costs and anticipated reproductive benefits shifts. For example, female western lowland gorillas (*Gorilla gorilla gorilla*) are more likely to leave ageing males, and this tendency is even stronger when males are infected with a skin disease [82]. This raises questions about whether infections could accelerate the dissolution of reproductive relationships by tipping the balance between parasite costs and benefits.

Infections should have minimal effects on parent-offspring relationships (figure 3C). Still, as offspring become increasingly independent, the effect of parasite costs on parent-offspring relationships should increase (figure 3C). This prediction is supported by observations that nematode-infected female European shag (*Phalacrocorax aristotelis*) provision their newly hatched chicks at similar rates to nematode-free mothers; however, as chicks age, nematode-infected mothers invest comparatively less in care than nematode-free mothers [74]. While this result supports the idea that parasites have different effects on early and late parent-offspring relationships, it remains unclear to what extent parasite-driven differences in parental care shape longer term parent-offspring relationships.

Competitive and dominance relationships are also influenced by social history and may therefore differentially respond to parasites over time. Most group-living animals form dominance hierarchies structured by a combination of phenotypic differences (e.g. body size), individual experiences (winner-loser effects) and pairwise interaction history [83–85]. Because dominance relationships are built from cumulative interactions, early infections may disrupt hierarchies, while established hierarchies may be robust to infections (figure 3D). Supporting this hypothesis, male mice infected with the tapeworm *Taenia*

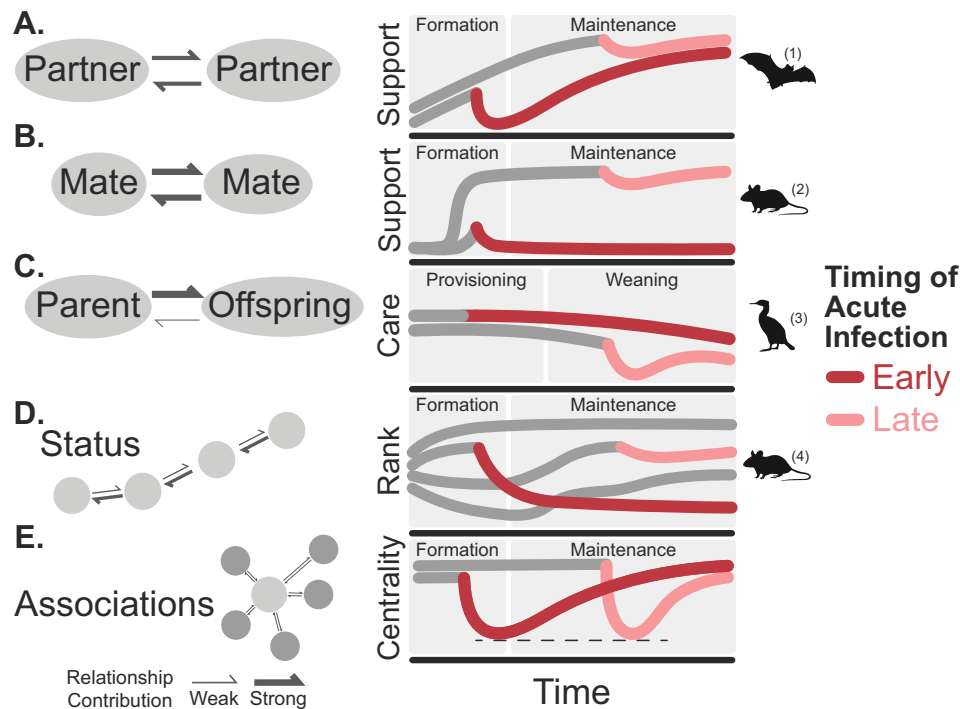


Figure 3. Acute infection effects on long-term social relationships might vary with relationship type and duration. Various social relationships, from enduring (A) same-sex affiliative relationships, (B) reproductive relationships, (C) parent–offspring relationships, (D) dominance relationships based on competition and status and (E) associations based on by-product benefits can have stages of investment or development (shaded regions). As fitness interdependence changes during relationship formation, relationships may respond differently to acute infections depending on whether they occur early (red lines) or late (pink lines). Icons represent systems where temporal changes in relationships are linked to different effects of parasites on relationships: (1) [72], (2) [73], (3) [74] and (4) [75]. Animal icons were sourced from PhyloPic.org.

crassiceps were less likely to become dominant during initial interactions, but infection had no clear effect on established hierarchies [75]. Infections may also affect competitive relationships by directly altering traits that influence competitive outcomes. In bighorn sheep (*Ovis canadensis*), for example, acute pneumonia infections, particularly during early development, can impair horn growth [86], which determines reproductive contests [87]. This suggests that early-life infections may therefore have lasting effects on competitive outcomes well into adulthood (figure 3D).

Unlike previously considered social relationships, some social associations are formed for short-term by-product benefits, like huddling for warmth or risk dilution. Because social history does not shape these by-product benefits, infections should have a similar magnitude of effect on these social associations, no matter the time of group formation (figure 3E). Instead, the trade-off is simply between the immediate by-product benefits and the parasite costs. For example, in the absence of predators, rainbow trout (*Oncorhynchus mykiss*) groups infected with an eye fluke form smaller shoals, but both uninfected and infected groups respond to predator cues by forming larger shoals [88]. This leads to the hypothesis that, for by-product associations, the duration of an association does not significantly alter parasite-driven behaviour, though to our knowledge, this remains to be explicitly tested.

2. Conclusion

The feedback between social interactions and parasitism has largely been studied with an epidemiological focus on contact rates [28,89–92]. However, different types of social relationships pose different costs and benefits for infected and uninfected hosts, likely shaping how infection alters social behaviour. And these changes might, in turn, shape long-term dynamics of relationships. This trade-off between social benefits and parasite costs can be context-dependent, dynamic and often asymmetric within dyads, leading individuals to have diverse social motivations in the face of parasitism. In turn, even short-term parasite-induced changes in social interactions might have long-term social consequences. The effect of infection on social relationships is likely to depend on the severity and timing of infection and the type of social relationship.

Testing cost–benefit trade-offs across social relationships requires integration of theory and experiments. Game theoretic models [93] and dynamic network models [7] that incorporate asymmetric costs and benefits for infected and uninfected partners could yield new insights into both behaviour and epidemic dynamics. Experiments that track changes in relationship formation in response to infection or immune challenge can measure how short-term physiological responses translate to long-term social dynamics (e.g. [80,81,94]). Operant choice tests can reveal relationship-specific motivations for infected and uninfected hosts that are difficult to detect in the wild (e.g. [81]). Although translating these approaches to natural systems is challenging because longitudinal sampling of both infection and relationship dynamics is difficult, field manipulations of infection status (e.g. anthelmintic interventions [74]) offer promising alternatives. Finally, new generative models of social networks [95,96] or the dynamics of social interactions [97] will be essential for disentangling infection effects on directed social behaviour from individual tendencies, dyadic interdependence and confounding factors such as space and time.

Animal behaviourists and disease ecologists have long appreciated that (i) social networks can shape parasite transmission [2,3,98,99], (ii) parasite-induced sickness behaviour or avoidance influences social networks [28,89–91], and (iii) social relationships are important determinants of fitness, particularly in the face of parasitism [35,100]. A critical challenge remains to more fully address the role social relationships play in behavioural and physiological responses central to epidemic dynamics. A particularly promising direction is to explore how such dynamics affect the persistence of diseases within social networks [7]. For instance, if parasites causing severe disease most strongly disrupt the formation of new relationships, the result may be the severing of ‘weak ties’ that bridge social clusters and facilitate population-wide transmission [3]. As a consequence, infection may lead to more fragmented, modular social networks and slow-burning epidemics rather than explosive outbreaks [13–15]. We suggest that modelling feedbacks between parasites and long-term social relationships is a challenging but worthwhile step towards predicting both the fitness consequences for individual hosts and the outcomes of epidemics.

Ethics. This work did not require ethical approval from a human subject or animal welfare committee.

Data accessibility. This article has no additional data.

Declaration of AI use. We have not used AI-assisted technologies in creating this article.

Authors' contributions. W.R.: conceptualization, visualization, writing—original draft; V.O.E.: supervision, writing—review and editing; G.G.C.: conceptualization, writing—original draft; S.S.: conceptualization, writing—original draft.

All authors gave final approval for publication and agreed to be held accountable for the work performed therein.

Conflict of interest declaration. We declare we have no competing interests.

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References

- Anderson RM, May RM. 1979 Population biology of infectious diseases: Part I. *Nature* **280**, 361–367. (doi:10.1038/280361a0)
- Lloyd-Smith JO, Schreiber SJ, Kopp PE, Getz WM. 2005 Superspreading and the effect of individual variation on disease emergence. *Nature* **438**, 355–359. (doi:10.1038/nature04153)
- Sah P, Leu ST, Cross PC, Hudson PJ, Bansal S. 2017 Unraveling the disease consequences and mechanisms of modular structure in animal social networks. *Proc. Natl Acad. Sci. USA* **114**, 4165–4170. (doi:10.1073/pnas.1613616114)
- Van Baalen M. 2002 Contact networks and the evolution of virulence. In *Adaptive dynamics of infectious diseases: in pursuit of virulence management*, pp. 85–103. Cambridge, UK: Cambridge University Press. (doi:10.1017/CBO9780511525728.010)
- Read JM, Keeling MJ. 2003 Disease evolution on networks: the role of contact structure. *Proc. R. Soc. B* **270**, 699–708. (doi:10.1098/rspb.2002.2305)
- Ferrari MJ, Perkins SE, Pomeroy LW, Bjørnstad ON. 2011 Pathogens, social networks, and the paradox of transmission scaling. *Interdiscip. Perspect. Infect. Dis.* **2011**, 267049. (doi:10.1155/2011/267049)
- Silk MJ, Fefferman NH. 2021 The role of social structure and dynamics in the maintenance of endemic disease. *Behav. Ecol. Sociobiol.* **75**, 1–16. (doi:10.1007/s00265-021-03055-8)
- Christley RM, Pinchbeck GL, Bowers RG, Clancy D, French NP, Bennett R, Turner J. 2005 Infection in social networks: using network analysis to identify high-risk individuals. *Am. J. Epidemiol.* **162**, 1024–1031. (doi:10.1093/aje/kwi308)
- Rushmore J, Caillaud D, Matamba L, Stumpf RM, Borgatti SP, Altizer S. 2013 Social network analysis of wild chimpanzees provides insights for predicting infectious disease risk. *J. Anim. Ecol.* **82**, 976–986. (doi:10.1111/1365-2656.12088)
- Silk MJ, Croft DP, Delahay RJ, Hodgson DJ, Boots M, Weber N, McDonald RA. 2017 Using social network measures in wildlife disease ecology, epidemiology, and management. *Bioscience* **67**, 245–257. (doi:10.1093/biosci/biw175)
- Moore J. 2002 *Parasites and the behavior of animals*. Oxford, UK: Oxford University Press.
- Stockmaier S, Stroeymeyt N, Shattuck EC, Hawley DM, Meyers LA, Bolnick DI. 2021 Infectious diseases and social distancing in nature. *Science* **371**, eabc8881. (doi:10.1126/science.abc8881)
- Ezenwa VO, Archie EA, Craft ME, Hawley DM, Martin LB, Moore J, White L. 2016 Host behaviour–parasite feedback: an essential link between animal behaviour and disease ecology. *Proc. R. Soc. B* **283**, 20153078. (doi:10.1098/rspb.2015.3078)
- Hawley DM, Gibson AK, Townsend AK, Craft ME, Stephenson JF. 2021 Bidirectional interactions between host social behaviour and parasites arise through ecological and evolutionary processes. *Parasitology* **148**, 274–288. (doi:10.1017/S0031182020002048)
- Hawley DM, Ezenwa VO. 2022 Parasites, host behavior, and their feedbacks. In *Animal behavior and parasitism* (eds VO Ezenwa, SM Altizer, R Hall), pp. 15–32. Oxford, UK: Oxford University Press. (doi:10.1093/oso/9780192895561.003.0002)
- Poulin R. 2018 Modification of host social networks by manipulative parasites. *Behaviour* **155**, 671–688. (doi:10.1163/1568539x-00003456)
- Hart BL. 1988 Biological basis of the behavior of sick animals. *Neurosci. Biobehav. Rev.* **12**, 123–137. (doi:10.1016/S0149-7634(88)80004-6)
- Dantzer R, Bluthé RM, Layé S, Bret-Dibat JL, Parnet P, Kelley KW. 1998 Cytokines and sickness behavior. *Ann. N. Y. Acad. Sci.* **840**, 586–590. (doi:10.1111/j.1749-6632.1998.tb09597.x)
- Lopes PC, French SS, Woodhams DC, Binning SA. 2021 Sickness behaviors across vertebrate taxa: proximate and ultimate mechanisms. *J. Exp. Biol.* **224**, jeb225847. (doi:10.1242/jeb.225847)
- Kiesecker JM, Skelly DK, Beard KH, Preisser E. 1999 Behavioral reduction of infection risk. *Proc. Natl Acad. Sci. USA* **96**, 9165–9168. (doi:10.1073/pnas.96.16.9165)
- Behringer DC, Butler MJ, Shields JD. 2006 Ecology: avoidance of disease by social lobsters. *Nature* **441**, 421. (doi:10.1038/441421a)
- Curtis VA. 2014 Infection-avoidance behaviour in humans and other animals. *Trends Immunol.* **35**, 457–464. (doi:10.1016/j.it.2014.08.006)

23. Poirotte C, Massol F, Herbert A, Willaume E, Bomo PM, Kappeler PM, Charpentier MJE. 2017 Mandrills use olfaction to socially avoid parasitized conspecifics. *Sci. Adv.* **3**, e1601721. (doi:10.1126/sciadv.1601721)
24. Cremer S, Pull CD, Fürst MA. 2018 Social immunity: emergence and evolution of colony-level disease protection. *Annu. Rev. Entomol.* **63**, 105–123. (doi:10.1146/annurev-ento-020117-043110)
25. Stockmaier S, Bolnick DI, Page RA, Carter GG. 2020 Sickness effects on social interactions depend on the type of behaviour and relationship. *J. Anim. Ecol.* **89**, 1387–1394. (doi:10.1111/1365-2656.13193)
26. Christensen C, Albert I, Grenfell B, Albert R. 2010 Disease dynamics in a dynamic social network. *Physica A* **389**, 2663–2674. (doi:10.1016/j.physa.2010.02.034)
27. Van Segbroeck S, Santos FC, Pacheco JM. 2010 Adaptive contact networks change effective disease infectiousness and dynamics. *PLoS Comput. Biol.* **6**, e1000895. (doi:10.1371/journal.pcbi.1000895)
28. Stroeymeyt N, Grasse AV, Crespi A, Mersch DP, Cremer S, Keller L. 2018 Social network plasticity decreases disease transmission in a eusocial insect. *Science* **362**, 941–945. (doi:10.1126/science.aat4793)
29. White LA, Forester JD, Craft ME. 2018 Covariation between the physiological and behavioral components of pathogen transmission: host heterogeneity determines epidemic outcomes. *Oikos* **127**, 538–552. (doi:10.1111/oik.04527)
30. Van Kerckhove K, Hens N, Edmunds WJ, Eames KTD. 2013 The impact of illness on social networks: implications for transmission and control of influenza. *Am. J. Epidemiol.* **178**, 1655–1662. (doi:10.1093/aje/kwt196)
31. Alexander RD. 1974 The evolution of social behavior. *Annu. Rev. Ecol. Syst.* **5**, 325–383. (doi:10.1146/annurev.es.05.110174.001545)
32. Schaller M, Park JH. 2011 The behavioral immune system (and why it matters). *Curr. Dir. Psychol. Sci.* **20**, 99–103. (doi:10.1177/0963721411402596)
33. Kramer P, Bressan P. 2021 Infection threat shapes our social instincts. *Behav. Ecol. Sociobiol.* **75**, 47. (doi:10.1007/s00265-021-02975-9)
34. Lochmiller RL, Deerenberg C. 2000 Trade-offs in evolutionary immunology: just what is the cost of immunity? *Oikos* **88**, 87–98. (doi:10.1034/j.1600-0706.2000.880110.x)
35. Kappeler PM, Cremer S, Nunn CL. 2015 Sociality and health: impacts of sociality on disease susceptibility and transmission in animal and human societies. *Phil. Trans. R. Soc. B* **370**. (doi:10.1098/rstb.2014.0116)
36. Lopes PC. 2014 When is it socially acceptable to feel sick? *Proc. R. Soc. B* **281**, 20140218. (doi:10.1098/rspb.2014.0218)
37. Amoroso CR, Antonovics J. 2020 Evolution of behavioural resistance in host-pathogen systems. *Biol. Lett.* **16**, 20200508. (doi:10.1098/rsbl.2020.0508)
38. Poirotte C, Charpentier MJE. 2022 Inter-individual variation in parasite avoidance behaviors and its epidemiological, ecological, and evolutionary consequences. In *Animal behavior and parasitism*, pp. 257–270. Oxford, UK: Oxford University Press. (doi:10.1093/oso/9780192895561.003.0015)
39. Stockmaier S, Ulrich Y, Albery GF, Cremer S, Lopes PC. 2023 Behavioural defences against parasites across host social structures. *Funct. Ecol.* **37**, 809–820. (doi:10.1111/1365-2435.14310)
40. Kavaliers M, Choleris E. 2018 The role of social cognition in parasite and pathogen avoidance. *Phil. Trans. R. Soc. B* **373**, 20170206. (doi:10.1098/rstb.2017.0206)
41. Stephenson JF, Perkins SE, Cable J. 2018 Transmission risk predicts avoidance of infected conspecifics in Trinidadian guppies. *J. Anim. Ecol.* **87**, 1525–1533. (doi:10.1111/1365-2656.12885)
42. Zylberberg M, Klasing KC, Hahn TP. 2013 House finches (*Carpodacus mexicanus*) balance investment in behavioural and immunological defences against pathogens. *Biol. Lett.* **9**, 20120856. (doi:10.1098/rsbl.2012.0856)
43. Yirmiya R, Avitsur R, Donchin O, Cohen E. 1995 Interleukin-1 inhibits sexual behavior in female but not in male rats. *Brain Behav. Immun.* **9**, 220–233. (doi:10.1006/brbi.1995.1021)
44. Lopes PC, Chan H, Demathieu S, González-Gómez PL, Wingfield JC, Bentley GE. 2013 The impact of exposure to a novel female on symptoms of infection and on the reproductive axis. *Neuroimmunomodulation* **20**, 348–360. (doi:10.1159/000353779)
45. Lopes PC, Faber-Hammond JJ, Siemonsma C, Patel S, Renn SCP. 2023 The social environment alters neural responses to a lipopolysaccharide challenge. *Brain Behav. Immun.* **110**, 162–174. (doi:10.1016/j.bbi.2023.03.004)
46. Ezenwa VO, Ghai RR, McKay AF, Williams AE. 2016 Group living and pathogen infection revisited. *Curr. Opin. Behav. Sci.* **12**, 66–72. (doi:10.1016/j.cobeha.2016.09.006)
47. De Moor D, Brent L. 2025 Quality, quantity, and the adaptive function of social relationships. *EcoEvoRxiv*. (doi:10.32942/X2MM15)
48. Hinde RA. 1976 Interactions, relationships and social structure. *Man* **11**, 1–17.
49. Carter GG, Farine DR, Crisp RJ, Vrtilek JK, Ripperger SP, Page RA. 2020 Development of new food-sharing relationships in vampire bats. *Curr. Biol.* **30**, 1275–1279. (doi:10.1016/j.cub.2020.01.055)
50. De Moor D, Hart JDA, Franks DW, Brent LJN, Silk MJ, Brask JB. 2025 Latent layers in social networks and their implications for comparative analyses. *Behav. Ecol.* **36**, araf113. (doi:10.1093/beheco/araf113)
51. Kawam B, Ostner J, McElreath R, Schülke O, Redhead D. 2025 A causal framework for the drivers of animal social network structure. *PLoS Comput. Biol.* **21**, e1013370. (doi:10.1101/2024.06.26.600748)
52. Roberts G. 2005 Cooperation through interdependence. *Anim. Behav.* **70**, 901–908. (doi:10.1016/j.anbehav.2005.02.006)
53. Bshary R, Zuberbühler K, van Schaik CP. 2016 Why mutual helping in most natural systems is neither conflict-free nor based on maximal conflict. *Phil. Trans. R. Soc. B* **371**, 20150091. (doi:10.1098/rstb.2015.0091)
54. Carter GG. 2024 Reciprocity versus pseudo-reciprocity: a false dichotomy. *Ethology* **130**, e13431. (doi:10.1111/eth.13431)
55. Fruteau C, Voelkl B, van Damme E, Noë R. 2009 Supply and demand determine the market value of food providers in wild vervet monkeys. *Proc. Natl Acad. Sci. USA* **106**, 12007–12012. (doi:10.1073/pnas.0812280106)
56. Borgeaud C, Bshary R. 2015 Wild vervet monkeys trade tolerance and specific coalitionary support for grooming in experimentally induced conflicts. *Curr. Biol.* **25**, 3011–3016. (doi:10.1016/j.cub.2015.10.016)
57. Kern JM, Radford AN. 2018 Experimental evidence for delayed contingent cooperation among wild dwarf mongooses. *Proc. Natl Acad. Sci. USA* **115**, 6255–6260. (doi:10.1073/pnas.1801000115)
58. Kings M, Arbon JJ, McIvor GE, Whitaker M, Radford AN, Lerner J, Thornton A. 2023 Wild jackdaws can selectively adjust their social associations while preserving valuable long-term relationships. *Nat. Commun.* **14**, 5103. (doi:10.1038/s41467-023-40808-7)
59. Sun P, Smith AS, Lei K, Liu Y, Wang Z. 2014 Breaking bonds in male prairie vole: long-term effects on emotional and social behavior, physiology, and neurochemistry. *Behav. Brain Res.* **265**, 22–31. (doi:10.1016/j.bbr.2014.02.016)
60. van der Marel A, Francis X, O'Connell CL, Estien CO, Carminito C, Moore VD, Lormand N, Kluever BM, Hobson EA. 2023 Perturbations highlight importance of social history in parakeet rank dynamics. *Behav. Ecol.* **34**, 457–467. (doi:10.1093/beheco/arad015)

61. Razik I, Brown BKG, Carter GG. 2022 Forced proximity promotes the formation of enduring cooperative relationships in vampire bats. *Biol. Lett.* **18**, 20220056. (doi:10.1101/2022.01.30.478382)
62. Lopes PC. 2022 Anticipating infection: how parasitism risk changes animal physiology. *Funct. Ecol.* **37**, 821–830. (doi:10.1111/1365-2435.14155)
63. Konrad M, Pull CD, Metzler S, Seif K, Naderlinger E, Grasse AV, Cremer S. 2018 Ants avoid superinfections by performing risk-adjusted sanitary care. *Proc. Natl Acad. Sci. USA* **115**, 2782–2787. (doi:10.1073/pnas.1713501115)
64. Almberg ES, Cross PC, Dobson AP, Smith DW, Metz MC, Stahler DR, Hudson PJ. 2015 Social living mitigates the costs of a chronic illness in a cooperative carnivore. *Ecol. Lett.* **18**, 660–667. (doi:10.1111/ele.12444)
65. Ezenwa VO, Worsley-Tonks KEL. 2018 Social living simultaneously increases infection risk and decreases the cost of infection. *Proc. R. Soc. B* **285**, 20182142. (doi:10.1098/rspb.2018.2142)
66. Langager MM, Arneson AG, Hawley DM. 2025 In sickness and in health: group-living augments behavioural responses to food and predation risk for sick house finches (*Haemorrhous mexicanus*). *R. Soc. Open Sci.* **12**, 251715. (doi:10.1098/rsos.251715)
67. Beltran-Bech S, Richard FJ. 2014 Impact of infection on mate choice. *Anim. Behav.* **90**, 159–170. (doi:10.1016/j.anbehav.2014.01.026)
68. Paciência FMD, Rushmore J, Chuma IS, Lipende IF, Caillaud D, Knauf S, Zinner D. 2019 Mating avoidance in female olive baboons (*Papio anubis*) infected by *Treponema pallidum*. *Sci. Adv.* **5**, eaaw9724. (doi:10.1126/sciadv.aaw9724)
69. Hamilton WD, Zuk M. 1982 Heritable true fitness and bright birds: a role for parasites? *Science* **218**, 384–387. (doi:10.1126/science.7123238)
70. Siva-Jothy JA, Vale PF. 2019 Viral infection causes sex-specific changes in fruit fly social aggregation behaviour. *Biol. Lett.* **15**, 20190344. (doi:10.1098/rsbl.2019.0344)
71. Merrell WN, Choi S, Ackerman JM. 2024 When and why people conceal infectious disease. *Psychol. Sci.* **35**, 215–225. (doi:10.1177/09567976231221990)
72. Razik I, Stockmaier S, Abou-Elias M, Carter GG. 2024 Opportunistic evidence of the impact of bacterial infections on social integration in vampire bats. *bioRxiv*. (doi:10.1101/2023.04.17.537180)
73. Kavaliers M, Colwell DD, Cloutier CJ, Ossenkopp KP, Choleris E. 2014 Pathogen threat and unfamiliar males rapidly bias the social responses of female mice. *Anim. Behav.* **97**, 105–111. (doi:10.1016/j.anbehav.2014.09.006)
74. Reed TE, Daunt F, Hall ME, Phillips RA, Wanless S, Cunningham EJA. 2008 Parasite treatment affects maternal investment in sons. *Science* **321**, 1681–1682. (doi:10.1126/science.1159466)
75. Gourbal BEF, Lacroix A, Gabrion C. 2002 Behavioural dominance and *Taenia crassiceps* parasitism in BALB/c male mice. *Parasitol. Res.* **88**, 912–917. (doi:10.1007/s00436-002-0691-7)
76. Wilkinson GS. 1984 Reciprocal food sharing in the vampire bat. *Nature* **308**, 181–184. (doi:10.1038/308181a0)
77. Carter GG, Wilkinson GS. 2013 Food sharing in vampire bats: reciprocal help predicts donations more than relatedness or harassment. *Proc. R. Soc. B* **280**, 20122573. (doi:10.1098/rspb.2012.2573)
78. Andersson M, Iwasa Y. 1994 *Sexual selection*, pp. 53–58. Princeton, NJ: Princeton University Press. (doi:10.1016/0169-5347(96)81042-1)
79. Clayton DH. 1990 Mate choice in experimentally parasitized rock doves: lousy males lose. *Am. Zool.* **30**, 251–262. (doi:10.1093/icb/30.2.251)
80. Love AC, Anthony AC, Nash A, Campos-Melara A, Kodali J, DuRant SE. 2023 Simulated infection alters the behavior of pair bonded songbirds and their healthy neighbors. *Behav. Ecol.* **34**, 251–260. (doi:10.1093/beheco/araa120)
81. Young GK, Chernyak D, Naik GA, Song SE, Beery AK. 2024 Prairie voles seek social contact with peer companions during immune challenge. *Horm. Behav.* **166**, 105653. (doi:10.1016/j.yhbeh.2024.105653)
82. Baudouin A *et al.* 2019 Disease avoidance, and breeding group age and size condition the dispersal patterns of western lowland gorilla females. *Ecology* **100**, e02786. (doi:10.1002/ecy.2786)
83. Chase ID, Tovey C, Spangler-Martin D, Manfredonia M. 2002 Individual differences versus social dynamics in the formation of animal dominance hierarchies. *Proc. Natl Acad. Sci. USA* **99**, 5744–5749. (doi:10.1073/pnas.082104199)
84. Hsu Y, Earley RL, Wolf LL. 2006 Modulation of aggressive behaviour by fighting experience: mechanisms and contest outcomes. *Biol. Rev.* **81**, 33–74. (doi:10.1017/S146479310500686X)
85. Tibbetts EA, Pardo-Sanchez J, Weise C. 2022 The establishment and maintenance of dominance hierarchies. *Phil. Trans. R. Soc. B* **377**, 20200450. (doi:10.1098/rstb.2020.0450)
86. Martin AM, Hogg JT, Manlove KR, LaSharr TN, Shannon JM, McWhirter DE, Miyasaki H, Monteith KL, Cross PC. 2022 Disease and secondary sexual traits: effects of pneumonia on horn size of bighorn sheep. *J. Wildl. Manage.* **86**, e22154. (doi:10.1002/jwmg.22154)
87. Coltman DW, Festa-Bianchet M, Jorgenson JT, Strobeck C. 2002 Age-dependent sexual selection in bighorn rams. *Proc. R. Soc. B* **269**, 165–172. (doi:10.1098/rspb.2001.1851)
88. Seppälä O, Karvonen A, Valtonen ET. 2008 Shoaling behaviour of fish under parasitism and predation risk. *Anim. Behav.* **75**, 145–150. (doi:10.1016/j.anbehav.2007.04.022)
89. Croft DP, Edenbrow M, Darden SK, Ramnarine IW, van Oosterhout C, Cable J. 2011 Effect of gyrodactylid ectoparasites on host behaviour and social network structure in guppies *Poecilia reticulata*. *Behav. Ecol. Sociobiol.* **65**, 2219–2227. (doi:10.1007/s00265-011-1230-2)
90. Lopes PC, Block P, König B. 2016 Infection-induced behavioural changes reduce connectivity and the potential for disease spread in wild mice contact networks. *Sci. Rep.* **6**, 31790. (doi:10.1038/srep31790)
91. Hamilton DG, Jones ME, Cameron EZ, Kerlin DH, McCallum H, Storfer A, Hohenlohe PA, Hamede RK. 2020 Infectious disease and sickness behaviour: tumour progression affects interaction patterns and social network structure in wild Tasmanian devils. *Proc. R. Soc. B* **287**, 20202454. (doi:10.1098/rspb.2020.2454)
92. Ripperger SP, Stockmaier S, Carter GG. 2020 Tracking sickness effects on social encounters via continuous proximity sensing in wild vampire bats. *Behav. Ecol.* **31**, 1296–1302. (doi:10.1093/beheco/araa111)
93. Leimar O, Bshary R. 2024 Social bond dynamics and the evolution of helping. *Proc. Natl Acad. Sci. USA* **121**, e2317736121. (doi:10.1073/pnas.2317736121)
94. Stockmaier S, Bolnick DI, Page RA, Josic D, Carter GG. 2020 Immune-challenged vampire bats produce fewer contact calls. *Biol. Lett.* **16**, 20200272. (doi:10.1098/rsbl.2020.0272)
95. Ross CT, McElreath R, Redhead D. 2024 Modelling animal network data in R using STRAND. *J. Anim. Ecol.* **93**, 254–266. (doi:10.1111/1365-2656.14021)
96. Ross CT, Kajokaite K, Pinkney S, Sosa S. 2025 Bayesian multiplex network models in R using STRAND: methods for biologists and social scientists. *R. Soc. Open Sci.* **12**, 250555. (doi:10.1098/rsos.250555)
97. Kawam B, McElreath R, Ostner J, Redhead D, Schülke O. 2025 Inferring the causes of animal social network structure from time-series data. *bioRxiv*. (doi:10.1101/2025.11.08.687336)
98. Gear DA, Luong LT, Hudson PJ. 2013 Network transmission inference: host behavior and parasite life cycle make social networks meaningful in disease ecology. *Ecol. Appl.* **23**, 1906–1914. (doi:10.1890/13-0907.1)

99. Briard L, Ezenwa VO. 2021 Parasitism and host social behaviour: a meta-analysis of insights derived from social network analysis. *Anim. Behav.* **172**, 171–182. (doi:[10.1016/j.anbehav.2020.11.010](https://doi.org/10.1016/j.anbehav.2020.11.010))
100. Snyder-Mackler N *et al.* 2020 Social determinants of health and survival in humans and other animals. *Science* **368**, eaax9553. (doi:[10.1126/science.aax9553](https://doi.org/10.1126/science.aax9553))