UNSOLVED MYSTERY

Why Do We Feel Sick When Infected—Can Altruism Play a Role?

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Abstract

When we contract an infection, we typically feel sick and behave accordingly. Symptoms of sickness behavior (SB) include anorexia, hypersomnia, depression, and reduced social interactions. SB affects species spanning from arthropods to vertebrates, is triggered non-specifically by viruses, bacteria, and parasites, and is orchestrated by a complex network of cytokines and neuroendocrine pathways; clearly, it has been naturally selected. Nonetheless, SB seems evolutionarily costly: it promotes starvation and predation and reduces reproductive opportunities. How could SB persist? Former explanations focused on individual fitness, invoking improved resistance to pathogens. Could prevention of disease transmission, propagating in populations through kin selection, also contribute to SB?

Sickness Syndrome and Sickness Behavior

Sickness syndrome is the generalized response of the host to infections. Its classical physiological signs include fever and anemia, but it also includes psychological symptoms—collectively termed "sickness behavior" (SB) [1–3]. These symptoms, familiar to anyone who has been sick, include fatigue, depression, irritability, discomfort, pain, nausea, and loss of interest in food, drink, social interactions, and sex. In animals, such changes can be quantified based on behavioral responses (Fig 1) [4,5].

A common misconception is that pathogens directly produce these behavioral symptoms, but in fact SB is orchestrated by the host's immune and neuroendocrine systems; mammals have evolved several parallel pathways to alert the brain of inflammation and trigger symptomatic behaviors (Fig 1) [4,5].

Although the specificities may vary, SB is widespread with respect to both pathogens and hosts: diverse pathogens, including viruses, bacteria, and protozoa [1], can trigger it, and equivalent behavioral responses characterize several vertebrate classes [1,2,7] as well as arthropods [8,9]. However, when closely examined, some genera exhibit significant variation in the extent of SB [10], which to date remains unexplained.

The Mystery—Why Do We Feel Sick?

Since SB is a conserved phenomenon that is mediated by complex immunological and neuroendocrine pathways, it clearly must have evolutionary benefits. Still, in the last 25 years, much
Fig 1. Information regarding inflammation is communicated to the brain through parallel neural and circulatory routes [4,5]. Leukocytes, such as dendritic cells (DCs) and macrophages, sense microbes through pathogen-recognition receptors such as toll-like receptors (TLRs) and NOD-like receptors (NLRs) and then release inflammatory cytokines such as interleukin-1 beta (IL-1β), IL-6, and tumor necrosis factor alpha (TNF-α). In the neural route, cytokines trigger activity in vagal afferents that innervate nuclei in the brain stem such as the nucleus of the solitary tract (NTS). These in turn relay the signal to various nuclei in the hypothalamus, thalamus, and amygdala [4]. In the circulatory route, microbial ligands and cytokines travel through the blood to reach the meninges, choroid plexus, and circumventricular organs (pink) where they can enter the brain. More recent data indicate that such ligands can also activate the epithelium in areas with an intact blood–brain barrier, causing it to synthesize various prostaglandins and release them into nuclei involved in specific behaviors [6].

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effort has been directed at understanding the proximate reasons for SB [3], but its ultimate causation—the reasons SB has evolved in the first place—attracted relatively little attention.

Unlike physiological symptoms of sickness, such as fever and hypoferremia, which likely boost resistance to pathogens (Box 1), behavioral symptoms remain poorly explained. Clearly, all of these symptoms impose significant costs to host fitness (Fig 2) [11,12]. Anorexia and adipsia increase the risk of starvation, loss of essential nutrients, and dehydration, particularly in the context of fever. Lethargy can lead to predation by slowing down prey and singling it out for predators [13,14]. Social disinterest decreases parental care [15,16], limits mating opportunities [17], and, together with fatigue, can lead to loss of territory and social status [7,18]. For SB to evolve, these costs must be offset by benefits—what can these benefits be?

**Box 1. Fever and Hypoferremia: Physiological Manifestations of Sickness Syndrome**

Physiological responses to sickness are initiated by the immune system and propagated mainly by the brain and liver. Many of these are believed to benefit host resistance to infections, and two, fever and anemia, have been linked to SB [1].

Fever is widely believed to improve survival following infection [19,20] by directly inhibiting the growth of various pathogens and by enhancing immune function (e.g., bacterial clearance, T cell proliferation, and neutrophil activation) [21]. The benefits of hyperthermia has been most convincingly demonstrated in ectoderms such as reptiles and fish, in which deliberate exposure to higher environmental temperatures improved survival [19]. Correspondingly, in rabbits, mice, and chicks, antipyretic drugs repeatedly increased mortality rates from bacterial [22] and viral [23] infections. The evidence in humans is less conclusive, as large-scale blinded trials have not been performed [24,25]. Nonetheless, several small randomized trials have reported that antipyrogenic agents delayed recovery from infections such as malaria [26–28] and chicken pox [29]. Consequently, it has been estimated that routinely treating influenza patients with antipyretics causes at least 700 extra deaths annually in the United States alone [20].

Another physiological component of sickness syndrome is anemia, which is a byproduct of “hypoferremia of infection.” Hypoferremia is a well-regulated process intended to deprive pathogens of the iron essential for their growth [30,31]. It affects several classes of pathogens, including many bacteria, some viruses, and several protozoa. Freely available iron can diminish normal resistance to bacteria in several diseases, and iron overload increased infection rates of pathogens such as tuberculosis, malaria, and brucellosis [31,32].

Infection elicits hypoferremia as part of the hepatic acute phase response [31]. Inflammatory cytokines such as IL–6, IL–22, and type-I interferons trigger the production of the peptide hormone hepcidin in the liver. Hepcidin then binds and internalizes the iron exporter protein ferroportin. As a result, macrophages trap the iron recycled from erythrocytes, and enterocytes stop transferring dietary iron to the circulation, rapidly reducing plasma iron.

**Can SB Improve Host Resistance?**

The concept that SB is a coordinated and adaptive response to infections has been established since the mid-1980s. Several comprehensive reviews have covered the historical development of this concept and considered various hypotheses regarding the adaptive role of SB [2–4,14,33].

Early findings suggesting that SB directly benefits the host examined anorexia. In a well-controlled study from 1979, Murray and Murray infected mice with *Listeria monocytogenes* and
Fig 2. The costs of SB to direct fitness. Behavioral (pink) and physiological (green) symptoms of SB can, either directly or indirectly, lead to maladaptive consequences (orange) that reduce individual fitness.

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force-fed them to compensate for the resultant anorexia [34]. The treated mice succumbed to the infection at high rates. Unlike the adaptive effects of fever, this remained a largely isolated study, and contesting theories still debate whether anorexia boosts resistance to pathogens and how it might do so [35]. Suggestions included deliberate restriction of nutritional elements, avoiding potentially contaminated food, and a decrease in risky foraging while weak [35]. Newly established routes linking nutrition, intestinal microbiota, and immunity [36] can now also be considered.

In 1988, a seminal paper by Benjamin L. Hart was the first to suggest that SB in its entirety is a coordinated response benefitting the host [1]. Realizing that fever and hypoferremia directly promote host defense (Box 1), Hart suggested that SB is primarily intended to serve these physiological adaptations. Specifically, he proposed that SB evolved to conserve energy needed to sustain metabolically demanding fever. Thus, immobility, lethargy, and reduced motivation to obtain food and drink could have developed to minimize muscle work and exposure to the cold. Anorexia, on the other hand, would promote hypoferremia by reducing iron intake. Other behaviors were viewed as subordinate to the primary ones that conserve energy and reduce iron. Reduced grooming, for example, could preserve fluids in the context of adipisia, whereas decreased foraging would protect a weak animal from predators.

Hart’s hypothesis remained the dominant theory in the field [2,37–40], as it parsimoniously explains a large range of symptoms. Since it was proposed, though, accumulated evidence has exposed some gaps in the hypothesis; it is now time to reassess it.

Conserving energy to maintain fever is central to Hart’s hypothesis. SB is definitely associated with reduced motivation for action—and therefore with less energy expenditure. However, in many cases, fever and SB are decoupled, the one arising without the other [10]. In humans, for instance, malaise and fatigue often characterize mild infections that do not elicit fever. More importantly, several aspects of SB can actually tip the energy balance in the wrong direction. Confinement to nests and dens does not always conserve heat. In warmer climates, dens are cooler than the outside environment and mobility increases body temperature, yet desert animals still remain inside [10]. Another counterproductive symptom is reduced grooming. When mammals and birds stop grooming, their fur and plumage gradually lose their insulating efficiency, requiring more energy to maintain fever [41,42].

The most counterintuitive symptom is anorexia, which, as Hart acknowledged, deprives sick animals of calories needed to fuel fever (especially in migratory animals that cannot reduce energy expenditure by retiring to protected environments). Recognizing this caveat, Hart suggested instead that anorexia evolved to reduce iron consumption, consequently assisting another important antimicrobial response—hypoferremia. It seems unlikely, though, that evolution would favor an indiscriminate reduction in food intake just to decrease iron consumption. Herbivores, for instance, can vary their diet to suit nutritional needs [43], so they could instead avoid only iron-rich foods or ingest clayey soil to interfere with iron absorption [44].

More importantly, physiologists have since gained much mechanistic insight into hypoferremia, rendering this notion less likely. Dietary iron absorption is dwarfed by the total iron reserves in the human body and the amount recycled through erythropoiesis [45]. Anorexia, therefore, can only mediate slow-acting changes in plasma iron [46]. In contrast, inflammatory agents such as lipopolysaccharide (LPS) can halve plasma iron within a few hours [47]. The direct mechanism through which infection elicits hypoferremia (Box 1) was only discovered 15 years ago [31] and involves the rapid production of hepcidin in the liver. This efficient mechanism obviates anorexia when infection requires the host to rapidly reduce plasma iron.

Sensing that Hart’s explanation cannot account for all the symptoms of SB, several complementary theories have since been proposed. Watkins and Maier [48] stressed the importance of allodynia and hyperalgesia (reduced threshold and increased intensity of pain) in SB. They
proposed that these symptoms, together with the reduced activity SB introduces, are intended to protect sensitive organs and tissues from further damage. Medzitov et al. [49] maintained that SB chiefly promotes tolerance towards parasites, rather than their clearance, although the details of this interaction remained unclear. All these theories focus on direct benefits that infected individuals may derive from SB; they disregard the indirect effects SB may have at the group level.

Overall, the evidence that all the symptoms of SB directly improve host resistance to infection remains incomplete, and after several decades of research in this field, writers still debate whether and how symptoms of SB benefit hosts [3,14,33,35,50]. What, then, could a complementary evolutionary explanation be?

Could Kin Selection Drive the Evolution of SB?

If gains to direct fitness cannot fully explain SB, perhaps inclusive fitness could come into play. We propose that reduced transmission of infectious disease among related individuals contributed to the evolution of SB. Although the idea that SB reduces transmission has been alluded to before [3,20,51,52], it was never recognized as a major organizing principle for SB in vertebrates. We name this theory "the Eyam hypothesis" after the English mining community that isolated itself to contain an outbreak of bubonic plague in 1666. Three-quarters of the villagers reportedly died, but the surrounding communities were saved [53].

The Eyam hypothesis relies on three premises:

Premise 1: SB Reduces Direct and Indirect Contacts between Infected Individuals and Their Conspecifics

Strikingly, most of the symptoms that constitute SB share a common denominator: they restrict contacts between sick individuals and their social groups (Fig 3). Symptoms of sickness achieve this feat using three containment strategies:

Containment Strategy #1: Restricting Physical Contacts

It is self-evident that salient symptoms of SB, such as social disinterest, depression, hyperalgesia, fatigue, and hypersomnia, reduce the mobility and social activity of infected individuals, limiting their contact with conspecifics. Likewise, sexual disinterest suppresses courtship and mating behaviors, whereas reduced parental care entails by definition less interaction with offspring. The contribution of anorexia and adipsia may be less apparent; by suppressing the motivation to eat and drink, they reduce the urge to travel in search of food and water, share meals with group members, and gather at water sources.

Self-imposed isolation may account for the folk observation that terminally ill dogs leave their owners to die alone. Similar behavior has been recorded in the wild among badgers, which, when infected with bovine tuberculosis, separated from their clan and settled in individual setts, where they died [54].

Tellingly, the opposite effect is observed when pathogens manipulate host behavior to their benefit. In such diseases, infected hosts become hyperactive and interact more with potential hosts: for example, rabid dogs become fearless and bite, and rodents infected with *Toxoplasma gondii* lose their fear of cats (the definitive hosts) [55].

Containment Strategy #2: Limiting Environmental Contamination

On top of restricting direct contacts, SB can also limit indirect contacts between conspecifics by reducing microbial contamination of shared resources: ground, food and water (Fig 3).
Symptoms such as hypersomnia, fatigue, and depression restrict the animal’s radius of activity, limiting environmental contamination to its immediate surroundings. Social and sexual disinterest, as well as anorexia and adipsia, further reduce the drive of animals to travel farther afield.

Anorexia and adipsia seem paramount in that respect as they prevent sick animals from contaminating shared food and water resources. Contamination of pastures (for herbivores) or carcasses (for carnivores) and contamination of water holes are undoubtedly major routes for oral and fecal-to-oral transmission in the wild. Finally, anorexia and adipsia also reduce defecation, diarrhea, and vomiting, which are the major means of spreading for enteric pathogens.

Fig 3. The benefits of SB to indirect fitness. Symptoms of SB (pink) can suppress (red connectors) or promote (green arrows) several mediating behaviors (yellow), consequently reducing pathogen transmission through several routes (blue).

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Containment Strategy #3: Advertising Infection to Conspecifics

Whereas strategies #1 and #2 involve self-imposed restrictions, SB can also act by provoking responses from conspecifics. In many species, group members can detect infected individuals through visual, olfactory, and chemical cues [56–59], distance themselves, and stop interacting with them [60]. Such signaling has been demonstrated most convincingly in eusocial insects in which chemical communication is used to coordinate social immunity (Box 2).

Box 2. The Case for Social Immunity in Eusocial Insects

Eusocial insects—social bees and wasps, ants, and termites—form colonies dubbed “superorganisms.” These contain few breeding individuals and many closely related sterile workers. Workers are dispensable, care collectively for brood, and are genetically investing in their siblings and parents. This situation encourages cooperation and altruism. Colonies of eusocial insects are ideal settings for the spread of pathogens, as their inhabitants live at high density, constantly touch one another, and exchange food orally. Low genetic diversity may pose an additional risk, as more individuals are susceptible to the same pathogens. Theoretically, these factors make eusocial insects optimal candidates to develop SB.

Empirically, it has long been recognized that eusocial insects exhibit social immunity, collective behaviors that promote parasite resistance [61] and limit contagious interactions among group members [62]. Many of these behaviors resemble SB in vertebrates, whereas some are idiosyncratic adaptations to the situation in insect colonies.

Specifically, among several species of ants, individuals that had been experimentally treated with live pathogens or pathogen-associated molecules (such as LPS) are less social [8], avoid contacting brood [8,63], stop transferring food to nest mates (trophallaxis) [9], become less motile [9], decrease allogrooming of nest mates [64], and spend most of their time outside the nest, where they eventually die [8,65]. Similarly, among honeybees, individuals whose health is compromised eat less, transfer less nectar to the hive [66], spend less time in the hive [67], and leave it to die in isolation [66]. This compulsion to leave the hive may explain sudden mass desertions observed in the recent epidemic of collapsed colony disease (CCD), regardless of the elusive pathogen that induces it [68]. The behavior of parasitized termites has been less studied, but infected individuals seem to migrate to bottom strata of mounds and die there [69].

Communicating health status is an important aspect of social immunity. The bulk of our knowledge concerns “hygienic behavior” in honeybees. In this process, infected larvae and pupae are detected and removed from the hive by workers, limiting the spread of infections [70,71]. Evidently, the brood communicates its health status chemically at the earliest sign of infection. Recently, it was shown that adult bees can also be expelled from the hive based on similar signals [71]. The behavioral component of such signaling is clearer in dampwood termites in which adults that have contacted fungal spores signal through vibration to repel colony members [60].

Studies in rodents implicated the vomeronasal organ in sensing infection [72] and discouraging social and sexual interactions [59]; importantly, immune activation with LPS was enough to mark animals as sick. Even in humans, mammals with an ill-reputed sense of smell, the clothes of LPS-treated subjects can be sniffed out [73].

It is easy to accept that the detection of infected conspecifics has evolved as a protective avoidance mechanism, but the transmission of such signals could also have been selected for.
Several symptoms of SB may act as infection cues: reduced self-grooming visibly distinguishes infected individuals as scruffy [1] and probably accentuates the olfactory signals they emit. Similar changes may affect vocal communication. In sparrows, for instance, the frequency and pattern of birdsong change during an inflammatory response [74]. Lastly, the stereotypic posture and motion that infected animals adopt because of fatigue and hyperalgesia can act as additional cues. Thus, LPS-treated subjects can be detected by observers based on their gait [75]. The signaling aspects of such behavioral changes are exposed by the response of sick animals to predators. Under the gaze of carnivores, sick members of a herd would attempt to disguise their vulnerability and suppress SB [14]. This observation suggests that animals can alert their kin of infection but suppress such signaling to predators.

**Premise 2: Reduced Contacts Limit the Spread of Infections**

Medicine has long acknowledged the importance of isolation for containing infectious disease in humans. Behavioral interventions such as quarantine, school closures, and bans on travel and public gathering have curtailed the spread of contagious diseases such as Ebola [76], vector-mediated diseases such as bubonic plague [77], and airborne ones such as severe acute respiratory syndrome (SARS) [78]. These successes demonstrate that, regardless of the route, social isolation can reduce transmission.

A question more relevant to the evolution of SB is whether self-imposed social isolation is effective in the wild. Several such examples exist: in the last decade, bat populations of many species in North America collapsed because of the "white nose" fungal disease. Although almost all of the colonies observed were decimated, some bat populations survived by adopting a solitary roosting pattern [79]. Conversely, a study in wild deer mice has shown that highly active individuals, which encountered more mice, exhibited higher viral infection rates [80].

Isolation of infected people based on clinical symptoms can be effective only when they overlap with the infectious period [81]. Empirical data suggest that, in the few infectious diseases studied (barring HIV), this is indeed the case. Thus, in SARS, smallpox, and foot-and-mouth disease, this overlap exceeds 80% [81,82], and estimates for influenza range between 50% and 90% [81,83]. Since behavioral symptoms typically precede specific clinical signs, these figures likely underestimate the overlap between SB and infectivity and the potential reduction in transmission.

**Premise 3: Behaviors That Reduce Pathogen Transmission Can Persist through Kin Selection**

If indeed SB favors the fitness of other group members at the expense of the individuals, then it can be considered an instance of biological altruism. It has long been debated how altruism can become an evolutionarily stable strategy (ESS). A likely mechanism is kin selection, the positive selection of traits that increase the fitness of the individual’s relatives. This initially controversial theory, put forward by W. D. Hamilton [84], has been mathematically validated and widely accepted since [85].

Kin selection is easy to accept when altruism is actively directed at relatives (e.g., birds feigning injury to lead predators away from their chicks), but how can it promote SB, a response that indiscriminately favors related and unrelated group members? This can only happen when the average relatedness within the social group is higher than within the entire population. Indeed, in many (although certainly not all) species, genetically related individuals are disproportionately represented in the immediate social groups in which most physical interactions occur [86–89]. Such bias develops because of high population viscosity, i.e., slow and spatially restricted dispersal of progeny.
Animal species vary in the degree of intergroup relatedness based on their life history. At one end of the spectrum are r-strategists whose offspring are neonatally independent and disperse widely. Under such conditions, social considerations are unlikely to drive SB. At the opposite end of the spectrum lie eusocial animals. Among these, eusocial hymenoptera have been studied most (Box 2). These insects indeed display a variety of collective disease defense behaviors, in part resembling SB, which are collectively termed “social immunity” [61]. Humans, classical K-strategists who cohabit most of their lives with first-degree relatives, seem to lie closer to this pole.

Intriguingly, some experimental evidence suggests that SB is actually not as universal as commonly assumed. Some birds can become infected, mount an immune response and develop fever without showing conspicuous signs of illness [90], leading to an apparently sudden death from infection [14]. In fish, administration of LPS triggers no observable behavioral changes [91]. Studies in wild mouse populations showed that the intensity of SB varies considerably among related species [10]. How this diversity relates to social structure is yet to be examined.

Where Do We Go from Here?

The Eyam hypothesis has never been directly tested, so the empirical evidence supporting it is still limited; nonetheless, it produces testable predictions. As stated above, an ESS for SB would counterbalance its cost to the infected individuals with the benefit of reducing transmission to their kin. This benefit should be proportional to pathogen virulence, the chances of transmission between individuals (infectivity), and the average relatedness of susceptible hosts. Consequently, several predictions can be examined either correlatively (1-3), experimentally (4, 5), or mathematically (6, 7).

1. **Virulence:** Different pathogens invoke SB of varying intensities. Our theory predicts that, through an evolutionary process, more virulent pathogens would come to provoke stronger behavioral responses. When a pathogen is deadly, the individual loses little (as it would die anyway) and gains much (as it saves its relatives from death) from a debilitating behavioral response. When a pathogen is avirulent, the optimal behavioral response would be a subclinical one, invoking no SB even if an immune response is activated.

2. **Disease transmission:** Increased odds for transmission would also favor a vigorous SB. Thus, more intense SB is expected among species that live in dense colonies and engage in close physical contact (especially in gregarious seasons). Likewise, highly contagious pathogens are expected to trigger a more pronounced SB.

3. **Genetic relatedness:** Higher relatedness within social groups should also promote SB (as it would other altruistic behaviors). Thus, SB would intensify with population viscosity and the length of care for offspring. This could be tested by comparing phylogenetically close species (e.g., social versus solitary wasps). Comparisons can also be made among individuals within communities: for example, mothers versus fathers in polygamous species and reproductive versus nonreproductive members of eusocial communities.

4. **Anti-inflammatory drugs:** Pharmacologically suppressing SB in experimentally infected individuals should accelerate the spread of infections even when the course of disease is unaltered. Such experiments, though, would require habitats that allow efficient self-isolation.

5. **Tracers:** To rule out immunological effects on pathogen clearance, the dispersal of innocuous tags such as pigments or radioisotopes can be traced.
6. **Mathematical models**: These could be developed to formally examine the feasibility of the hypothesis.

7. **Computer simulations**: These could be applied to test the hypothesis numerically.

We are so used to malaise being the essence of infection that we often forget to ask why it evolved. The social implications of SB may not be the only selective force driving it, but they clearly contribute and have been disregarded for too long. The mystery of SB is not only intellectually provoking but also clinically significant. This is because behavioral symptoms are routinely relieved using anti-inflammatory drugs such as cyclooxygenase (COX) inhibitors. According to the Eyam hypothesis, such use could prove socially irresponsible. By enabling infected people to travel widely and socialize, it interferes with a natural mechanism that prevents pathogen spread. In contrast, SB that accompanies medical procedures (such as cytokine treatment) and noninfectious diseases (such as cachexia in cancer) is a side effect that could be treated safely.

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**References**

1. Hart BL. Biological basis of the behavior of sick animals. Neurosci Biobehav Rev. 1988; 12: 123–137. PMID: 3050629
2. Aubert A. Sickness and behaviour in animals: A motivational perspective. Neurosci Biobehav Rev. 1999; 23: 1029–1036. PMID: 10580315
3. Dantzer R, Kelley KW. Twenty years of research on cytokine-induced sickness behavior. Brain Behav Immun. 2007; 21: 153–160. PMID: 17088043
4. McCusker RH, Kelley KW. Immune-neural connections: how the immune system’s response to infectious agents influences behavior. J Exp Biol. 2013; 216: 84–98. doi: 10.1242/jeb.073411 PMID: 23225871
5. Dantzer R, O’Connor JC, Freund GG, Johnson RW, Kelley KW. From inflammation to sickness and depression: when the immune system subjugates the brain. Nat Rev Neurosci. 2008; 9: 46–56. PMID: 18073775
6. Saper CB, Romanovsky AA, Scammell TE. Neural circuitry engaged by prostaglandins during the sickness syndrome. Nat Neurosci. 2012; 15: 1088–1095. doi: 10.1038/nn.3159 PMID: 22837039
7. Owen-Ashley NT, Turner M, Hahn TP, Wingfield JC. Hormonal, behavioral, and thermoregulatory responses to bacterial lipopolysaccharide in captive and free-living white-crowned sparrows (Zonotrichia leucophrys gambelii). Horm Behav. 2006; 49: 15–29. PMID: 15967447
8. Bos N, Lefèvre T, Jensen AB, D’Ettorre P. Sick ants become unsocial. J Evol Biol. 2012; 25: 342–351. doi: 10.1111/j.1420-9101.2011.02425.x PMID: 22122288
9. Aubert A, Richard F-JJ. Social management of LPS-induced inflammation in Formica polyctena ants. Brain Behav Immun. 2008; 22: 833–837. doi: 10.1016/j.bbi.2008.01.010 PMID: 18331785
10. Martin LB, Weil ZM, Nelson RJ. Fever and sickness behaviour vary among congeneric rodents. Funct Ecol. 2008; 22: 68–77.
11. Hanssen SA, Hasselquist D, Foslund I, Erikstad KE. Costs of immunity: immune responsiveness reduces survival in a vertebrate. Proc Biol Sci. 2004; 271: 925–930. PMID: 15255047
12. Morét Y, Schmid-Hempel P. Survival for immunity: the price of immune system activation for bumblebee workers. Science. 2000; 290: 1166–1168. PMID: 11073456
13. Eraud C, Jacquet A, Faivre B. Survival cost of an early immune soliciting in nature. Evolution. 2009; 63: 1036–1043. doi: 10.1111/j.1558-5646.2008.00540.x PMID: 19055677
14. Tizard I. Sickness behavior, its mechanisms and significance. Anim Health Res Rev. 2008; 9: 87–99. doi: 10.1017/S1466253308001448 PMID: 18423072
15. Bonneau A, Mazuc J, Gonzalez G, Haussey C, Chastel O, Faivre B, et al. Assessing the cost of mounting an immune response. Am Nat. 2003; 161: 367–379. PMID: 12703483
16. Aubert A, Goodall G, Dantzer R, Gheusi G. Differential effects of lipopolysaccharide on pup retrieving and nest building in lactating mice. Brain Behav Immun. 1997; 11: 107–118. PMID: 9299060
17. Avitsur R, Yirmiya R. The immunobiology of sexual behavior: gender differences in the suppression of sexual activity during illness. Pharmacol Biochem Behav. 1999; 64: 787–796. PMID: 10593202
18. Cohn DWH, Gabanyi I, Kinoshita D, de Sá-Rocha LC. Lipopolysaccharide administration in the dominant mouse destabilizes social hierarchy. Behav Processes. 2012; 91: 54–60. doi: 10.1016/j.beproc.2012.05.008 PMID: 22664349
19. Kluger MJ, Kozak W, Conn CA, Leon LR, Soszynski D. Role of Fever in Disease. Ann N Y Acad Sci. 1998; 856: 224–233. PMID: 9917881
20. Earn DJD, Andrews PW, Bolker BM. Population-level effects of suppressing fever. Proc R Soc B-Biological Sci. 2014; 281: 20132570.
21. Evans SS, Repasky EA, Fisher DT. Fever and the thermal regulation of immunity: the immune system feels the heat. Nat Rev Immunol. 2015; 15: 335–349. doi: 10.1038/nri3843 PMID: 25976513
22. Jefferies S, Weatherall M, Young P, Eyers S, Beasley R. Systematic review and meta-analysis of the effects of antipyretic medications on mortality in Streptococcus pneumoniae infections. Postgrad Med J. 2012; 88: 21–27. doi: 10.1136/postgradmedj-2011-130217 PMID: 22121249
23. Eyers S, Weatherall M, Shirtcliffe P, Perrin K, Beasley R. The effect on mortality of antipyretics in the treatment of influenza infection: systematic review and meta-analysis. J R Soc Med. 2010; 103: 403–411. doi: 10.1258/jrsm.2010.090441 PMID: 20929891
24. Pursell E, While AE. Does the use of antipyretics in children who have acute infections prolong febrile illness? A systematic review and meta-analysis. J Pediatr. 2013; 163: 822–827. doi:10.1016/j.jpeds.2012.05.008 PMID: 22664349
25. Meremikwu MM, Odigwe CC, Akudo Nwagbara B, Udoh EE. Antipyretic measures for treating fever in malaria. Cochrane database Syst Rev. 2012; 9: CD002151. doi:10.1002/14651858.CD002151.pub2 PMID: 22972057
26. Tarimo DS, Minjas JN, Bygbjerg IC. Sulfadoxine-pyrimethamine monotherapy in Tanzanian children gives rapid parasite clearance but slow fever clearance that is improved by chloroquine in combination therapy. Trop Med Int Heal. 2002; 7: 592–598.
27. Brandts SIRC, Ndjave M, Graninger W. Effect of paracetamol on parasite clearance time in Plasmodium falciparum malaria. Cutaneous T-cell lymphoma severity index and T-cell gene rearrangement. Lancet. 1997; 350: 704–709. PMID: 9291905
28. Krishna S, Supananarond W, Pukrittayakamee S, ter Kuile F, Supputamangkol Y, Attatamsoonthorn K, et al. Fever in uncomplicated Plasmodium falciparum infection: effects of quinine and paracetamol. Trans R Soc Trop Med Hyg. 1995; 89: 197–199. PMID: 7778148
29. Doran TF, De Angelis C, Baumgardner RA, Mellits ED. Acetaminophen: more harm than good for chickenpox? J Pediatr. 1989; 114: 1045–1048. PMID: 2656959
30. Weinberg ED. Iron availability and infection. Biochim Biophys Acta—Gen Subj. 2009; 1790: 600–605.
31. Drakesmith H, Prentice AM. Hepcidin and the iron-infection axis. Science. 2012; 338: 768–772. doi:10.1126/science.1224577 PMID: 23139325
32. Schaible UE, Kaufmann SHE. Iron and microbial infection. Nat Rev Microbiol. 2004; 2: 946–953. PMID: 15550940
33. Ashley NT, Wingfield JC. Sickness behavior in vertebrates. In: Demas Gregory, editor. Ecoimmunology. Oxford University Press; 2011. pp. 45–91.
34. Murray MJ, Murray AB. Anorexia of infection as a mechanism of host defense. Am J Clin Nutr. 1979; 32: 593–596. PMID: 283688
35. Adamo SA. Comparative psychoneuroimmunology: evidence from the insects. Behav Cogn Neurosci Rev. 2006; 5: 128–140. PMID: 16891555
36. Maslowski KM, Mackay CR. Diet, gut microbiota and immune responses. Nat Immunol. 2011; 12: 5–9. doi:10.1038/ni0111-5 PMID: 21169997
37. Lopes PC. When is it socially acceptable to feel sick? Proc R Soc B. 2014; 281: 20140218. doi: 10.1098/rspb.2014.0218 PMID: 24943375
38. Adelman JS, Martin LB. Vertebrate sickness behaviors: Adaptive and integrated neuroendocrine immune responses. Integr Comp Biol. 2009; 49: 202–214. doi:10.1093/icb/icp028 PMID: 21665814
39. Exton MS. Infection-induced anorexia: active host defence strategy. Appetite. 1997; 29: 369–383. PMID: 9468766
40. Kent S, Bluthé RM, Kelley KW, Dantzer R. Sickness behavior as a new target for drug development. Trends Pharmacol Sci. 1992; 13: 24–28. PMID: 1542935
41. Thiessen DD. Body temperature and grooming in the Mongolian gerbil. Ann N Y Acad Sci. 1988; 525: 27–39. PMID: 3291667

42. Shanas U, Terkel J. Grooming secretions and seasonal adaptations in the blind mole rat (Spalax ehrenbergi). Physiol Behav. 1996; 60: 653–656. PMID: 8840931

43. Hutchings MR, Athanasiadou S, Kyriazakis I, Gordon IJ. Can animals use foraging behaviour to combat parasites? Proc Nutr Soc. 2003; 62: 361–370. PMID: 14506883

44. Hooda PS, Henry CJK, Seyoum T a., Armstrong LDM, Fowler MB. The potential impact of soil ingestion on human mineral nutrition. Sci Total Environ. 2004; 333: 75–87. PMID: 15364520

45. Ganz T, Nemeth E. Hepcidin and disorders of iron metabolism. Annu Rev Med. 2011; 62: 347–360. doi: 10.1146/annurev-med-050109-142444 PMID: 20887198

46. Palmblad J. Fasting (acute energy deprivation) in man: effect on polymorphonuclear granulocyte functions, plasma iron and serum transferrin. Scand J Haematol. 1976; 17: 217–226. PMID: 968452

47. Kemna E, Pickens P, Nemeth E, Van Der Hoeven H, Swinkels D. Time-course analysis of hepcidin, serum iron, and plasma cytokine levels in humans injected with LPS. Blood. 2005; 106: 1864–1866. PMID: 15986319

48. Watkins LR, Maier SF. The pain of being sick: implications of immune-to-brain communication for understanding pain. Annu Rev Psychol. 2000; 51: 29–57. PMID: 10751964

49. Medzhitov R, Schneider DS, Soares MP. Disease Tolerance as a Defense Strategy. Science. 2012; 335: 936–941. doi: 10.1126/science.1214935 PMID: 22363001

50. Shattuck EC, Muehlenbein MP. Human sickness behavior: Ultimate and proximate explanations. Am J Phys Anthropol. 2015; 157: 1–18. doi: 10.1002/ajpa.22698 PMID: 25639499

51. Johnson RW. The concept of sickness behavior: a brief chronological account of four key discoveries. Vet Immunol Immunopathol. 2002; 87: 443–450. PMID: 12072271

52. Hawley DM, Altizer SM. Disease ecology meets ecological immunology: Understanding the links between organismal immunity and infection dynamics in natural populations. Funct Ecol. 2011; 25: 48–60.

53. Massad E, Coutinho FAB, Burattini MN, Lopez LF. The Eyam plague revisited: Did the village isolation change transmission from fleas to pulmonary? Med Hypotheses. 2004; 63: 911–915. PMID: 15488668

54. Cheeseman C, Mallinson P. Behaviour of badgers (Meles meles) infected with bovine tuberculosis. J Zool. 1981; 42: 284–289.

55. Poulin R. Parasite Manipulation of Host Behavior: An Update and Frequently Asked Questions. In: Brockmann HJ, Roper TJ, Naguib M, Wynne-Edwards KE, Mitani JC, Simmons LW, et al., editors. Advances in the Study of Behavior. 1st ed. Elsevier Inc.; 2010. pp. 151–186.

56. Schaller M, Park JH. The Behavioral Immune System (and Why It Matters). Curr Dir Psychol Sci. 2011; 20: 99–103.

57. Behringer DC, Butler MJ, Shields JD. Avoidance of disease by social lobsters. Nature. 2006; 441: 421. PMID: 16724051

58. Kiesecker JM, Skelly DK, Beard KH, Preisser E. Behavioral reduction of infection risk. Proc Natl Acad Sci U S A. 1999; 96: 9165–9168. PMID: 10430913

59. Arakawa H, Cruz S, Deak T. From models to mechanisms: odorant communication as a key determinant of social behavior in rodents during illness-associated states. Neurosci Biobehav Rev. 2011; 35: 1916–1928. doi: 10.1016/j.neubiorev.2011.03.007 PMID: 21414355

60. Rosengaus R, Jordan C, Lefebvre M, Traniello J. Pathogen alarm behavior in a termite: A new form of communication in social insects. Naturwissenschaften. 1999; 86: 544–548. PMID: 10551951

61. Cremer S, Armitage S a O, Schmid-Hempel P. Social Immunity. Curr Biol. 2007; 20: 159–160.

62. Stroeymeyt N, Casillas-Pérez B, Cremer S. Organisational immunity in social insects. Curr Opin Insect Sci. 2014; 5: 1–15.

63. Ugelvig LV, Cremer S. Social prophylaxis: group interaction promotes collective immunity in ant colonies. Curr Biol. 2007; 17: 1967–1971. PMID: 17980590

64. Theis FJ, Ugelvig LV, Marr C, Cremer S. Opposing effects of allogrooming on disease transmission in ant societies. Philos Trans R Soc Lond B Biol Sci. 2015; 370: 20140108. doi: 10.1098/rstb.2014.0108 PMID: 25870394

65. Heinze J, Walter B. Moribund ants leave their nests to die in social isolation. Curr Biol. 2010; 20: 249–252. doi: 10.1016/j.cub.2009.12.031 PMID: 20116243

66. Rueppell O, Hayworth MK, Ross NP. Altruistic self-removal of health-compromised honey bee workers from their hive. J Evol Biol. 2010; 23: 1538–1546. doi: 10.1111/j.1420-9101.2010.02022.x PMID: 20500363
67. Dussaubat C, Maisonnasse A, Crusater D, Beslay D, Costagliola G, Soubeyrand S, et al. Flight behavior and pheromone changes associated to Nosema ceranae infection of honey bee workers (Apis mellifera) in field conditions. J of Invertebr Pathol. 2013; 42–51.

68. Core A, Runceel C, Ivers J, Quock C, Siapno T, DeNault S, et al. A new threat to honey bees, the parasitic phorid fly Apocephalus borealis. PLoS One. 2012; 7: 1–9.

69. Lenz M. Biological Control in Termite Management: the Potential of Nematodes and Fungal Pathogens. Fifth International Conference on Urban Pests. 2005. 47–52.

70. Wilson-Rich N, Spivak M, Fefferman NH, Starks PT. Genetic, individual, and group facilitation of disease resistance in insect societies. Annu Rev Entomol. 2009; 54: 405–423. doi: 10.1146/annurev.ento.53.103106.093031 PMID: 18793100

71. Baracchi D, Fadda A, Turillazzi S. Evidence for antiseptic behaviour towards sick adult bees in honey bee colonies. J Insect Physiol. 2012; 58: 1589–1596. doi: 10.1016/j.jinsphys.2012.09.014 PMID: 23068993

72. Boillat M, Challet L, Rossier D, Kan C, Carleton A, Rodriguez I. The vomeronasal system mediates sick conspecific avoidance. Curr Biol. 2015; 25: 251–255. doi: 10.1016/j.cub.2014.11.061 PMID: 25578906

73. Olsson MJ, Lundström JN, Kimball BA, Gordon AR, Kaschikoff B, Hosseini N, et al. The scent of disease: human body odor contains an early chemosensory cue of sickness. Psychol Sci. 2014; 25: 817–23. doi: 10.1177/0956797613515681 PMID: 24452606

74. Munoz NE, Blumstein DT, Foufopoulos J. Immune system activation affects song and territorial defense. Behav Ecol. 2010; 21: 788–793.

75. Sundelin T, Kaschikoff B, Axelsson E, Höglund CO, Lekander M, Axelsson J. Sick man walking: Perception of health status from body motion. Brain Behav Immun. 2015; 48: 53–56. doi: 10.1016/j.bbi.2015.03.007 PMID: 25801061

76. Chowell G, Nishiura H. Transmission dynamics and control of Ebola virus disease (EVD): a review. BMC Med. 2014; 12: 1–16.

77. Tognotti E. Lessons from the history of quarantine, from plague to influenza A. Emerg Infect Dis. 2013; 19: 254–259. doi: 10.3201/eid1902.120312 PMID: 23343512

78. Riley S, Fraser C, Donnelly CA, Ghani AC, Abu-Raddad LJ, Hedley AJ, et al. Transmission dynamics of the etiological agent of SARS in Hong Kong: impact of public health interventions. Science. 2003; 300: 1961–1966. PMID: 12766206

79. Langwig KE, Bried JT, Hicks AC, Kunz TH, Marm Kilpatrick A. Sociality, density-dependence and microclimates determine the persistence of populations suffering from a novel fungal disease, white-nose syndrome. Ecol Lett. 2012; 15: 1050–1057. doi: 10.1111/j.1461-0248.2012.01829.x PMID: 22747672

80. Dizney L, Dearing MD. The role of behavioural heterogeneity on infection patterns: Implications for pathogen transmission. Anim Behav. 2013; 86: 911–916.

81. Fraser C, Riley S, Andersson RM, Ferguson NM. Factors that make an infectious disease outbreak controllable. Proc Natl Acad Sci U S A. 2004; 101: 6146–6151. PMID: 15071187

82. Charleston B, Bankowski BM, Gubbins S, Chase-Topping ME, Schley D, Howey R, et al. Relationship between clinical signs and transmission of an infectious disease and the implications for control. Science. 2011; 332: 726–729. doi: 10.1126/science.1199884 PMID: 21951063

83. Carrat F, Vergu E, Ferguson NM, Lemaitre M, Cauchemez S, Leach S, et al. Time lines of infection and disease in human influenza: a review of volunteer challenge studies. Am J Epidemiol. 2008; 167: 775–785. doi: 10.1093/aje/kwm375 PMID: 18230677

84. Hamilton WD. The genetical evolution of social behaviour. II. J Theor Biol. 1964; 7: 17–52. PMID: 5875340

85. Foster KR, Wenseleers T, Ratnieks FLW. Kin selection is the key to altruism. Trends Ecol Evol. 2006; 21: 57–60. PMID: 16701471

86. Archie EA, Moss CJ, Alberts SC. The ties that bind: genetic relatedness predicts the fission and fusion of social groups in wild African elephants. Proc Biol Sci. 2006; 273: 513–522. PMID: 16537121

87. Gompper ME, Gittleman JL, Wayne RK. Genetic relatedness, coalitions and social behaviour of white-nosed coats, Nasua narica. Anim Behav. 1997; 53: 781–797.

88. Madden JR, Nielsen JF, Clutton-Brock TH. Do networks of social interactions reflect patterns of kinship? Curr Zool. 2012; 58: 319–328.

89. Dierkes P, Heg D, Taborsky M, Skubic E, Achmann R. Genetic relatedness in groups is sex-specific and declines with age of helpers in a cooperatively breeding cichlid. Ecol Lett. 2005; 8: 968–975.

90. Marais M, Maloney SK, Gray DA. Sickness behaviours in ducks include anorexia but not lethargy. Appl Anim Behav Sci. 2013; 145: 102–108.
91. Swain P, Nayak SK, Nanda PK, Dash S. Biological effects of bacterial lipopolysaccharide (endotoxin) in fish: A review. Fish Shellfish Immunol. 2008; 25: 191–201. doi: 10.1016/j.fsi.2008.04.009 PMID: 18603445