Individual differences in boldness influence patterns of social interactions and the transmission of cuticular bacteria among group-mates

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Despite the importance of host attributes for the likelihood of associated microbial transmission, individual variation is seldom considered in studies of wildlife disease. Here, we test the influence of host phenotypes on social network structure and the likelihood of cuticular bacterial transmission from exposed individuals to susceptible group-mates using female social spiders (Stegodyphus dumicola). Based on the interactions of resting individuals of known behavioural types, we assessed whether individuals assorted according to their behavioural traits. We found that individuals preferentially interacted with individuals of unlike behavioural phenotypes. We next applied a green fluorescent protein-transformed cuticular bacterium, Pantoea sp., to individuals and allowed them to interact with an unexposed colony-mate for 24 h. We found evidence for transmission of bacteria in 55% of cases. The likelihood of transmission was influenced jointly by the behavioural phenotypes of both the exposed and susceptible individuals: transmission was more likely when exposed spiders exhibited higher ‘boldness’ relative to their colony-mate, and when unexposed individuals were in better body condition. Indirect transmission via shared silk took place in only 15% of cases. Thus, bodily contact appears key to transmission in this system. These data represent a fundamental step towards understanding how individual traits influence larger-scale social and epidemiological dynamics.

1. Background

The transmission of microorganisms from exposed to susceptible hosts represents one of the most crucial forces regulating infectious disease dynamics. Recent attention [1,2], however, has scrutinized historical attempts to quantify and predict transmission dynamics. For example, a foundational principle of epidemiology, ‘the mass action principle’, relies on the assumption that the course of an epidemic is determined by the rate of random contacts between infected and susceptible individuals [3]. Of course, populations are not comprised individuals who interact randomly, and interactions during epidemics do not always result in transmission [4]. Variation in host behavioural phenotypes and social interactions remain putative explanations of the immense transmission heterogeneity observed in human and wildlife diseases [5–7]. Consequently, an important topic of current discourse in disease ecology has focused on understanding how consistent behavioural variation among individuals (i.e. behavioural types, syndromes or personality [8,9]) may influence the dynamics of microbial transmission [10].

Many of the most prevalent and deadly diseases threatening wildlife and human populations are characterized by intense variation in the degree to which infected individuals produce secondary cases of infection [5]. In fact, host
heterogeneity in infection susceptibility [11] and pathogen transmission [5,6,12] are becoming increasingly emphasized in disease ecology and epidemiology. In addition, a group’s behavioural composition may also have consequences for the spread of infectious agents [13]. A recent study on behavioural variation and disease dynamics in house finches found that an individuals’ propensity to use experimental feeders increased both the acquisition and transmission of a bacterial pathogen [14]. Such studies are sorely needed because they employ experimental infections and characterize interindividual variation in behaviours that are predicted to influence transmission dynamics.

Variation among individuals in how frequently and with whom they interact has important implications for epidemiological dynamics. For example, individuals that interact with others more frequently can have an inordinately large effect on driving the spread of an epidemic [5]. Both individual-level (e.g. infection status) and group-wide processes (e.g. disease outbreaks) are influenced by the connectivity of individuals [15–17]. The incorporation of social network theory into models of disease dynamics has shown that the structure of the interaction network and the processes that underlies its formation can affect transmission rate [18–20]. Therefore, empirical studies of the processes that drive social network formation are crucial for our understanding of disease dynamics. A common mechanism of network formation is assortativity according to a certain trait, that is, individuals often tend to interact with others of similar phenotypes [21]. Just as common is the complementary mechanism, disassortativity, where individuals tend to interact with individuals of opposing phenotypes [22,23]. These interaction preferences will shape network structure and may influence the transmission rate of microbes among individuals [7,18].

Some prior studies have identified natural relationships between host personality type and infection status [24–26], though fewer studies have experimentally differentiated behaviour-mediated infection from parasite manipulation of behaviour (notable exceptions include: [27,28]) (reviewed in [29]). Others report evidence for associations between host behavioural traits and virus transmission [30], and experimental infections of laboratory animals have generated enormous variation in pathogen shedding rates associated with host traits like co-infection status and immunocompetence [31,32]. Despite these advances, studies rarely, if ever, evaluate the role of behavioural phenotypes in both infected and susceptible individuals simultaneously to test their joint effects on multiple modes of transmission.

Here, we examine how variation among individuals in consistent behavioural traits influences contact network formation and bacterial transmission in the social spider Stegodyphus dumicola (Araneae, Eresidae). In colonies of S. dumicola, the execution of collective behaviours such as prey capture [33–36] and antipredator defences [37] is associated with heterogeneity among individuals in their boldness, suggesting that boldness is a reliable indicator of individuals’ role in these societies. Therefore, we examined assortativity according to boldness by observing marked individuals in experimental colonies. To examine how the boldness of both exposed and susceptible individuals influences the likelihood of direct and indirect (i.e. environmental) transmission, we experimentally exposed individual spiders with a green fluorescent protein (GFP)-transformed cuticular bacterium. We focused on cuticular bacteria, because the integument represents the barrier between the host and constant bombardment by microbes from its environment, representing a primary line of defence against invading microbes, and body contact represents a likely mode of transmission for cuticle-associated microbes [38,39]. We hypothesize that cuticular bacterial transmission will be more likely to occur between individuals who are more likely to interact in observed contact networks. That is, if colony contact networks are assortative, then cuticular bacterial transmission should be more likely between similar individuals. In contrast, if colony networks are disassortative, then transmission should be more likely to occur between dissimilar individuals. Lastly, we hypothesize that cuticular bacteria will be primarily transmitted through direct interactions (i.e. bodily contact) rather than indirectly through the environment.

2. Material and methods

(a) Animal collection and maintenance

Stegodyphus dumicola is a southwestern African social spider that lives in age-structured colonies of up to several hundred individuals that exhibit cooperative behaviours and alliparental care, and spend the majority of their time in close contact with colony-mates, either in the colony retreat or during co-feeding [40–43]. We collected 16 S. dumicola colonies along roadside Acacia trees in the Northern Cape of South Africa in January 2015. After transport to the laboratory, individual adult females were isolated into 30 ml plastic cups containing a piece of chicken wire to facilitate web-building. Only adult female spiders were used in the present study. Spiders were each fed one two-week-old cricket weekly until the onset of behavioural assays.

(b) Behavioural assays

In Stegodyphus, ‘boldness’ (the latency to resume movement after experiencing an aversive stimulus [44]) and aggressiveness are highly consistent behavioural metrics (repeatability = 0.63 and 0.55, respectively; [45]) and are linked with an individual’s propensity to participate in a variety of collective tasks [45–48]. To determine individuals’ boldness, we subjected them to an antipredator behaviour assay developed by Riechert & Hedrick [49]. The spider is placed in a clear plastic arena (12 cm diameter), given a 30 s acclimation period, and then administered two rapid puffs of air to the anterior prosoma using an infant nose-clearing bulb which causes them to ‘huddle’ by halting movement and pulling the legs close to the body. We then measured the latency for spiders to ‘unhuddle’ and move one full body length. ‘Bold’ individuals unhuddle and resumed movement more quickly, whereas ‘shy’ individuals have longer latencies to resume activity. We subtract the latency for a spider to resume movement from the maximum latency allowed (600 s) such that a higher boldness score represents bolder behaviour.

For the bacterial transmission experiments, we also assessed individuals’ aggressiveness by placing the spider in a plastic arena (12 cm diameter), allowing it a 30 s acclimation period, and then pricking their foremost left leg with a blunt metal probe. We scored their immediate response to this stimulus with a nominal categorization described previously [45,46]. ‘Non-aggressive’ behaviours included a ‘huddle’ response and moving away from stimulus, whereas ‘aggressive’ behaviours included turning or walking towards the stimulus, raising their anterior legs and shifting the abdomen in place. Aggressive-ness assays took place the same day as boldness assays, approximately 6 h later.
and growing it in 1 ml LB broth supplemented with 100 μg ml\(^{-1}\) ampicillin and 20% arabinose ('LB amp/ara'; electronic supplementary material, text S2) for 15 h with agitation. This solution was vortexed at 2500 r.p.m. for 3 s with a vortex to disrupt the hydrophobic barrier of the spider’s cuticle. These bacteria remain viable and detectable on the cuticles of spiders at least 72 h after exposure (electronic supplementary material, text S3). Henceforth, we will refer to experimentally exposed individuals as ‘exposed’ and those that were exposed only to PBS and marked with a green paint dot as ‘susceptible’. Prior to bacterial exposure, we measured each spider’s mass and prosoma width. All bacterial exposures were carried out by the same two researchers (C.N.K. and D.A.A.) to minimize methodological inconsistencies.

(c) Social interactions
We observed interactions in laboratory colonies to determine whether individuals assort according to boldness and whether there is a relationship between boldness and the number of associates of resting spiders. Nine colonies of 10–30 adult female spiders of known boldness (see boldness assay above), and individually marked with acrylic paint dots atop their dorsal abdomen, were kept in plastic containers with chicken wire that allowed them to build a retreat and a capture web. Each experimental colony was constructed from a different source colony, without mixing individuals from multiple source colonies. These spiders spend much of their time resting in the colony retreat, often in groups. We defined interactions between resting group members as a physical contact between any body parts of two spiders. We manually noted the resting interaction patterns of all individuals in each colony two to four times a week (figure 1). Repeated observations of the same colony occurred either on different days or on the same day if the colony had an opportunity to re-assort. For example, an observation was conducted before a colony was fed or provided with water and, if the colony responded to the prey or water, another resting network observation was conducted after the collective response ended and the spiders resumed resting (electronic supplementary material, table S1).

(d) Bacterial exposure
We used electroporation [50] to transform a strain of Pantoaea (CNK01) collected from the cuticle of an adult female spider in the field in January 2014 (methods described in Keiser et al. 2016 – in Current Zoology) with the pGLO plasmid (BioRad, Hercules, CA) that encodes β-lactamase (confering ampicillin resistance) and green fluorescent protein (henceforth CNK02; electronic supplementary material, text S1). Experimental bacterial cultures were prepared by selecting a single colony of CNK02 and growing it in 1 ml LB broth supplemented with 100 μg ml\(^{-1}\) ampicillin and 20% arabinose ('LB amp/ara'; electronic supplementary material, text S2) for 15 h with agitation. This solution was vortexed at 2500 r.p.m. for 25 min, washed with 1 ml phosphate-buffered saline (PBS, pH 7.4; Sigma-Aldrich, St Louis, MO 63103), and then diluted in 1 ml PBS. Ninety spiders marked with a blue paint dot were placed in 1 ml of liquid culture of CNK02 at approximately 10^8 CFU ml\(^{-1}\), and shaken at 1500 r.p.m. for 3 s with a vortex to disrupt the hydrophobic barrier of the spider’s cutae. These bacteria remain viable and detectable on the cuticles of spiders at least 72 h after exposure (electronic supplementary material, text S3). Henceforth, we will refer to experimentally exposed individuals as ‘exposed’ and those that were exposed only to PBS and marked with a green paint dot as ‘susceptible’. Prior to bacterial exposure, we measured each spider’s mass and prosoma width. All bacterial exposures were carried out by the same two researchers (C.N.K. and D.A.A.) to minimize methodological inconsistencies.

(e) Transmission experiments
To test for the transmission of bacteria between exposed and susceptible individuals when allowed to interact directly (i.e. via cuticle-to-cuticle contact), we exposed one spider as described above, and allowed it 24 h in its housing container to dry. We then transferred the exposed individual to the housing container of a susceptible individual with a different boldness value than the exposed individual, and allowed them to interact naturally for 24 h (n = 66 pairs). The identities of the spiders chosen for each pair were chosen using a random-number generator, and paired spiders always originated from the same source colony. Care was taken to place the exposed individual away from the susceptible individual in its home container to allow natural contact between spiders. After 24 h, we removed the susceptible individual and vortexed it in 1 ml sterile LB amp/ara broth for 10 s. We removed the spider, transferred 40 μl of this solution onto LB amp/ara agar, and incubated this plate and the remaining solution (960 μl) at 30°C for 20 h. We visually counted the number of LB amp/ara broth solutions that fluoresced under a long-wave UV light pen (BioRad) to assess successful transmission, and counted the number of colony forming units (CFUs) that grew and fluoresced on each LB amp/ara plate to approximate the relative bacterial load that had been transferred to susceptible individuals. We performed a series of side experiments to verify that (i) the pGLO plasmid is required to observe fluorescence from spider-collected bacterial cultures, (ii) that amplification at 30°C is required to visually detect fluorescence from GFP-transformed Pantoaea collected from spiders, (iii) that 20 h of amplification under 30°C is sufficient to detect fluorescence of bacteria collected from bold and shy spiders and that (iv) fluorescent bacteria are detectable on the cuticles of spiders for at least 72 h after exposure (electronic supplementary material, text S3).

(f) Statistical analyses
(i) Social interactions
To determine assortativity according to boldness of the resting networks, we used the igraph package in R [51]. Positive
assortativity values indicate assortative networks (i.e. homophily) and negative assortativity values indicate disassortative networks. To determine whether the observed assortativity values are different than expected at random, we created 10 000 randomized networks for each observed network maintaining the connectivity (degree) of each individual (node) and permuting who it interacted with. We then calculated the probability that an observed assortativity value differed from the 10 000 randomized assortativity values as the proportion of cases in which the absolute value of the observed assortativity was smaller than the absolute value of the randomized assortativity. To determine whether the difference between observed and randomized assortativity values was statistically significant, we compared the average assortativity of all observed networks with the average assortativity of each of the 10 000 sets of randomized networks. We deemed the observed assortativity significantly different from random if the absolute value of the observed average assortativity was smaller than the absolute value of the average randomized assortativity in less than 0.05 of the 10 000 randomizations. We examined the relationship between connectivity (degree) and boldness in each interaction network using Pearson’s correlation and Bonferroni correction for multiple hypotheses testing which set the significant p-value at 0.002.

(ii) Transmission experiments
We used two nominal logistic regressions and one GLMM with a log link function with identical independent variables to test for transmission via direct interaction, indirect transmission and the number of CFUs ml⁻¹ collected from susceptible individuals in the direct interaction experiment, respectively. The independent variables in each model were the difference in boldness between the two individuals, the aggressiveness of the exposed individual, the aggressiveness of the susceptible individual, the body condition of the susceptible individual and the body condition of the exposed individual. Values for the difference in boldness between exposed and susceptible individuals ranged from ~600 to 600 (electronic supplementary material, figure S2). Body condition was estimated using the residuals of a linear regression of individual body mass and body size [52]. We treated aggressiveness as a categorical variable here (following the methods of [45]). In a supplemental analysis (electronic supplementary material, text S4), we treated aggressiveness as an ordinal variable (following the methods of [46]), and used a ‘difference in aggressiveness’ value for each spider pair. We tested for correlations between measures of boldness and aggressiveness with nominal logistic regression. Experimental pair ID nested in source colony ID was included as a random effect in each model.

3. Results

(a) Social interactions
Observed networks were significantly more disassortative than expected at random. The observed average assortativity value was significantly smaller than that obtained from 10 000 randomizations (figure 2). Of the 36 resting interaction networks, 35 exhibited disassortative mixing, while individuals preferentially interacted with individuals of different boldness than their own. The probability that an observed assortativity value differed from the randomized assortativity for each network is provided in electronic supplementary material, table S1. We did not detect a significant relationship between the boldness of an individual and the number of individuals it contacted while resting in any of the interaction networks (electronic supplementary material, table S1).

(b) Bacterial transmission
Transmission of cuticular bacteria via direct interaction was influenced by the boldness of both the exposed and susceptible individuals. In the direct contact experiment, we detected the presence of CNK02 on the cuticles of 36/66 susceptible spiders (55%) that were allowed to interact with exposed spiders (figure 3). We detected the presence of CNK02 in 36/66 susceptible spiders (55%) that were allowed to interact with exposed spiders (figure 3). In the direct contact experiment, we detected the presence of CNK02 on the cuticles of 36/66 susceptible spiders (55%) that were allowed to interact with exposed spiders (figure 3). We detected the presence of CNK02 on the cuticles of 36/66 susceptible spiders (55%) that were allowed to interact with exposed spiders (figure 3). We detected the presence of CNK02 on the cuticles of 36/66 susceptible spiders (55%) that were allowed to interact with exposed spiders (figure 3). We detected the presence of CNK02 on the cuticles of 36/66 susceptible spiders (55%) that were allowed to interact with exposed spiders (figure 3). We detected the presence of CNK02 on the cuticles of 36/66 susceptible spiders (55%) that were allowed to interact with exposed spiders (figure 3). We detected the presence of CNK02 on the cuticles of 36/66 susceptible spiders (55%) that were allowed to interact with exposed spiders (figure 3). We detected the presence of CNK02 on the cuticles of 36/66 susceptible spiders (55%) that were allowed to interact with exposed spiders (figure 3). We detected the presence of CNK02 on the cuticles of 36/66 susceptible spiders (55%) that were allowed to interact with exposed spiders (figure 3). We detected the presence of CNK02 on the cuticles of 36/66 susceptible spiders (55%) that were allowed to interact with exposed spiders (figure 3). We detected the presence of CNK02 on the cuticles of 36/66 susceptible spiders (55%) that were allowed to interact with exposed spiders (figure 3). We detected the presence of CNK02 on the cuticles of 36/66 susceptible spiders (55%) that were allowed to interact with exposed spiders (figure 3). We detected the presence of CNK02 on the cuticles of 36/66 susceptible spiders (55%) that were allowed to interact with exposed spiders (figure 3). We detected the presence of CNK02 on the cuticles of 36/66 susceptible spiders (55%) that were allowed to interact with exposed spiders (figure 3).
We detected CNK02 on the cuticles of susceptible spiders in only 5/33 cases (15%) when exposed spiders interacted with a susceptible spider’s silk alone and never with the susceptible spider directly. We did, however, detect CNK02 on the silk with which exposed spiders interacted in 12/33 cases (36%). Thus, 5/12 cases (42%) where the silk became contaminated with CNK02 resulted in exposure to the susceptible individual. Evidence of indirect transmission was not influenced by the traits of either the exposed or susceptible individual (logistic regression: all $p > 0.10$; table 1).

4. Discussion

Interindividual variation in behaviour and contact networks of infected and susceptible individuals have vast consequences for many emerging diseases in wildlife [53] and humans [5]. Here, we did not find support for our original hypothesis, but rather found that *S. dumicola* contact networks are behaviourally disassortative, and that the transmission of cuticular bacteria is more likely when exposed individuals were bolder than their susceptible colony-mates. Thus, under some conditions, this system might be poised for rapid and widespread transmission of cuticular microbes, harmful or otherwise. Presuming that at least a subset of resident cuticular bacteria can be harmful under some circumstances, as is the case for this species [54], the observed social network pattern may help to explain the high incidence of idiopathic colony extinction in *S. dumicola* and other species of social spiders [41,55,56].

(a) Network patterns and bacterial transmission

In our observed spider contact networks, behavioural disassortativity was more prevalent than expected at random. Many animal and human networks, however, are characterized by positive assortativity, (i.e. homophily) [22,23,57], where individuals preferentially interact with others like themselves. It has been suggested that infectious agents spread more slowly when hosts engage in disassortative networks [22]. For instance, simulated outbreaks of foot and mouth disease in livestock are shortened owing to disassortative contacts [58]. Additionally, individuals often tend to avoid visibly infected conspecifics [59] whose recognition may be heightened via their altered behaviours, further reducing the likelihood of transmission. Whether, and to what degree, our observed social interaction patterns in *S. dumicola* would afford colonies reduced overall transmission among individuals (i.e. a form of ‘social immunity’; [20,60,61]), or alternatively, enhanced susceptibility to transmission is yet unresolved; we discuss both possibilities below.

Our data suggest that the ability of the disassortative nature of these spiders’ interactions to prevent or facilitate the transmission of cuticular microbes would depend on the traits of the exposed and susceptible individuals.
Transmission was more likely when there was a difference in the boldness of the exposed versus susceptible individual, but more so in one direction. When the exposed individual was bolder than the susceptible individual, the incidence of transmission was lower (42%). Conversely, when the susceptible individual was bolder than the exposed individual, transmission was more likely (69%). Thus, disassortative contact networks in *S. dumicola* colonies could, depending on the situation considered, either intensify or reduce the incidence of bacterial transmission throughout the colony. If the index case (aka ‘patient zero’) for a transmission event is a bold individual, this could beget rapid transmission to shyer colony members, which is the more common behavioural phenotype within these societies. However, if the index case is a shy individual, bacterial transmission to bolder individuals could be constrained. Although reduced, the observed incidence of transmission from shyer to bolder spiders was still considerable, and colony-wide bacterial transmission would not be completely quelled if the index case were a shy spider.

The aggregate effects that these interaction networks and patterns of transmission have on colony performance would thus depend on whether bold individuals or shy individuals are more likely to be the index case, and whether different behavioural phenotypes are differentially likely to encounter, and become colonized by, novel environmental microbes. In *S. dumicola*, the available data suggest that bold individuals more readily interact with the environment outside of the colony’s nest during foraging [33,62], suggesting that these individuals may be the pathway by which environmental microbes are subsequently transmitted to shy colony-mates.

(b) Transmission through contact versus via shared silk
Notably, we found that the incidence of transmission was greater when individuals were allowed to interact directly compared to cases where individuals only interacted indirectly via shared experience with the same silk. This suggests an important role of body contact or affiliative behavioural interactions in the transmission of cuticular bacteria. A recent experiment using freshwater snails demonstrated that bodily contact is a major determinant of the dispersal of a defensive symbiont from ‘donor’ to ‘receiver’ hosts, and similar (in terms of trait disassortativity) to the data presented here, the degree of transmission was greater when donor hosts were larger than their unexposed receivers, with the opposite being true if receivers were larger than donors [63]. This highlights the need to test the generality of trait disassortativity and transmission across many different host–symbiont systems.

(c) Future directions
For studies of wildlife disease, variation in the behavioural traits of infected and susceptible individuals is rarely explored in conjunction, despite ample evidence of its importance from the biomedical and human epidemiological fields. More comprehensive experiments should test how syndromes of behavioural traits can combine to influence the likelihood of individuals’ acquisition, colonization and

| Table 1. (a,b) Two nominal logistic regressions predicting the presence or absence of CNK02 on the cuticles of susceptible individuals after interacting with exposed individuals directly (i.e. co-habitating) or indirectly (i.e. only in contact with the same substrate 1 day later). (c) A GLMM predicting the estimated bacterial load collected from the cuticles of susceptible spiders after directly interacting with exposed spiders. Effects with significant *p*-values are denoted with an asterisk. |
|---|---|---|---|
| **(a) transmission after direct interaction** |  |  |  |
| difference in boldness between individuals | 1 | 7.58 | 0.006** |
| aggressiveness of exposed individual | 7 | 9.00 | 0.25 |
| aggressiveness of susceptible individual | 9 | 16.74 | 0.06 |
| body condition of susceptible individual | 1 | 5.70 | 0.02* |
| body condition of exposed individual | 1 | 0.51 | 0.47 |
| **(b) transmission after indirect interaction** |  |  |  |
| difference in boldness between individuals | 1 | 0.53 | 0.46 |
| aggressiveness of exposed individual | 6 | 10.58 | 0.10 |
| aggressiveness of susceptible individual | 7 | 8.63 | 0.28 |
| body condition of susceptible individual | 1 | 0.81 | 0.37 |
| body condition of exposed individual | 1 | 0.39 | 0.53 |
| **(c) estimated bacterial load transmitted (direct interaction)** |  |  |  |
| difference in boldness between individuals | 1 | 0.42 | 0.52 |
| aggressiveness of exposed individual | 6 | 9.30 | 0.16 |
| aggressiveness of susceptible individual | 5 | 6.95 | 0.22 |
| body condition of susceptible individual | 1 | 0.37 | 0.54 |
| Body condition of exposed individual | 1 | 3.33 | 0.07 |

*p < 0.05, **p < 0.01.
transmission of microbes (i.e. 'behavioural competence'; [64]). Future experiments could also incorporate how the composition of resident cuticular microbial communities can combine with behavioural traits to drive the likelihood of bacterial colonization and transmission [65,66]. More specifically in regards to this system, it would be informative to identify the role that males play in social contact networks, including sexual interactions, and their influence on bacterial transmission. Although colony sex ratios are strongly female-biased [39], males could have a high impact on bacterial transmission if they are highly interactive and/or susceptible to exposure. Further, it would be informative to observe matched pairs of exposed and susceptible individuals and record the frequency and nature of their interactions to determine if bacteria are transmitted simply by bodily contact or if other affiliative/aggressive interactions are linked to transmission. Our data here verify that the behavioural traits of exposed and susceptible individuals jointly influence the likelihood of interindividual transmission of cuticular bacteria, and represent a fundamental step towards understanding how individual traits can explain larger-scale epidemiological processes. These data thus reinforce the growing sentiment that comprehensive models of epidemiological processes must account for behavioural variation at the level of the individual.

Data accessibility. All data associated with this manuscript have been deposited at Dryad digital repository: http://dx.doi.org/10.5061/dryad.60an7.

Author’s contributions. C.N.K. transformed the bacteria under the advise of J.G.L. and performed statistical analyses; C.N.K., D.A.A. and M.J.Z. carried out experiments; N.P.W. and L.H. performed colony observations and network analyses; J.N.P. assisted in experimental design and manuscript preparation. All authors gave final approval for publication.

Competing interests. We declare that we have no competing interests.

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