

## Unraveling the disease consequences and mechanisms of modular structure in animal social networks

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Disease risk is a potential cost of group living. Although modular organization is thought to reduce this cost in animal societies, empirical evidence toward this hypothesis has been conflicting. We analyzed empirical social networks from 43 animal species to motivate our study of the epidemiological consequences of modular structure in animal societies. From these empirical studies, we identified the features of interaction patterns associated with network modularity and developed a theoretical network model to investigate when and how subdivisions in social networks influence disease dynamics. Contrary to prior work, we found that disease risk is largely unaffected by modular structure, although social networks beyond a modular threshold experience smaller disease burden and longer disease duration. Our results illustrate that the lowering of disease burden in highly modular social networks is driven by two mechanisms of modular organization: network fragmentation and subgroup cohesion. Highly fragmented social networks with cohesive subgroups are able to structurally trap infections within a few subgroups and also cause a structural delay to the spread of disease outbreaks. Finally, we show that network models incorporating modular structure are necessary only when prior knowledge suggests that interactions within the population are highly subdivided. Otherwise, null networks based on basic knowledge about group size and local contact heterogeneity may be sufficient when data-limited estimates of epidemic consequences are necessary. Overall, our work does not support the hypothesis that modular structure universally mitigates the disease impact of group living.

modularity | sociality | community structure | wildlife disease | infection

**S** ocial behavior is ubiquitous in vertebrates and arthropods. An evolutionary product of sociality in animal populations is group living. Group-living societies benefit from improved success in intergroup competition, protection from predation, mating opportunities, cooperative care of young, foraging success, and group defense. Conversely, group living is also associated with the costs of elevated disease burden due to a higher frequency of contact between hosts (1). Recently, it has been suggested that modular subdivisions in social interactions alleviate the association between group size and disease burden (2, 3).

Modular organization emerges in animal groups when subsets of conspecifics consistently interact with each other more often than they do with other individuals in the group, forming subgroups. Such modular structure is widespread across social networks in wildlife species, including relatively solitary species (4, 5), and can emerge from a combination of social behavior, demography, environmental, and landscape factors. However, theoretical investigations of the impact of modularity on spreading processes have produced mixed results in the past. Although some studies have suggested that modular subdivisions reduce transmission by creating structural bottlenecks (6–8), a few have found modular structure to increase outbreak size due to higher connectivity within subgroups (9, 10). This discrepancy in past research implies the need to consider realistic ranges of modularity (because not all animal social networks exhibit the high levels of modular subdivision typically investigated in past studies) and the need to mechanistically understand how modular structure in social networks affects the dynamics of disease spread.

Thus, we propose an empirically driven investigation of the structural features of interaction patterns that are associated with modular subdivisions in animal groups, including the ones that change the degree of network fragmentation (i.e., the number of subgroups), promote subgroup cohesion (i.e., individual preferences to interact with subgroup members), and drive variation in subgroup size (Fig. 1).

We have three objectives: (i) identify features of animal interaction patterns that increase the strength of modular organization in their social networks; (ii) explore the conditions under which modular organization influences the dynamics of disease spread and explain the mechanisms behind observed effects; and (iii) investigate whether knowledge of modular organization in animal social networks is necessary (and sufficient) to build accurate contact network models of disease spread. Overall, our study provides mechanistic insights on the context in which modular structure influences infectious disease spread in animal populations. We further suggest the use of appropriate null models when data-limited estimates of epidemic consequences in animal groups are necessary.

#### **Results and Discussion**

Modular Organization and Disease in Animal Social Networks. We obtained published (and publicly available) social networks for 69 groups across 43 animal species where edges can serve as realistic routes of infection spread. Newman modularity, Q, is a commonly used measure to estimate the strength of subdivision in networks (11). When Q = 0, the density of interactions within subgroups is equivalent to the density of interactions

### **Significance**

Modular organization in animal social networks is hypothesized to alleviate the cost of disease burden in group-living species. However, our analysis of empirical social networks of 43 animal species along with theoretical networks demonstrates that infectious disease spread is largely unaffected by the underlying modular organization, except when social networks are extremely subdivided. We show that high fragmentation and high subgroup cohesion, which are both associated with high modularity in social networks, induce structural delay and trapping of infections that spread through these networks, reducing disease burden. We validate our results using real animal social networks and recommend the use of appropriate null network models when data-limited estimates of epidemic consequences are necessary.

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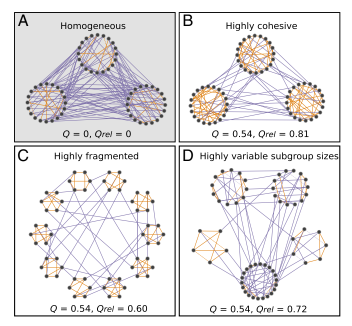


Fig. 1. Schematic representation of a homogeneous social network with no modular subdivision (A) and highly modular social networks with high cohesion within subgroups (B), high network fragmentation (C), and high subgroup size variation (D). Each network has 60 nodes. Orange edges indicate interactions within a subgroup, and purple edges are between-subgroup interactions. Q is Newman modularity (11) and  $Q_{rel}$  is relative modularity.

between subgroups. Higher values of Newman modularity indicate stronger subdivisions of social networks. However, using Newman modularity to compare modular subdivision across networks is problematic because the measure is inherently correlated to the size and number of links in the network (12).

This is because, in principle, we expect the maximum modularity  $Q_{max}$  of all networks to be equal to one. However, for smaller networks,  $Q_{max}$  tends to be a much smaller value (Fig. 2A). We therefore estimated relative modularity (obtained by normalizing Q by  $Q_{max}$ ) to facilitate comparison of modular subdivision strength across different animal groups. Animal social networks in our database ranged from homogeneous ( $Q_{rel} = 0$ ) to highly subdivided structure (Fig. 2A). Animal interactions are dynamic in nature, and therefore social networks of the same set of individuals can also fluctuate between being relatively homogeneous to highly modular over time (Fig. 2B).

To gain intuition about how modular organization in animal social networks influences epidemiological outcomes, we first generated homogeneous null networks for each empirical animal social network in our database. In homogeneous null networks, higher-order organization in social networks (such as modular structure, clustering coefficient, cliquishness, and degree homophily) are randomized, weakening relative modularity (SI Appendix, Fig. S1), but individual contact heterogeneity (degree distribution) of the empirical network is preserved. We next performed susceptible-infected-recovered (SIR) disease simulations of a moderately transmissible infectious disease (basic reproduction number,  $R_0 = 1.2$ ) through these empirical networks and their homogeneous null network counterparts. Based on previous work suggesting a protective effect of modular structure on disease risk (3, 6), we expected a lower disease burden (measured as the proportion of population infected) in empirical networks than in homogeneous null networks. Reduced disease burden, however, was apparent only in social networks with  $Q_{rel} > 0.6$  for moderately spreading infectious disease (Fig. 2C and SI Appendix, Fig. S2). Additionally, none of the empirical networks demonstrated a major reduction in disease burden for a highly transmissible infectious disease ( $R_0 = 4.8$ , SI Appendix, Fig. S3). This implies that the protective effect of modular subdivisions is realized only at high values of relative modularity, but the presence of a threshold depends on pathogen transmissibility (SI Appendix, Fig. S3).

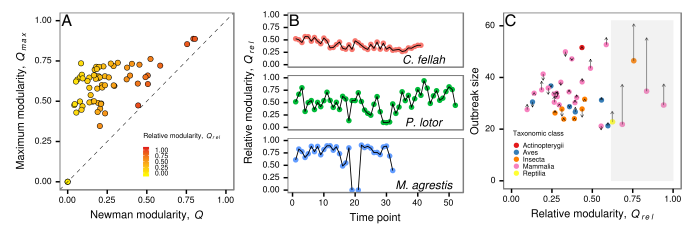


Fig. 2. Animal social networks with modular subdivisions. (A) Values of Newman modularity, Q, and maximum possible modularity given the network configuration,  $Q_{max}$ , across all animal groups in the database. Point color represents the relative modularity estimated as the ratio of Q over  $Q_{max}$ . The dashed line represents  $Q = Q_{max}$  and the maximum value of  $Q_{rel} = 1$ . (B) Relative modularity values of dynamic interactions in ants (Camponotus fellah), raccoons (Procyon lotor), and field voles (Microtus agrestis), where time points represent consecutive days, weeks, and trapping sessions, respectively. (C) Comparisons between real (filled points) and their homogeneous null networks (tips of arrows) with respect to the percentage of infected individuals (outbreak size) due to an outbreak with basic reproduction number,  $R_0 = 1.2$ . Point color corresponds to the taxonomic class of the animal group. Social networks with nonsignificant modular subdivision (as indicated by t test analysis) have been excluded in C. The generated homogeneous null networks preserve only the local heterogeneity of contacts among individuals; the arrows therefore indicate the change in direction and magnitude of outbreak size under the scenario in which all higher-order structural complexities (including modular subdivisions) are removed from animal social networks. The shaded area represents the region where empirical networks tend to experience reduced outbreak size (at  $Q_{rel} > 0.6$ ; SI Appendix, Fig. S2). The networks that experienced lower disease burden included social networks of raccoons (P. lotor), elephant seals (Mirounga angustirostris), ants (Camponotus pennsylvanicus), bottlenose dolphins (Tursiops truncatus), and Australian sleepy lizards (Tiliqua rugosa). In A and C, we used a randomly selected network snapshot for groups with temporal interaction data.

#### Determinants of Modular Organization in Animal Social Networks.

Fig. 2C suggests that the epidemiological consequences of modular structure are context-dependent, depending on the strength of relative modularity, and only above a certain threshold. Additionally, the extent of outbreak size reduction above the modularity threshold was fairly inconsistent across networks. This could be due to varying features of interaction patterns driving modular subdivision in animal social networks. We therefore first turned our attention to some of these network features, including the number of individuals present in the social network (network size); (log of the) number of subgroups in the social network (network fragmentation); preferential association with own subgroup (subgroup cohesion, measured as the proportion of total contacts that occur within subgroups); variation in subgroup cohesion across the network; average and variation in subgroup sizes; average and variation in number of individual contacts (degree); and variation in contacts among subgroups (subgroup degree variation). For groups that had multiple temporal snapshots of a social network, we randomly selected one snapshot network for the analysis to avoid intragroup correlation in network metrics (and we analyze these temporal networks separately below).

The mixed-effects beta-regression model identified three network features driving modular organization in animal social networks: subgroup cohesion, network fragmentation, and subgroup size variation (Table 1). Of the three factors, subgroup cohesiveness and network fragmentation had strong positive effects on modular organization. Cohesiveness can be a direct result of strong social bonds due to similar nutrient requirements [e.g., in spotted hyenas (13)], time spent foraging together, or matrilineal descent [e.g., in African elephants (14)]. High network fragmentation, conversely, can be caused by constraints on resource density, resource distribution, or high intragroup competition [e.g., in brown spider monkeys (15)]. In contrast to network fragmentation and subgroup cohesion, we found variation in subgroup sizes to have a weak negative effect on network modularity. Fluid subgroup sizes are common in many social species (e.g., in spotted hyenas, elks, chimpanzees, bottlenose dolphins, and African lions) and can be brought about by changes in resource availability, intragroup aggression, demographic factors (such as sex ratio), dominance hierarchy, and female reproductive state (13, 15, 16).

Thus, far, we have considered only a single snapshot of interactions for each animal group in our database for consistent comparison across species with unequal sampling efforts. However, dynamic interactions between group members in fission–fusion species, species with nomadic individuals, or species with seasonally driven social interactions can change the level of network

Table 1. Multivariable analysis on determinants of modular organization ( $Q_{rel}$ ) across groups of different species

Explanatory variable	Effect size	95% confidence intervals	
Intercept	-0.42	−0.59 to −0.25	
Network size	-0.07	-0.17 to 0.03	
Network fragmentation*	0.66	0.51 to 0.82	
Subgroup cohesion*	1.14	1.03 to 1.25	
Subgroup cohesion variation	-0.03	-0.14 to 0.08	
Subgroup size average	_	_	
Subgroup size variation*	<b>-0.13</b>	−0.26 to −0.01	
Individual degree average	_	_	
Individual degree variation	0.07	0.04 to 0.18	
Subgroup degree variation	0.01	-0.08 to 0.10	
Random effects	Variance estimate ( $\sigma^2$ )		
Taxonomic order	0.066		
Sociality	< 0.001		

Asterisks and bold text indicate significance. Explanatory variables with VIF > 5 were dropped from the model, and therefore their effect sizes were not estimated.

modularity over time (Fig. 2B). We therefore repeated the analysis by considering dynamic interactions for a few animal groups for which such data are available. Consistent with our previous analysis of aggregated interactions, we found that animals groups become increasingly modular with higher subgroup cohesion or network fragmentation (SI Appendix, Table S1). In addition, variation in subgroup size and variation in contacts among subgroups had a weak negative correlation on relative modularity within animal groups.

The Impact of Modularity on Dynamics of Disease Spread. We have shown that network fragmentation and cohesion within subgroups are associated with modular structure in animal social networks. Do the two factors also determine the disease consequences of modular subdivisions in animal social networks? To answer this question, we used synthetic modular networks generated using our model proposed in ref. 17. The model generates networks with a tunable strength of modularity, while keeping other higher-order network features (such as degree homophily, clustering coefficient, and path length) close to the configuration realized in homogeneous networks (17). Because several structural features of networks can influence disease spread, this approach is appropriate to isolate the epidemiological consequences of modular organization in social networks. Performing SIR disease simulations on synthetic modular networks, we found that the magnitude of outbreak size experienced by low modular networks ( $Q_{rel} < 0.6$ ) was similar to that of homogeneous networks ( $Q_{rel} = 0$ ). Outbreak size was reduced only in extremely modular networks for moderately transmissible pathogens (Fig. 3 A and B). We corroborated our finding that highly (but not low) modular networks inhibit global parasite transmission by using empirical data to investigate the spread of protozoan parasites *Babesia* in the social networks of field voles (18) (SI Appendix, Fig. S11). In addition, we found no empirical evidence of structural trapping of a gastrointestinal parasitic infection in the low modular social network of brown spider monkeys (19) (SI Appendix, Fig. S12).

We next quantified the robustness of modular social networks toward disease spread in two ways. First, we estimated the minimum level of pathogen contagiousness below which there is no risk of a large outbreak (epidemic threshold; solid line in Fig. 3A). We found networks with  $Q_{rel} > 0.6$  to have a higher epidemic threshold compared with homogeneous and low modular networks, implying that highly modular networks alleviate the risk of large disease outbreaks for low contagious pathogens. Second, for pathogen transmissibility beyond the epidemic threshold, we calculated the minimum level of relative modularity in which networks experience at least a 10% reduction in outbreak size compared with homogeneous ( $Q_{rel} = 0$ ) networks (modularity threshold; solid line in Fig. 3B). We found that networks with  $Q_{rel} \ge 0.5$  mitigated disease spread for moderately contagious (transmissibility = 0.08-0.16) but not highly contagious pathogens (transmissibility > 0.16). Highly modular networks  $(Q_{rel} > 0.8)$  had a striking impact on disease transmissionoutbreak sizes of moderately transmissible pathogens were reduced up to 58% compared with homogeneous networks. Such highly modular networks, although rare, do occur in some host interaction networks. In our database, three animal species demonstrated  $Q_{rel}$  values of >0.8—raccoons (P. lotor), field voles (M. agrestis), and northern elephant seals (M. angustirostris).

The question then is why do not all levels of modularity reduce disease burden in animal social networks? The relationship between disease burden and modularity depends on the tradeoff between local disease transmission within subgroups and global disease spread (Fig. 3C). Compared with homogeneous networks (with  $Q_{rel}=0$ ), modular networks experience higher local transmission due to a high density of contacts within subgroups; conversely, low intersubgroup contacts in these networks reduce global disease transmission (Fig. 3C). This tradeoff balances the overall disease spread, and thus modular structure,

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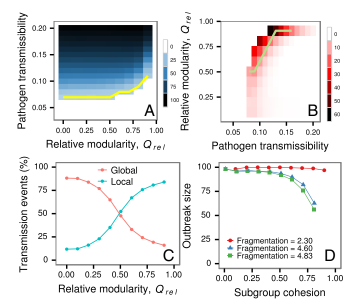


Fig. 3. Overall disease implications of modular subdivisions in synthetic modular networks. (A) Average outbreak size, measured as the percentage of infected individuals, over increasing subdivided social networks and pathogen transmissibility. Outbreak size values have been normalized to the maximum observed outbreak. The solid line indicates epidemic threshold namely, the threshold value of pathogen contagiousness below which there is no risk of a large outbreak (> 10% outbreak size; Materials and Methods). (B) Epidemic robustness of networks with increasing value of relative modularity, measured as the percentage reduction in outbreak size compared with outbreak size experienced by homogeneous ( $Q_{rel} = 0$ ) networks. The solid line indicates the modularity threshold, where networks experience at least a 10% reduction in outbreak size. (C) Infection transmission events, expressed as the percentage of total outbreak size, within subgroups (local) and between subgroups (global); pathogen transmissibility = 0.18. (D) Disease implications of modular subdivisions as a function of subgroup cohesion and network fragmentation (measured as the log-number of subgroups present in the network).

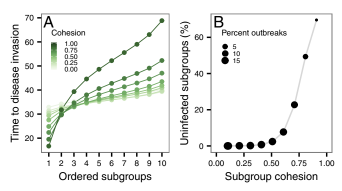
under most conditions, does not lower disease burden. The conditions under which modularity does reduce disease burden of social networks is largely determined by the two factors underlying network modularity—network fragmentation and subgroup cohesion (Table 1). Synthetic networks that possess both high subgroup cohesion and high network fragmentation experience lower outbreak size compared with homogeneous and low modular social networks (Fig. 3C and SI Appendix, Fig. S6). In addition, high variation in subgroup sizes along with high subgroup cohesion and fragmentation further reduce outbreak size and increase outbreak duration in social networks (SI Appendix, Fig. S7). Disease consequences in real animal social networks, however, tend to be driven by subgroup cohesion because they typically exhibit a much lower range of fragmentation and subgroup size variation compared with subgroup cohesion (SI Appendix, Figs. S4 and S5).

Mechanisms That Drive the Impact of High Modular Structure on Disease Spread: Structural Delay and Trapping. We next examined the mechanisms by which highly fragmented social networks with cohesive subgroups experience reduced disease burden. The presence of only a few interactions between highly cohesive subgroups increased the amount of time it took for an infection to spread from one subgroup to another (the structural delay effect; Fig. 4A), causing longer disease outbreaks. When networks were fragmented into highly cohesive subgroups, the structural delay imposed by cohesion caused infection to be localized to a small proportion of subgroups before dying out (Fig. 4B). Popularly known as structural trapping (20), this effect also reduces the overall likelihood of a major outbreak by a novel

infection in modular social networks. The structural trapping effect (albeit weak) has been observed in a field study of pneumonia transmission in highly subdivided networks of bighorn lambs (SI Appendix, Fig. S13) (21) and others (SI Appendix, Table S2). We also found empirical evidence of structural delay in the spread of mycoplasma in the low-modular network, but highly cohesive subgroups, of house finches described in ref. 22 (SI Appendix, Fig. S14). A high probability of extinction associated with the structural trapping effect has been reported in theoretical studies of disease spread in spatially structured metapopulation models (7, 8). We note that the magnitude of structural trapping and delay depends on the contagiousness of the infection—moderately transmissible pathogens spread slowly through populations and are therefore able to perceive modular structures present in networks. For rapidly spreading pathogens, highly fragmented and cohesive networks do not contain, but delay, the spread of disease to other subgroups, creating only a structural delay effect.

Is Knowledge of Modular Structure Necessary (and Sufficient) to Predict Disease Outcomes? Our investigations of disease consequences on synthetic networks suggest that only modular subdivision beyond a certain threshold can reduce disease burden of moderately transmissible pathogens. Other topological features of the network—and, in particular, heterogeneity in host contact (23)—are, however, bound to confound the effect of modularity on infectious disease spread. Our work thus far assumes high variation in contacts among individuals (24), although factors including sociality (25) and nonpersistent space use (26) are known to reduce variation in individual contact rate. Does individual contact heterogeneity then affect the way modular organization influences disease spread? We found that for social networks with low contact heterogeneity, high modularity brings about a greater reduction in disease burden compared with networks with high contact heterogeneity (SI Appendix, Fig. S8). These results imply that the local contact patterns should also be considered when estimating the reduction of disease burden in extremely subdivided social networks.

Although modular subdivision is a common feature observed in many animal groups, it is often correlated with other higher-order network structures, such as clustering coefficient and degree homophily (27). In empirical networks, it is therefore important to ask whether modular organization has any epidemiological consequence above and beyond these network properties, after accounting for contact heterogeneity. To answer this



**Fig. 4.** Mechanisms behind the effect of modular organization on disease spread of a moderately transmissible pathogen (= 0.1). (A) Structural delay effect in a moderately fragmented network (fragmentation = 2.30): We considered the time to disease invasion for a subgroup as the number of time steps it takes for at least 2% of individuals to acquire infection. Subgroups at the x axis are sorted according to the increasing disease invasion time (B) Structural trapping effect in a highly fragmented network (fragmentation = 4.83): High fragmentation and subgroup cohesion localize infection to a small proportion of subgroups in the social network. Structural trapping also increases the likelihood of stochastic extinction of disease outbreaks.

question, we performed disease simulations through 19 animal social networks in our database and their modular null networks. Modular null networks preserve the modularity level and degree distribution of their empirical network counterparts, but randomize all other network structure. We expected a high error rate in outbreak size prediction from modular null networks if other structural features mask the effect of modular subdivisions. We also compared the performance of modular null networks with that of homogeneous null networks, where all network structure, except degree distribution, was randomized (similar to the networks used in Fig. 2C). We found low (but not zero) percentage error in outbreak size predictions from modular null networks (Fig. 5 and SI Appendix, Fig. S9). Homogeneous null networks, however, performed equally well in predicting disease outcome for most animal social networks, except when relative modularity of empirical networks was high ( $Q_{rel} > 0.6$ ). We note that this  $Q_{rel}$  value is similar to the threshold identified above (Fig. 2C and SI Appendix, Fig. S2). For moderately transmissible infections, animal social networks experience low outbreak size at this level of modular structure due to the structural trapping effect. Homogeneous null networks do not preserve the modular structure of the empirical networks and, therefore, overestimate the true disease burden. In contrast to moderately spreading infectious diseases, slowly  $(R_0 < 1)$  and rapidly spreading infections are unaffected by higher-order complexities of the networks, and therefore outbreak size estimates of homogeneous and modular null networks are similar across all levels of relative modularity (SI Appendix, Fig. S10). To empirically confirm the results of our disease simulations, we compared the performance of modular and homogenous null networks to predict outbreak size of two Salmonella enterica strains in the Australian sleepy lizards, T. rugosa, using data described in ref. 28 (SI Appendix, Fig. S15). Congruent to the theoretical predictions, homogeneous and modular networks produced identical and accurate outbreak size predictions for the low-transmissibility strain. For the moderately transmissible strain, conversely, modular null networks performed better in estimating the true disease burden.

#### **Conclusions**

While modular social organization has been postulated to alleviate the disease burden of group-living species, past evidence sup-

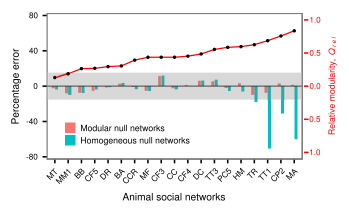


Fig. 5. Percentage error in outbreak size predictions using modular and homogeneous null networks for 19 animal social networks due to an outbreak with  $R_0=1.2$ . Percentage error is calculated as  $(S_{emp}-S_{null})/S_{emp}\times 100$ , where S= outbreak size, emp= empirical network, and null are modular or homogeneous null networks of the empirical network. The social networks are ordered according to the increasing value of relative modularity (red solid curve, secondary y axis). The shaded region indicates the range of percentage error values <15%. BA, B rachyteles B rachnoides; BB, B rach B roc, C cercopithecus C campbelli; C CCR, C rocuta C crocuta; C CF, C fellah; C CP, C pennsylvanicus; C DC, dairy cattle; C DR, C rocuta C r

porting this hypothesis has been equivocal. More importantly, the features of interaction patterns that cause animal social networks to be subdivided have not been identified before. In this study, we resolved the ambiguity of past research on disease consequences of network modularity in three ways. First, we introduced a normalized version of Newman modularity called relative modularity  $(Q_{rel})$  to allow comparison across networks of different sizes and identified two distinct mechanisms that can lead to modular structure. Second, we systematically studied the epidemiological consequences of modular organization by using a model (17) that allows for the mechanistic generation of synthetic modular networks and null networks (of empirical animal social networks), while minimizing the presence of higher-order network properties. Third, we combined published social networks across 43 animal species with this theoretical network model and biologically realistic disease simulation to derive insights into disease consequences of mechanisms of modular organization in animal societies.

It has been hypothesized that subgroups in animal social networks reduce the disease costs of group living (2, 3). Contrary to this hypothesis, our study shows that disease burden is largely unaffected by modular subdivisions in animal groups, except when social networks are extremely fragmented and have high cohesion within subgroups. High subgroup cohesion and fragmentation in networks structurally trap moderately transmissible infections to a few subgroups and therefore reduce the overall disease burden. We note that, although overall disease burden is unaffected at most levels of modularity, the design of control strategy will vary for different levels of network modularity. An effective intervention should aim to minimize global spread of infection for low network modularity, but reduce local transmission at high network modularity. Because our analysis is based on a systematic investigation of contagion spread in synthetic modular networks, the results of this study are potentially applicable across a wide range of systems, including spreading processes at higher ecological scales (SI Appendix, Table S2), infectious disease spread in human interaction networks and transportation networks, and perturbation spread in metabolic and ecological networks, as well as spread of information through communication networks.

Because this study involved meta-analysis of social networks across a broad range of taxonomic groups, we made a number of simplifying assumptions that should be considered before extending the results to a specific system. First, we filtered the weights assigned to social interactions between individuals to construct unweighted networks. This was because the impact of weights (whether they represent frequency, duration, or intensity of contact) on transmission potential is generally unclear and usually context-dependent. We advise leveraging transmission studies in captive populations or historical disease spread data, where available, to identify the appropriate weighting criterion of contact networks. Second, we assume that the aggregated static networks and temporal "snapshots" of dynamic networks in our database coincide with the temporal scale relevant to infection spread in our disease simulations. Consideration of the relevant time interval of animal interactions relative to the transmission mode and infectious period of the pathogen, however, is crucial to developing accurate network models of disease spread (7, 29). Because many animal interactions can be infrequent or intermittent, particularly between subgroups, aggregating interactions over a small time window may ignore these fleeting contacts and overestimate the modularity of social networks. Conversely, pooling interactions over large time windows may amplify the role of temporary contact, and, consequently, the social subgroups might appear more connected. In addition, chronic infections with a long infectious period will be relatively unaffected by high network modularity if the time scale of network fluctuations is shorter than the average infectious period. Acute infections with short infectious periods, however, will be highly sensitive to the state of network modularity at the time of disease outbreaks.

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This study suggests the presence of a high modularity threshold, above which social networks experience reduced outbreak size. However, we caution against the use of a single modularity threshold that is applicable to all systems. Multiple behavioral, ecological, and epidemiological factors-including pathogen infectiousness (Fig. 3B), infectious period, local contact heterogeneity, seasonality of contacts, and transmission, coupled with recruitment of susceptible through births—can influence the  $Q_{rel}$  threshold and the extent to which network fragmentation and subgroup cohesiveness structurally trap and delay infection spread. Therefore, to assess the disease implications of modular structure observed in specific social groups, we recommend the use of null networks, as presented in this study. Comparison of disease consequences on empirical network data with those on a hierarchy of null networks clarifies whether modular structure is important. If natural history and limited observations suggest a species' network will be highly modular, then models that incorporate modularity should be used; otherwise, homogeneous null models (based on basic knowledge about network size and local heterogeneity) may be sufficient when datalimited estimates of epidemic consequences are necessary.

#### **Materials and Methods**

Extended methods are provided in SI Appendix, SI Text.

Measuring Modularity. We used modularity (Q) proposed by Newman (11) to measure the strength of modular organization in networks. Modularity can be defined as  $Q = \sum_{k=1}^{K} \left[ \frac{l_k^w}{L} - \left( \frac{l_k}{L} \right)^2 \right]$ ), where  $L_k$  is the total number of edges in a subgroup k, of which  $L_k^w$  are the edges within the subgroup, and L is the number of total edges in the network. Community structure, or the number and composition of subgroups, for each animal social network was estimated by using the Louvain method (30). The highest possible modularity in a network ( $Q_{max}$ ) is achieved when all individuals in a subgroup konly interact with each other and no edges are present between subgroups (i.e., subgroups are disjointed). In other words, Qmax of a network is when  $L_k^W = L_k$ , and can be written as  $Q_{max} = \sum_{k=1}^K \frac{L_k}{L} \left(1 - \frac{L_k}{L}\right)$ . We measured the relative modularity of networks as  $Q_{rel} = \frac{Q}{Q_{max}}$ .

Mechanisms of Modular Organization. To examine the relative contribution of factors such as network size, network fragmentation, average subgroup

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size, etc., toward network modularity, we ran a mixed-effects beta regression model using the almmADMB package (Version 0.8.3.3) in R (Version 3.2.3). We treated the species nested within the taxonomic order of the animal group as a random effect in the model to account for any correlations in network metrics. However, the variance accounted for by the nesting of species was negligible ( $\sigma^2 \approx 0$ ), and we therefore considered only taxonomic order as the random effect in the analysis. We also accounted for differences between the social organization of different species by defining three broad levels of sociality—relative solitary, social, and fission-fusion and included sociality as a random effect in the model.

Generation of Synthetic Modular and Null Networks. We used the network generator described in ref. 17 to generate networks of a specified modularity and degree sequence, while keeping other network properties close to homogeneous null networks. For Figs. 3 and 4, we generated synthetic modular networks with 10,000 nodes, with an exponential degree distribution with a mean degree of 10. Two types of null networks were generated for each animal social network. Modular null networks were created by randomizing within- and between-subgroup connections. The modular null networks therefore had identical modular configuration and degree sequence as empirical networks, but were random with respect to other higher-order network properties. We generated homogeneous null networks by performing simultaneous edge swaps over within- and between-subgroup connections, which preserves the local contact heterogeneity, but randomizes all higherorder features of the network, including network modularity.

Disease Simulations. We performed Monte Carlo simulations of a discretetime SIR model of disease spread. Because we were interested only in major outbreaks, we considered only those simulations in our calculations where at least 10% of the population acquired infection. Transmission occurred according to a pathogen transmissibility, defined as the probability of transmission from an infected to susceptible host during the period when the infected host is infectious. The recovery probability was assumed to be con-

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# SI Appendix for: Unraveling the disease consequences and mechanisms of modular structure in animal social networks

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#### SI Materials and Methods

Animal social networks. We obtained previously published (and publicly available) social networks of 69 groups across 43 animal species. The networks were selected based on the criterion that edge connections can serve as actual routes of infectious disease spread. For groups that had multiple temporal snapshots of social network, we randomly selected one network for the analysis to avoid intra-group correlation in network metrics. Few networks had weighted edges. To enable consistent comparison across all networks, we transformed the edges of these networks to binary edges by employing an edge filtering approach (1). To this end, edges with weights less than 50<sup>th</sup> percentile of edge weights were removed from the network, and the remaining edges were assigned an edge weight of one.

**Measure of relative modularity.** We used modularity (Q) proposed by Newman (2, 3) to measure the strength of modular organization in networks. Modularity can be defined as:

$$Q = \sum_{k=1}^{K} (e_{kk} - a_k^2)$$
 [1]

where  $e_{kk}$  denotes the fraction of edges within the subgroup k and  $a_k$  is the fraction of the total edges present within the subgroup. If  $\overline{d^k}$  is the average degree of individuals in subgroup k,  $\overline{d^k_w}$  is the average within subgroup degree and  $s_k$  is the total number of individuals in the subgroup, then the equation 1 can be rewritten as:

$$Q = \sum_{k=1}^{K} \left[ \frac{\overline{d_w^k} s_k}{\overline{d}n} - \left( \frac{\overline{d^k} s_k}{\overline{d}n} \right)^2 \right]$$
 [2]

where  $\overline{d}$  is the average degree of the network, and the total number of individuals and subgroups in the network are n and K, respectively.

Let  $L_k$  be the total edges of individuals of a subgroup k out of which  $L_k^w$  are the edges present within the subgroup. Equation (2) can then be further reduced to:

$$Q = \sum_{k=1}^{K} \left[ \frac{L_k^w}{L} - \left( \frac{L_k}{L} \right)^2 \right]$$
 [3]

where L is the total edges present in the network.

We estimated Newman modularity, Q, for the largest connected component of each animal social network using the Louvain method as described in (4).

The highest possible modularity in a network is achieved when all individuals in a subgroup k only interact with each other and no edges are present between subgroups (i.e., subgroups are disjointed). In other words, Qmax of a network is when  $L_k^w = L_k$ . Equation S3 can be therefore written as

$$Q_{max} = \sum_{k=1}^{K} \left[ \frac{L_k}{L} - \left( \frac{L_k}{L} \right)^2 \right]$$
 [4]

A network is more modular if its observed Q value is closer to the maximum possible modularity. We thus normalize the strength of modular organization with respect to its  $Q_{max}$  by computing relative modularity of networks as  $Q_{rel} = \frac{Q}{Q_{max}}$ .

Mechanisms of modular organization. We test the role of several features of social organization in animal societies that may contribute towards the observed magnitude of relative modularity. These features include network size (n); (log of the) number of subgroups in the social network (network fragmentation); preferential association with own subgroup (subgroup cohesion, measured as the proportion of total contacts that occur within subgroups); variation in subgroup cohesion across the network; average and variation in subgroup sizes; average and variation in number of individual contacts (degree); and variation in contacts among subgroups (subgroup degree variation). We note that this is not an exhaustive list but includes factors that we believe can be important in the context of animal social networks. Average contacts of subgroups was not included in the analysis because mathematically it is equal to the average degree of individuals.

To examine the relative contribution of these factors towards network modularity, we ran the following mixed effects beta regression model using glmmADMB package (version 0.8.3.3) in R (version 3.2.3). All predictors were centered and scaled to unit variances to assign same prior importance to each predictor in the analysis (5). The distributions of network size and number of subgroups were skewed to the right and therefore were natural log-transformed for further analyses. For all the variance predictors, we estimated coefficient of variation ( $CV = \sigma/\mu$ ) to avoid possible correlation with their average predictor counterparts. To avoid multicollinearity, we estimated the variance inflation factor (VIF) implemented in the package 'car' in R, and removed predictors with VIF >5 (6). We treated the species nested within the order of the social group and sociality as random effects in the model.

Generation of null networks. Two types of null networks were generated for each animal social network. Modular null networks were created in two steps: first, within-subgroup connections were randomized and the second step involved randomization of between-subgroup connections. Edges were randomized using the double-edge swap operation in the NetworkX package (7). The modular null networks, therefore, had the same modular subdivision and degree sequence as the empirical networks, but were random with respect to other higher-order network properties. We generated homogeneous null networks by performing simultaneous edge swaps over

within- and between-subgroup connections, which preserves the local contact heterogeneity but randomizes all higher-order features of the network including network modularity (8).

Generation of synthetic modular networks. We generated synthetic modular networks using modular random network generator described in (8). This model allows generation of modular networks by varying the level of network fragmentation or subgroup cohesion while keeping other network properties (such as degree homophily, clustering coefficient) close to homogeneous null network with no modular structure. In this study we generated modular networks with 10,000 nodes. The strength of modular organization was varied by adjusting either subgroup cohesion or network fragmentation. As the total number of individuals was fixed across all generated synthetic networks. increasing network fragmentation (i.e., the number of subgroups) also coincided with decreasing the average subgroup size in the network. Unless otherwise noted, we assumed local contact heterogeneity in social groups to follow an exponential degree distribution with mean contact (degree) of 10.

**Disease simulations.** We performed Monte-Carlo simulations of discrete-time susceptible-infected-recovered (SIR) model of disease spread. Each disease simulation was initiated by infecting a randomly chosen individual in the social group. At subsequent time steps every infected individual(s) in the population could either transmit infection to a susceptible neighbor with probability parameter  $\beta$  or recover with probability parameter  $\gamma$ . The disease simulation was terminated when there were no remaining infected nodes in the network. As we were interested only in major outbreaks, we considered only those simulations in our calculations where at least 10% individuals in the network acquired infection. Pathogen contagiousness was measured in terms of infection transmissibility. Transmissibility was defined as the probability of pathogen transmission from an infected to susceptible host during the period when the host is infectious. Assuming the individual's recovery and infection transmission to be a Poisson process, transmissibility was calculated as:  $T = \frac{\beta}{\beta + \gamma}$ .

Disease simulations in Figure 2C and 5: To compare epidemiological consequences of empirical networks, we simulated infectious disease spread with basic reproduction number,  $R_0 = 1.2$ . Basic reproduction number is defined as the average number of secondary infections caused by a single infected host in a completely susceptible population. Following ref. (9), we estimate transmissibility of pathogen corresponding as:

$$T = R_0 \left( \frac{\langle k \rangle}{\langle k^2 \rangle - \langle k \rangle} \right)$$
 [5]

where  $\langle k \rangle$  and  $\langle k^2 \rangle$  are the mean degree and mean square degree of the networks, respectively. Since empirical networks had varying values of  $\langle k \rangle$  and  $\langle k^2 \rangle$ , we obtained a different value of pathogen transmissibility corresponding to a  $R_0=1.2$  for each empirical network. For consistent comparison, disease simulations on null networks were performed using the same value of pathogen transmissibility that was estimated for their empirical network counterpart using equation 5.

Threshold analysis. We identified the threshold of pathogen contagiousness below which there is a minimal risk of large outbreak in modular networks using the formula proposed in

ref. (10). Specifically, the threshold is estimated numerically as:

$$\Delta = \frac{\sqrt{\langle \rho^2 \rangle - \langle \rho \rangle^2}}{\langle \rho \rangle}$$
 [6]

where  $\rho$  denotes the outbreak sizes of Monte-Carlo disease simulations, and  $\Delta$  is a variability measure. The epidemic threshold is estimated by performing disease simulations over a wide range of pathogen transmissibility and estimating the corresponding  $\Delta$  measure. The pathogen transmissibility where  $\Delta$  peaks is considered to be the epidemic threshold (10).

Table S1. Multivariable analysis on determinants of modular organization in the dynamic networks of four species (*C. fellah*, *C. pennsylvanicus*, *P. lotor*, and *M. agrestis*)

Explanatory variable	Effect size	95% confidence intervals
Intercept	0.05	0.03 - 0.07
Network size	-0.03	-0.05 - 0.00
Network fragmentation*	0.48	0.44 - 0.52
Subgroup cohesion*	0.87	0.85 - 0.90
Subgroup cohesion variation	_	_
Subgroup size average	_	_
Subgroup size variation*	-0.10	-0.120.08
Individual degree average	_	_
Individual degree variation	-0.02	-0.06 - 0.02
Subgroup degree variation*	-0.04	-0.07 — -0.02
Random effects	Variance estimate ( $\sigma^2$ )	
Group identification	0.0002	
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Response variable is the relative modularity,  $Q_{rel}$ , of the social networks. Asterisks and bold text indicate significance. We treated the social group identity of each network as the random effect. Explanatory variables with VIF>5 were dropped from the model, and therefore their effect sizes therefore were not estimated.

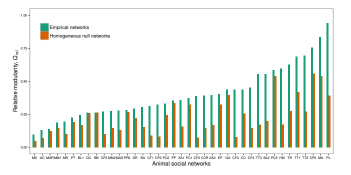
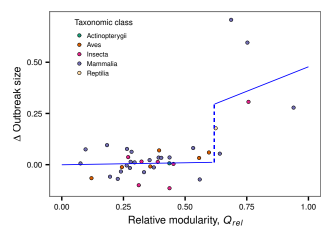
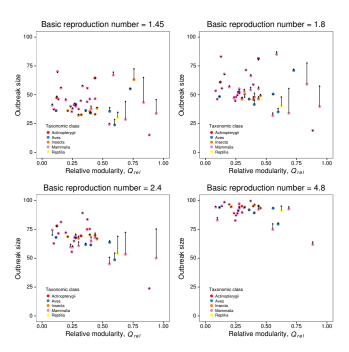


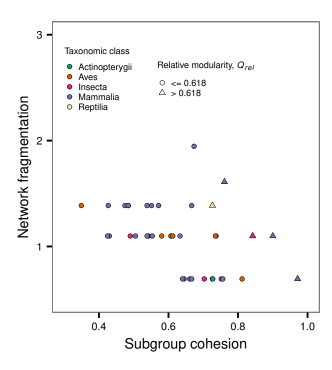
Fig. S1. Empirical animal social networks and the corresponding homogeneous null networks included in Fig.2C. Animal social network abbreviations: AC = Acanthiza spp., CC = Cercopithecus campbelli, CCR = Crocuta crocuta, CF = Camponotus fellah, CP = Camponotus pennsylvanicus, BB= Bison bison, BL = Branta leucopsis, EP = Erythrocebus patas, GA = Gasterosteus aculeatus, HM = Haemorhous mexicanus, MA = Mirounga angustirostris, MAR = Macaca arctoides , MM = Macaca mulatta, MS = Myotis sodalis, OC = Ovis canadensis, PC = Papio cynocephalus , PL = Procyon lotor, PP = Pan paniscus, PT = Pan troglodytes, TR = Tiliqua rugosa, TT = Tursiops truncatus, ZA = Zonotrichia atricapilla. Numbers next to the species acronym indicate a separate social group.



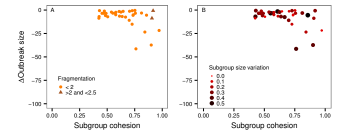
**Fig. S2.** Threshold value of relative modularity above which the empirical networks tend to experience lower outbreak sizes as compared to homogeneous null networks in Fig.2C. The breakpoint is at  $Q_{rel} \approx 0.618$ .  $\Delta$  Outbreak size is the relative outbreak size and is calculated as  $\frac{S_{null} - S_{emp}}{S_{emp}}$  where S = average outbreak size, emp = empirical network, null = homogeneous null networks. We recursively fit a piecewise linear mixed regression model with a sequence of increasing breakpoint values. The optimum breakpoint value was selected from the model with minimum deviance. To take into account any correlation within the taxonomic groups, we used the species nested with genus and taxonomic order of the groups as a random effect in the model. Sociality of the animal groups analyzed (relative solitary, social, and fission-fusion) was also included as a random effect.



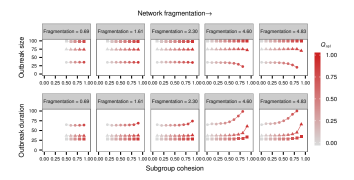
**Fig. S3.** Extended figure of Fig.2C Comparisons between real (filled points) and their homogeneous null networks (tip of arrows) with respect to the percentage of infected individuals (outbreak size) due to an outbreak with basic reproduction number,  $R_0$  equal to 1.45, 1.8, 2.4 and 4.8. Point color corresponds to the taxonomic class of the animal group. Social networks with non-significant modular subdivision (as indicated by t test analysis) have been excluded. The generated homogeneous null networks preserve only the local heterogeneity of contacts among individuals; the arrows therefore indicate the change in direction and magnitude of outbreak size under the scenario where all higher-order structural complexities (including modular subdivisions) are removed from animal social networks.



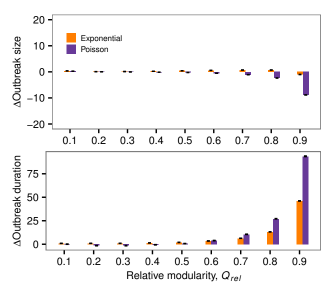
**Fig. S4.** The values of network fragmentation and subgroup cohesion present in empirical animal social networks. Network fragmentation is the log-transformed value of the total subgroups present in the social network. Circles are the networks with  $Q_{rel}$  <0.618 (the threshold value shown in Fig.2c of the main text and SI Appendix, Fig.S2) and triangles represent social networks above the modularity threshold. Animal social networks above the threshold experienced lower outbreak size as compared to homogeneous null networks in Fig.2*C*. In this study, we show that the disease consequences of modular subdivision in animal social networks are driven by network fragmentation and subgroup cohesion.



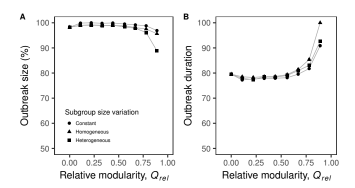
**Fig. S5.** Reduction in outbreak size experienced by empirical animal social networks as compared to their homogeneous null networks (as shown in Fig.2C) as a function of subgroup cohesion and (A) network fragmentation, and (B) subgroup size variation.



**Fig. S6.** Extended figure of Fig.3D. Disease implications of modular subdivisions in synthetic modular networks as a function of subgroup cohesion and network fragmentation. Outbreak size and duration have been normalized to the maximum observed outbreak size and duration respectively. Subgroup cohesion measures the tendency of individuals to interaction with members of their own subgroup, and is measured as the proportion of contacts within-subgroups to total contacts across the entire social network. In theory, a social network with two subgroups is considered to have  $Q_{rel}$ =0 when the proportion of contacts within subgroups is equal to the proportion of contacts between subgroups (=0.5). The minimum subgroup cohesion realized in a network is therefore a function of the number of subgroups (fragmentation) present in the network. Circles, triangles and squares summarize disease outbreaks of pathogen transmissibility of 0.1, 0.18 and 0.3, respectively.



**Fig. S8.** Disease burden in modular social networks with Poisson (purple bars) and exponential (orange bars) contact heterogeneity. The y-axis measures the change in outbreak size and duration of modular networks as compared to disease burden in homogeneous networks (with  $Q_{rel}$ =0), and is calculated as  $O_{modular} - O_{non-modular}$ , where  $O_{modular}$  and  $O_{non-modular}$  is the average outbreak size or duration of networks with  $Q_{rel} > 0$  and  $Q_{rel} = 0$ , respectively. Errors bars represent standard error of the means.



**Fig. S7.** (A) Outbreak size and (B) outbreak duration in networks with different heterogeneity in subgroup sizes. Constant = all subgroups have equal number of individuals; homogeneous = subgroup size distribution follows a Poisson distribution; heterogeneous = subgroup sizes follow an exponential distribution. For homogeneous and heterogeneous subgroup size distribution, the average subgroup size = N/network fragmentation, where N is total number of individuals in the social group. Results summarize disease simulations of a pathogen with transmissibility = 0.1 over networks with 10 subgroups. Outbreak size and duration have been normalized to the maximum observed outbreak size and duration respectively.

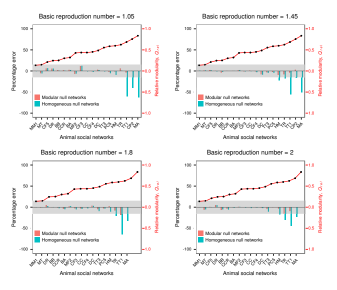
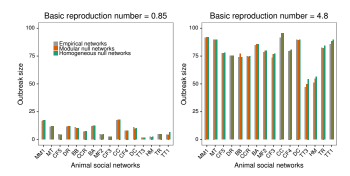


Fig. S9. Extended figure of Fig.5. Percentage error in outbreak size predictions using modular and homogeneous null networks for 19 animal social networks due to an outbreak with  $R_0$  = 1.05, 1.45, 1.8 and 2.0. Percentage error is calculated as  $(S_{emp} - S_{null})/S_{emp} \times 100$ , where S = outbreak size, emp = empirical network, null are modular or homogeneous null networks of the empirical network. The social networks are ordered according to the increasing value of relative modularity (red solid curve, secondary y-axis). The shaded region indicates the range of percentage error values below 15%. BA, Bachyteles Bachnoides; BB, Bison Bison; CC, Cercopithecus Cercopithecus



**Fig. S10.** Outbreak size predictions from modular and homogeneous null networks as compared to the "true" outbreak size (empirical networks, gray vertical bars). Results are shown for simulations of a slowly ( $R_0$ =0.85) and rapidly ( $R_0$ =4.8) spreading infectious disease. Outbreak size is averaged across all disease simulations. Both the null networks (in most cases) yield identical and accurate outbreak size predictions.

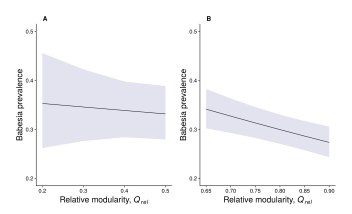


Fig. S11. Highly modular networks are able to reduce outbreak size. Empirical evidence from the spread of Babesia parasite in the field voles (Microtus agrestis): Babesia are spread by ticks and therefore networks based on common space use are appropriate to model the parasite transmission (11, 12). We therefore used the bipartite networks described in (13) to examine the relationship between network modularity and parasite transmission. Specifically, generalized linear mixed model with binomial response distribution was used, where Babesia prevalence was entered as the response and the relative modularity of networks in the previous sampling period was entered as the explanatory variable (for details see (13)). The number of nodes and the total edges were also entered as explanatory variables in the model to control for variation in the sampling effort. Additionally, the site of data collection was included as a random effect. To control for autocorrelation, we included the parasite prevalence in the previous sampling period as an explanatory variable. Two models were fit - one that explored the relationship between parasite prevalence and relative modularity in networks where  $Q_{rel} \leq 0.6$ , and other that examined association between parasite prevalence and relative modularity for networks where  $Q_{rel}>0.6.$  No significant association was found between network modularity and Babesia prevalence when  $Q_{rel} \leq 0.6$  ( $\chi^2$  = 0.21, P=0.65). However, Babasia prevalence decreased with increasing modularity when  $Q_{rel} > 0.6$  ( $\chi^2 = 12.50$ , P < 0.001).

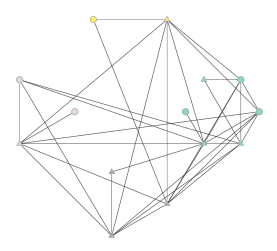


Fig. S12. Low modular networks do not limit global disease transmission. Empirical evidence from the spread of a gastrointestinal parasite in brown spider monkeys (Ateles hybridus) (14). Gastrointestinal parasites have been empirically shown to spread through the networks of physical interaction in brown spider monkeys (14). In the social networks of brown spider monkeys described in ref. (14), we identified four connected subgroups with a  $Q_{rel}$  of 0.24. Node colors indicate different subgroups. The triangle shaped nodes are the individuals that were reported infected with  $Strongyloides\,sp.$  in (14). The parasite prevalence in the connected component of the network was 57%. No structural trapping was observed in the network (i.e., infections were reported in all subgroups), which suggests that the low modularity of spider monkey social network does not inhibit the global spread of parasites.

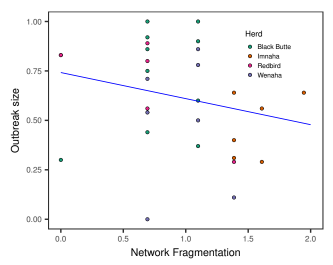


Fig. S13. Structural trapping of infectious disease increases with network fragmentation. Empirical evidence from the spread of pneumonia in the big horn lambs ( $Ovis\ canadensis$ ). Subgroups in big-horn lambs in (15) were identified based on direct or indirect association between animals to study the spread of pneumonia between individuals. This definition of subgroup implies that between-group interaction was minimal (i.e., high subgroup cohesion). Lamb mortality was considered to be a reasonable proxy of infection status (15); we therefore calculated the outbreak size as the proportion of deaths within each lamb herd (population). In the figure, each population is represented by a different point color. We found negative (albeit weak) relationship between network fragmentation and outbreak size ( $\chi^2=3.29$ , P = 0.069).

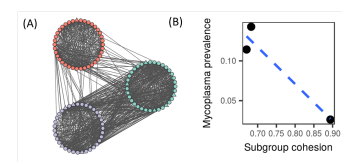


Fig. S14. High subgroup cohesion can induce structural delay of infection spread. Empirical evidence from the spread of *Mycoplasma gallisepticum* in house finches (*Haemorhous mexicanus*) (16). The social network of the songbirds based on common feeder use has been associated with spread of *Mycoplasma gallisepticum* (16). We estimated the relative modularity of the songbird social network to be 0.29 with three connected subgroups. Node colors indicate different subgroups. The triangle shaped nodes are the individuals that were reported infected with *Mycoplasma gallisepticum* in (16). The infection prevalence in the connected component of network was 9%. No structural trapping was observed (i.e., infections were reported in all the subgroups) in the network, which suggests that low modularity does not inhibit global transmission. However, the local disease prevalence (B) was inversely correlated to subgroup cohesion in the interaction networks of songbirds. This suggests that high local (subgroup) cohesion induces structural delay of infection spread within the social network.

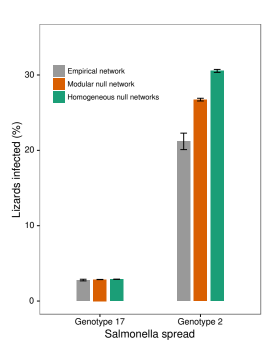


Fig. S15. Modular networks influence disease transmission of moderately, but not low, contagious pathogens/parasites. Networks based on physical proximity are associated with the spread of Salmonella in Australian sleepy lizards (17). Our investigation of modular subdivisions in the social network revealed network modularity to be moderately high ( $Q_{rel}$  = 0.624). We used the transmission data of two Salmonella strains (published in ref. (17)) to validate our theoretical prediction that network modularity influences disease transmission of moderately, but not of low contagious pathogen. To do so, we first estimated pathogen transmissibility of the two Salmonella strains (Genotype 17 and 2) to be 0.043 ( $R_0$  = 0.48) and 0.1788 ( $R_0$  = 2.02), respectively. The estimated transmissibility value was used to perform simulations of Salmonella spread in the modular and homogeneous null networks of sleepy lizards. Finally, the outbreak size of disease simulations was compared to the real transmission data of the two genotypes in the sleepy lizard social network. For the low transmissible strain (Genotype 17), we found that the homogeneous and modular null networks produced identical and accurate outbreak size predictions. This suggests that modular subdivision does not influence transmission of low contagious parasites. Conversely, for the moderately transmissible strain (Genotype 2), modular null networks performed better in estimating the true outbreak size as compared to homogeneous null networks, which suggests that modular subdivision does influence the transmission of moderately transmissible parasites.

Table S2. Empirical (or data-driven modeling) evidence that matches the predictions of our study

Key findings of this study	Evidence	Specific findings	Notes	Reference
Structural trapping by highly modular network localizes infection to a small proportion of subgroups	Plague (Yersinia pestis) infection in the great gerbil (Rhombomys opimus)	Plague spread easily within burrow systems (defined as subpopulations in the study) than between them. Probability that an animal was infectious or recovered was higher if its subpopulation was known to harbour infected animals in the recent past than if there was no evidence of recent infection	Although network modularity was not explicitly calculated, limited movement of gerbil or infected fleas between burrow systems is indicative of a high network modularity	(18)
Structural delay of infection spread	Canine distemper virus outbreak in Greater Yellowstone ecosystem's wolves ( <i>Canis lupus</i> ) and coyotes ( <i>Canis Latrans</i> )	Low connectivity (indicative of high modularity) between host population increases outbreak duration	Data driven model	(19)
	Epidemic of phocine distemper virus in the North Sea population of harbour seals ( <i>Phoca vitulina</i> )	When coupling between host patches is weak (indicative of high modularity), higher number of host patches (fragmentation) increases outbreak duration	Data-driven model	(20)
Low levels of modular organization does not structurally trap infection	Esherichia coli transmission in wild elephants (Loxodonta africana)	No evidence that animals were more likely to be infected with <i>E. coli</i> from members of their own subgroup than members of other subgroups	Aggregation of animals in single large groups suggests social networks to be homogeneous (or low modular) during rainy periods. In addition, the authors observe high overlap in home-ranges around water sources, which indicates social networks are not highly modular during non-rainy seasons	(21)
Effect of modular structure on disease trans- mission depends on pathogen contagious- ness	Cryptosporidium spread in wild lemurs	Increasing network modularity reduces outbreak size for moderately transmissible pathogen, but not for low and high pathogen transmissibility	Pathogen transmissibility can be derived from Figure 7 based $\beta$ and $\gamma$ values. Data-driven model	(22)

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