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Revealing mechanisms of infectious disease transmission through empirical contact networks

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11 Abstract

10

- 12 The spread of pathogens fundamentally depends on the underlying contacts between individuals.
- 13 Modeling infectious disease dynamics through contact networks is sometimes challenging,
- 14 however, due to a limited understanding of pathogen transmission routes and infectivity. We
- 15 developed a novel tool, INoDS (Identifying Network models of infectious Disease Spread) that
- ¹⁶ estimates the predictive power of empirical contact networks to explain observed patterns of
- ¹⁷ infectious disease spread. We show that our method is robust to partially sampled contact
- networks, incomplete disease information, and enables hypothesis testing on transmission
- ¹⁹ mechanisms. We demonstrate the applicability of our method in two host-pathogen systems:
- ²⁰ *Crithidia bombi* in bumble bee colonies and Salmonella in wild Australian sleepy lizard populations.
- ²¹ The performance of INoDS in synthetic and complex empirical systems highlights its role in
- ²² identifying transmission pathways of novel or neglected pathogens, as an alternative approach to
- laboratory transmission experiments, and overcoming common data-collection constraints.
- 24

25 Introduction

²⁶ Host contacts, whether direct or indirect, play a fundamental role in the spread of infectious ²⁷ diseases (*Newman, 2002; Rohani et al., 2010; Bansal et al., 2007; Sah et al., 2017a*). Traditional

- ²⁸ epidemiological models make assumptions of homogeneous social structure and mixing among
- hosts which can yield unreliable predictions of infectious disease spread (*Shirley and Rushton, 2005*;
 Volz and Meyers, 2007; Bansal et al., 2007; Chen et al., 2014). Network approaches to modeling the
- ³⁰ Volz and Meyers, 2007; Bansal et al., 2007; Chen et al., 2014). Network approaches to modeling the ³¹ spread of infectious diseases provide an alternative by explicitly incorporating host interactions that
- mediate pathogen transmission. Formally, in a contact network model, individuals are represented
- as nodes, and an edge between two nodes represents an interaction that has the potential to
- ³⁴ transmit infection. Constructing a complete contact network model requires (*i*) knowledge about
- the transmission routes of a pathogen, (*ii*) a sampling of all individuals in a population, and (*iii*) a
- ³⁶ sampling of all disease-causing interactions among the sampled individuals. In addition, accuracy
- of disease predictions depends on the quantification of the epidemiological characteristics of the
- ³⁸ pathogen, including the rate of pathogen transmission given a disease-causing contact between
- ³⁹ two individuals, and the rate of recovery of infected individuals.
- ⁴⁰ The use of modern technology in recent years, including RFID, GPS, radio tags, proximity loggers

and automated video tracking has enabled the collection of detailed movement and contact data,

42 making network modeling feasible. Despite the technology, logistical and financial constraints

43 still prevent data collection on all individuals and their social contacts (Welch et al., 2011; Cross

44 et al., 2012; Godfrey, 2013; Krause et al., 2013; Farine and Whitehead, 2015; Silk et al., 2015). More

45 importantly, limited knowledge about a host-pathogen system makes it challenging to identify the

- 46 mode of infection transmission, define the relevant disease-causing contacts between individuals,
- and measure per-contact rate of infection transmission (Craft and Caillaud, 2011; White et al., 2015;
- 48 Manlove et al., 2017). Laboratory techniques of unraveling transmission mechanisms usually take
- 49 years to resolve (Velthuis et al., 2007; Aiello et al., 2016; Antonovics et al., 2017). Defining accurate
- 50 contact networks underlying infection transmission in human infectious disease has been far from
- 51 trivial (Bansal et al., 2007; Pellis et al., 2014; Eames et al., 2015). For animal infectious disease,
- ⁵² limited information on host behavior and the epidemiological characteristics of the spreading
- pathogen makes it particularly difficult to define a precise contact network, which has severely
 limited the scope of network modeling in animal and wildlife epidemiology (*Craft and Caillaud*,
 2011: Craft, 2015).
- Lack of knowledge about disease transmission mechanisms has prompted the use of several 56 indirect approaches to identify the link between social structure and disease spread. A popular 57 approach has been to explore the association between social network position (usually quantified 58 as network degree) of an individual and its risk of acquiring infection (Godfrey et al., 2009, 2010; 59 Leu et al., 2010: MacIntosh et al., 2012). Another approach is to use proxy behaviors, such as 60 movement, spatial proximity or home-range overlap, to measure direct and indirect contact net-61 works occurring between individuals (Danon et al., 2011: Hamede et al., 2009: Fenner et al., 2011). 62 A recent approach, called the k-test procedure, explores a direct association between infectious 63 disease spread and a contact network by comparing the number of infectious contacts of infected 64 cases to that of uninfected cases (VanderWaal et al., 2016). However, several challenges remain in 65 identifying the underlying contact networks of infection spread that are not addressed by these 66 approaches. First, it is often unclear how contact intensity (e.g. duration, frequency, distance) relate 67 to the risk of infection transfer unless validated by transmission experiments (Aiello et al., 2016). 68 Furthermore, the role of weak ties (i.e., low intensity contacts) in pathogen transfer is ambiguous (Pellis et al., 2014: Sah et al., 2017b). The interaction network of any social group will appear as a 70 fully connected network if monitored for a long period of time. As fully-connected contact networks 71 rarely reflect the dynamics of infectious disease spread through a host population, one may ask 72 whether weak ties can be ignored, or what constitutes an appropriate intensity threshold below 73 which interactions are epidemiologically irrelevant? Second, many previous approaches ignore the 74 dynamic nature of host contacts. The formation and dissolution of contacts over time is crucial in 75 determining the order in which contacts occur, which in turn regulates the spread of infectious 76 diseases through host networks (Bansal et al., 2010: Fefferman and Ng, 2007: Farine, 2017). Finally, 77 none of the existing approaches allow direct comparison of competing hypotheses about disease 78 transmission mechanisms which may generate distinct contact patterns and consequently different 79 contact network models. 80

All of these challenges demand an approach that can allow direct comparison between com-81 peting contact network models while taking into account the dynamics of host interactions and 82 data-constraints of network sampling. In this study, we introduce a computational tool called INoDS 83 (Identifying Network models of infectious Disease Spread) that quantifies the predictive power 84 of empirical contact networks in explaining infectious disease spread, and enables comparison of 85 competing hypotheses about transmission mechanisms of infectious diseases. Our tool can also 86 infer the per-contact transmission rate of various infectious disease types (SI, SIS, and SIR), and 87 can be easily extended to incorporate other complex models of disease spread. The INoDS tool 88 provides inference on dynamic and static contact networks, and is robust to common forms of 89 missing data. Using two empirical datasets, we highlight the two-fold application of our approach – (i) to identify whether observed patterns of infectious disease spread are likely given an empirical



Figure 1. A schematic of our algorithm. **Observed data**: INoDS utilizes an observed infection time-series to estimate evidence towards a static or dynamic contact network hypothesis (or hypotheses) using a three step procedure. Shown here is an example of two competing contact network hypothesis based on different definitions of disease-causing contact (quantified by behavior A and behavior B). **Inferential steps**: In the first step, the tool estimates per-contact transmission rate parameter β , and an error parameter ϵ which captures the components of infection propagation unexplained by the edge connections of the network hypothesis. Second, the likelihood that the infectious disease spreads through the edge connections of the contact network hypothesis is compared to a distribution of likelihoods obtained from an ensemble of randomized networks. The predictive power of the empirical network hypothesis is considered to be high when its likelihood is higher than the null likelihood distribution at 5% significance level. Third, the marginal likelihood for the contact network hypothesis is calculated, which is then used to perform model comparison (using Bayes Factor, BF) between multiple contact network hypotheses, wherever available.

- ⁹² contact network, and (*ii*) to identify transmission routes, the role of the contact intensity, and the per
- ⁹³ contact transmission rate of a host-pathogen system. The epidemiological mechanisms of infection
- ⁹⁴ transmission identified by INoDS can therefore provide invaluable insights during implementation
- ⁹⁵ of immediate disease control measures in the event of an epidemic outbreak.

96 **Results**

The primary purpose of INoDS is to assess whether an empirical contact network is likely to generate 97 an observed infection time-series from a particular host population. INoDS also provides epidemio-98 logical insights into the spreading pathogen by estimating the per-contact rate of transmission. In 99 practice, the structure of a contact network model depends on the mode of infection transmission. 100 and is sensitive to the amount of missing data on nodes and edges. The tool therefore treats 101 empirically collected contact network models as network hypotheses, and facilitates hypothesis 102 testing between different contact networks. The INoDS algorithm follows a three step procedure 103 (Figure 1). First, the tool estimates a per-contact pathogen transmission rate (β) and an error 104 parameter (ϵ). The β parameter quantifies the rate of pathogen transmission through each edge of 105 the contact network, and the ϵ parameter quantifies components of infection transmission that are 106 unexplained by the edge connections of the contact network. In the second step, the likelihood of 107 the observed infection time-series under the network hypothesis and pathogen transmission rate 108 is compared to the null likelihood distribution based on an ensemble of randomized networks. The 109 randomized networks are generated by permuting the edge connections of the network hypothesis, 110

- while controlling the number of nodes and edges present. We consider the network hypothesis 111 to have high predictive power if the likelihood of thr infection time-series given the hypothesis is 112
- higher than the null likelihoods at the 5% significance level. In the final step, the marginal (Bayesian) 113
- evidence is calculated for the network hypothesis, which can be used to perform model selection 114
- between multiple network hypotheses.
- 115
- In the sections that follow, we evaluate the 116
- accuracy of the tool in recovering the transmis-117
- sion parameter, β , and its robustness to missing 118 data (missing individuals, missing contacts and 119
- missing infection cases). We further demonstrate
- 120
- the application of INoDS by using two empirical 121
- datasets: (i) spread of an intestinal pathogen in 122
- bumble bee colonies, and (ii) salmonella spread 123 in Australian sleepy lizards.
- 124

INoDS performance 125

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We evaluated the performance of INoDS on 126

- multiple infection time-series data generated by 127
- performing numerical simulations of infection 128
- spread on a synthetic dynamic network, for a 129
- wide range of pathogen transmission rates. 130

We found that INoDS accurately estimates the 131 true value of pathogen transmission rate, β , and 132 the accuracy is independent of the spreading rate 133 of the pathogen (Figure 2). The error parame-134 ter, ϵ , specified in the algorithm improves the 135 estimate of transmission rate when either the 136 network data or disease surveillance is incom-137 plete (Appendix Figure 1). The estimated rate of 138 pathogen transmission is therefore accurate even 130 when substantial network or infection time-series 140 data is missing (Appendix Figure 2). The expected 141 value of ϵ is zero when all infection transmission 142 events are explained by the edge connections in 143 the contact network hypothesis (Figure 2). Values 144 greater than zero, on the other hand, indicate 145 unexplained transmission events due to either 146 missing or inaccurate data (Appendix Figure 2). 147 Next, we tested the performance of INoDS



Figure 2. INoDS performance in recovering the per-contact pathogen transmission rate (β) for simulated infection time series under an susceptible-infected (SI) model. Each boxplot summarizes the results of 10 independent disease simulations; the black horizontal lines are the means of the estimated parameter values, the top and bottom black horizontal lines represent the standard deviation, and the tip of the black vertical line represents the maximum/minimum value. The solid red line represents one-to-one correspondence between the true value of the pathogen transmission rate (used to generate the simulated data), β^* , and the β value estimated by INoDS. Since the simulations were performed on a known synthetic network, the expected value of the error parameter, ϵ_i (represented by the green lines) is zero.

in establishing the epidemiological relevance of 149 a hypothesized contact network against three potential sources of error in data-collection: (a) 150 incomplete sampling of individuals in a population (missing nodes); (b) incomplete sampling of 151 interactions between individuals (missing edges); and (c) infrequent health diagnosis of individuals 152 (missing cases). The performance of the tool was quantified in terms of a true positive rate (i.e., the 153 proportion of times an epidemiologically relevant contact network with missing data was correctly 154 distinguished as statistically significant from an ensemble of randomized networks with the same 155 amount of missing data) and a true negative rate (i.e., the proportion of times a network with the 156 same degree distribution as the epidemiologically relevant contact network, but with randomized 157 edge connections, was correctly classified as statistically insignificant). We found INoDS to be both 158 sensitive and specific (with a high true positive and true negative rate) across a range of missing 159 data scenarios (Figure 3). The true positive rate of the tool remains close to one even when as low 160

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Figure 3. Robustness of INoDS in establishing the epidemiological significance of a hypothesized contact network under three common forms of missing data - missing nodes, missing edges and missing infection cases. True positive rate is calculated as the proportion of times (n = 20) the epidemiologically relevant (true) dynamic network was detected as statistically significant relative to a null distribution of randomized networks. True negative rate is calculated as the proportion of times (n = 20) a network with similar degree distribution as the epidemiologically relevant network, but randomized edge connections, was identified as statistically indistinguishable from the null distribution. The null distribution for a network hypothesis was generated by permuting all its edge connections, but preserving the number of nodes and edges.

Figure 3-Figure supplement 1. Robustness plot of (A) INoDS, (B) *k*-test and (C) network position test to three common forms of missing data - missing nodes, missing edges and missing infection cases. The null expectation in INoDS and the network position test was generated by permuting network edges, creating an ensemble of null networks. In the *k*-test, the location of infection cases within the observed network are permuted, creating a permuted distribution of the *k*-statistic (*VanderWaal et al., 2016*).

161 as 10% of infection cases are documented (missing cases, Figure 3). For incompletely sampled

contact networks, the true positive rate remains close to one when at least 50% nodes or 30%
 edges are documented.

The performance of INoDS in discriminating the epidemiological contact network from null 164 network hypotheses also surpasses two previous approaches – the k-test procedure and the 165 network position test (Figure 3 - figure supplement). In comparison to INoDS, the k-test and 166 network position test are sensitive to all three types of missing data. The true positive rate of the 167 k-test declines with an increasing number of missing nodes, edges, or infection cases. Of the three 168 approaches, the network position test has the lowest sensitivity (true positive rate). Since the k-test 169 procedure and network position test have been primarily used in the context of non-dynamic 170 networks, we repeated this analysis with simulated disease data from a static synthetic network. 171 Appendix Figure 4 demonstrates that even for observed networks that are not dynamic, INoDS has 172

¹⁷³ greater sensitivity and specificity than the *k*-test procedure and the network position test.

174 Applications to empirical data-sets

¹⁷⁵ We next demonstrate the application of INoDS to perform hypothesis testing on contact networks,

identify transmission mechanisms and infer transmission rate using two empirical datasets. The first
 dataset is derived from the study by *Otterstatter and Thomson (2007)* that examined the spread of

dataset is derived from the study by *Otterstatter and Thomson (2007)* that examined the spread of an intestinal pathogen (*Crithidia bombi*) within colonies of the social bumble bee, *Bombus impatiens*.

¹⁷⁹ The second dataset documents the spread of *Salmonella enterica* within two wild populations of

180 Australian sleepy lizards *Tiliqua rugosa* **Bull et al. (2012)**.

Determining transmission mechanism and the role of contact intensity: case study
 of *Crithidia bombi* in bumble bees

183 **Otterstatter and Thomson (2007)** showed that the transmission of *C. bombi* infection in bumble

¹⁸⁴ bee colonies was associated with the frequency of contacts with infected nest-mates rather than

the duration of contacts. However, the dynamic contact network models had a small number of



Figure 4. Identifying the contact network model of *Crithidia* spread in bumble bee colony (colony UN2). Edges in the contact network models represent physical interaction between the bees. Since the networks were fully connected, a series of filtered contact networks were constructed by removing weak weighted edges in the network. The x-axis represents the edge weight threshold that was used to remove weak edges in the network. Two types of edge weights were tested - duration and frequency of contacts. In addition, across all ranges of edge weight threshold, the weighted networks were converted to binary networks. The results shown are estimated values of the per contact rate transmission rate β , and estimated values of error ϵ , for the (A-B) the two types of binary network, (C) contact duration weighted network, (D) contact frequency weighted network. The faded bars correspond to networks where β parameter is statistically insignificant. Numbers above bars indicate the log Bayesian (marginal) evidence of the networks that were detected to have statistically significant higher predictive power as compared to an ensemble of null networks (P < 0.05, corrected for multiple comparisons).

nodes, and were fully connected (i.e., all individuals were connected to each other in the network) at 186 all time steps. Because predictions of infection transmission is sensitive to the size and edge density 187 of the contact network model, we extended the previous analysis by answering three specific 188 questions: (1) Do physical contact networks have higher predictive power to explain the spread of 189 C. bombi than random networks?, (2) Do the value of contact intensities (edge weights) matter in 190 transmission?, and (3) Do weak ties between individuals contribute to infection transfer? To validate 191 our tool, we performed analyses on two types of contact network models – those described by 192 frequency of contacts and those described by duration of contacts – and compared the results with 193 the findings reported in (Otterstatter and Thomson, 2007). 194 To answer the three questions, we constructed dynamic contact networks where edges represent 195 close proximity between individuals. Since fully connected networks rarely describe the dynamics of 196 infection spread, we sequentially removed edges with weights less than 10-50% of the highest edge 197 weight to generate contact network hypotheses at different edge weight thresholds. Corresponding 198 to the two types (frequency and duration) of weighted networks, unweighted contact networks 199 were also constructed by replacing weighted edges in the thresholded weighted networks with 200 binary edges (i.e., edges with an edge weight of one). 201 Figure 4 shows the estimates of pathogen transmission rate β_{ℓ} and error ϵ for the four types of 202

contact network hypotheses at different edge weight thresholds. Only a subset of contact network 203 hypotheses had statistically significant estimates of β (non faded bars). Two network hypotheses 204 summarizing frequency of contacts - binary frequency network at 15% edge weight threshold 205 and weighted frequency network at 5% edge weight threshold – demonstrated higher predictive 206 power than an ensemble of null networks. Of the two, the weighted frequency network had slightly 207 higher Bayesian evidence, although the binary frequency network was equally supported. In a 208 separate colony (UN1), only the weighted frequency network at 5% edge weight threshold had 209 higher predicted power compared to an ensemble of null networks (Appendix Figure 5). Together, 210 our results therefore show that (1) contact networks capturing frequency (but not duration) of 211 contacts have statistically high predictive power to explain the spread of C. bombi in bumble bee 212 colonies, (2) the contact networks should be weighted, and (3) weak ties (i.e., edges with weights 213 less than 5% of the highest weighted edge) are epidemiologically unimportant. 214

Identifying transmission mechanisms with imperfect disease data: case study of Salmonella enterica Australian sleepy lizards

Spatial proximity is known to be an important factor in the transmission of Salmonella enterica 217 within Australian sleepy lizard populations (**Bull et al., 2012**). However, it is not known whether 218 the transmission risk increases with frequency of proximate encounters between infectious and 219 susceptible lizards. We therefore tested two contact network hypotheses to explain the spread 220 of salmonella at two sites of wild sleepy lizards populations. The first contact network hypothesis 221 placed binary edges between lizards if they were ever within 14m distance from each other during a 222 day (24 hours). We constructed the second contact network by assigning edge weights proportional 223 to the number of times two lizards were recorded within 14m distance of each other during a day. 224 Because disease sampling was performed at regular fortnightly intervals, the true infection 225 time (day) of individuals at both study sites was unknown. We therefore used a data augmentation 226 method in INoDS (see Methods) to sample unobserved infection timings along with the per contact 227 transmission rate, θ_{i} and error, ϵ_{i} . We found that the likelihood of salmonella infection spreading 228 through the weighted contact network was significantly greater than the null expectation at both 220 sites (Figure 5). Compared to unweighted networks, networks with edges weighted by contact 230 frequency had higher marginal (Bayesian) evidence at both sites. This indicates that the occurrence 231 of repeated contacts between two spatially proximate individuals, rather than just the presence of 232 contact between individuals is important for Salmonella transmission. 233

234 **Discussion**

Network modeling of infectious disease spread is becoming increasingly popular, because modern 235 technology has dramatically improved the quality of data that can be collected from animal popula-236 tions. However, the concepts of power analysis and hypothesis testing are still underdeveloped 237 in network modeling, even though such approaches are widely recognized as key elements to 238 establish how informative and appropriate a model is (lennions and Møller, 2003: Johnson et al., 230 2015). Our ability to define a contact network relies on our understanding of host behavior, and 240 the dominant transmission mode for a given pathogen. Since such information is either derived 241 from expert knowledge (which can be subjective) or laboratory experiments (which are time- and 242 resource-intensive), it is essential to conduct an a priori analysis of contact network models to avoid 243 uninformative or misleading disease predictions. 244

In this study we therefore present INoDS as a tool that performs network model selection and 245 establishes the predictive power of a contact network model to describe the spread of infectious 246 diseases. Our method also provides epidemiological insights about the host-pathogen system 247 by enabling hypothesis testing on different transmission mechanisms, and estimating pathogen 248 transmission rates (transmission parameter, β). Unlike previous approaches, our method is robust 249 to missing network data, imperfect disease surveillance, and can provide network inference for a 250 range of disease spread models. The tool can thus be used to provide inference on contact networks 251 for a variety of pathogen types occurring both in wildlife and human populations. Inferring the role 252 of dynamic contacts on infectious disease spread requires the knowledge of either order or timing 253 of infection of individuals in the network. In practice, constraints on data collection (e.g., due to 254 infrequent health assessments), or infection diagnostics (e.g., due to sub-clinical infection, poor 255 diagnostics) precludes precise knowledge of infection timing. To overcome this challenge, our tool 256 assumes the infection times in a host population to be unobserved, and uses data on infection 257 diagnosis instead to provide inference on contact networks. 258 Our work thus addresses a growing subfield in network epidemiology that leverages statistical

Our work thus addresses a growing subfield in network epidemiology that leverages statistical
 tools to infer contact networks using all available host and disease data (*Welch et al., 2011; Stack et al., 2013; VanderWaal et al., 2016; Groendyke et al., 2011*). Our approach can be used to tackle
 several fundamental challenges in the field of infectious disease modeling (*Eames et al., 2015; Pellis et al., 2014*). First, INoDS can be used to perform model selection on contact network models



Figure 5. Identifying transmission mechanisms of Salmonella spread in Australian sleepy lizards. Dynamic network of proximity interactions for a total duration of 70 days between (A) 43 lizards at site 1, and (B) 44 lizards at site 2. Each temporal slice summarizes interactions within a day (24 hours). Edges indicate that the pair of individuals were within 14m distance of each other, and the edge weights are proportional to the frequency of physical interactions between the node pair. Green nodes are the animals that were diagnosed to be uninfected at that time-point, red are the animals that were diagnosis to be infected and grey nodes are the individuals with unknown infection status at the time-point. We hypothesized that the spatial proximity networks could explain the observed spread of *Salmonella* in the population. The results are summarized as a table. Bold numbers indicate that the network hypothesis was found to have high predictive power compared to an ensemble of randomized networks. The network hypothesis with the highest log Bayesian (marginal) evidence at each site is marked with an asterisk (*)

that guantify different transmission modes; this approach therefore facilitates the identification 264 of infection-transmitting contacts and does not rely on laboratory experimentation (or subjective 265 expert knowledge). Second, INoDS can be used to establish the predictive power of proxy measures 266 of contact in cases where limited interaction data is available. For example, spatial proximity, 267 home-range overlap or asynchronous refuge use are commonly used as a proxy for disease-causing 268 contact in wild animal populations (Godfrev et al., 2010: Leu et al., 2010: Sah et al., 2016). INoDS 269 establishes the epidemiological significance of such assumptions by comparing the likelihood 270 of infection spread occurring along the edges of the proxy contact network to the likelihoods 271 generated from an ensemble of random networks. Third, it is well known that not all contacts 272 between hosts have the same potential for infection transmission. The heterogeneity of host 273 contacts in a network model is typically captured through edge weights, but it is often not clear 274 which type of edge weights (frequency, duration or intensity) is relevant in the context of a specific 275 host pathogen system (*Pellis et al., 2014*). Through model selection of contact networks with similar 276 edge connection but different edge weighting criterion, INoDS can help establish a link between 277 edge weight and the risk of transmission across an edge in a contact network. 278

We demonstrate the application of INoDS using two real-world datasets. In the first dataset, we 279 used INoDS to determine the role of edge weight type and edge weight value on the predictive 280 power of the contact network. To accurately model the spread of the *Crithidia* gut protozoan in 281 bumble bee colonies, we show that the contact networks weighted with respect to frequency, 282 rather than duration, have higher predictive power given observed patterns of transmission. Our 283 results therefore support the original finding of the study (Otterstatter and Thomson, 2007), where 284 individual risk of infection was found to be correlated with contact rate with infected nest-mates. 285 Our analysis further extends previous findings in this system by comparing the observed patterns 286 of transmission against null expectations from random networks, and assessing the likelihood 287 of competing network hypotheses. We find weak ties below a certain threshold do not play an 288 important role in infection transfer. Contact networks where such weak weighted edges have 289 been removed, therefore, demonstrate higher predictive power than fully connected networks. In 290 the next empirical example, we explore the transmission mechanisms of a commensal bacterium 291 in wild populations of Australian sleepy lizards. We find that taking repeated contacts between 292 closely located lizards into account allows better, i.e. more consistent, predictions on Salmonella 293 transmission. 294

The current version of INoDS assumes the infection process has no latent period, and that the infectiousness of infected hosts and susceptibility of naive hosts is equal for all individuals in the population. These assumptions can be relaxed to incorporate more complex disease progression. For instance, heterogeneity in infectiousness of infected hosts and the susceptibility of naive hosts can be incorporated as random effects in the model by assuming the two follow a Gaussian distribution. Disease latency can also be incorporated using a data-augmentation technique, similar to what we use for inferring infection times.

Our results show that the data-collection efforts should aim to sample as many individuals in 302 the population as possible, since missing nodes have the greatest impact (rather than missing 303 edges) on the predictive power of network models. Since data-collection for network analysis can be 30/ labor-intensive and time-consuming, our approach can be used to make essential decisions on how 305 limited data collection resources should be deployed. Our approach can also be used to improve 306 targeted disease management and control by identifying high-risk behaviors and super-spreaders 307 of a novel pathogen without relying on expensive transmission experiments that take years to 308 resolve. 300

310 Methods

Here we describe INoDS, a computational tool that (*i*) estimates per contact transmission rate (β) of infectious disease for empirical contact networks, (*ii*) establishes predictive power of a contact network by comparisons with an ensemble of randomized networks, and (*iii*) enables discrimination of competing contact network hypotheses, including those based on pathogen transmission mode,

edge weight criteria and data collection techniques. Two types of data are required as input for

³¹⁶ INoDS – infection time-series data, which include infection diagnoses (coded as 0 = not infected and

1 = infected), and time-step of diagnosis for all available individuals in the population; and an edge-

list of a dynamic (or static) contact network. An edge-list format is simply the list of node pairs (each

node pair represents an edge of the network), along with the weight assigned to the interaction, and

time-step of interaction, with one node pair per line. The tool can be used for unweighted contact networks - an edge weight of one is assigned to all edges in this case. Time-steps of interactions are

networks - an edge weight of one is assigned to all edges in this case. Time-steps of interactions are not required when analysis is performed on static contact networks. The software is implemented in

not required when analysis is performed on static contact networks. The software is implemented in
 Python, is platform independent, and is freely available at https://bansallab.github.io/INoDS-model/.

324 INoDS formulation

³²⁵ We assume that at each instance the potential of acquiring infection for a susceptible individual *i* ³²⁶ depends on the per contact transmission rate β , the total strength of interactions with its infected ³²⁷ neighbors at the previous time-step, and an error parameter ϵ that captures the force of infection ³²⁸ that is not explained by the individual's social connections. The infection receiving potential, $\lambda_i(t_i)$,

of individual i at time t is thus calculated as:

$$\lambda_i(t) = 1 - \exp\{-\beta w_i(t-1) - \epsilon\},\tag{1}$$

where both transmission rate β and error parameter ϵ are > 0; $w_i(t-1)$ denotes the total strength of association between the focal individual *i* and its infected associates at the previous time-step (t-1). For binary (unweighted) contact network models $w_i = k_i$, where k_i is the total infected connections of the focal individual.

The log-likelihood for all observed timings of infection in a population given the contact network hypothesis (H_A) can therefore be estimated as:

$$\log(D|H_A, \beta, \epsilon) = \sum^n \log[\lambda_n(t_n)] + \sum^t \left(\sum^m \log[1 - \lambda_m(t)]\right),$$
(2)

where t_n is the time of infection of individual *n*. The first part of equation 2 estimates the log likelihood of all observed infection acquisition events. The second part of the equation represents the log-likelihood of susceptible individuals *m* remaining uninfected at time *t*.

³³⁹ Parameter estimation and data augmentation of infection timings

We adopted a Bayesian MCMC framework to estimate the unknown model parameters. Calculation 340 of the likelihood in equation 2 requires knowledge of exact timing of infection, t_1, \dots, t_n , for *n* infected 3/11 individuals in the population. However in many cases, the only data that is available are the timings 342 of when individuals in a populations were *diagnosed* to be infected, $d_1, ..., d_n$. We therefore employ 3/13 a Bayesian data augmentation approach to estimate the actual infection timings in the disease 344 dataset (Tanner and Wong, 1987). Since in this case the actual infection time t, for an individual i is 345 unobserved, we only know that the timing of infection for the individual lies between the interval 346 (L_i, d_i) , where L_i is the last negative diagnosis of individual i before infection acquisition. Within this 347 interval, the individual could have potentially acquired infection at any time-step where it was in 348 contact with other individuals in the network. Assuming incubation period to be one time-step, we 340 can therefore represent the potential set of infection timings as $t_i \in \{g_i(t_i-1) > 0, L_i < t_i \le d_i\}$, where 350 $g_i(t_i-1)$ is the degree (number of contacts) of individual i at time $t_i - 1$. For infections that follow a 351 SIS or SIR disease model, it is also essential to impute the recovery time of infected individuals for 352 accurate estimation of infected degree. To do so, we adopt a similar data augmentation approach 353 as described to sample from the set of possible recovery time-points. 354

The joint posterior distribution of augmented data and the set of parameters is proportional to:

$$P(\Theta|D,H) = \frac{\mathcal{L}(D|H,\Theta)\mathcal{P}(\Theta|H)}{\mathcal{E}(D|H)} \propto \mathcal{L}(D|H,\Theta)\mathcal{P}(\Theta|H)$$
(3)

where D is the infection time-series data. H is the contact network hypothesis, and P. L. P. \mathcal{E} 356 are the shorthands for the posterior, the likelihood, the prior and the evidence, respectively. The 357 data augmentation proceeds in two steps. In the first step, the missing infection times are imputed 358 conditional on the possible set of infection times. In the next step the posterior distributions of 359 the unknown parameters are sampled based on the imputed data. We performed data imputation 360 using inverse transform sampling method, which is a technique of drawing random samples from 361 any probability distribution given its cumulative distribution function (Robert and Casella, 2004). 362 We used a used a uniform prior on [0, 1000] for the per contact transmission rate and the error 363 parameter. 364

MCMC sampling of the unknown parameters is performed using the *PTsampler* function of *emcee* 365 package implemented in Python (Foreman-Mackey et al., 2013). PTsampler is an implementation 366 of the affine-invariant ensemble MCMC algorithm which provides efficient sampling of highly cor-367 related parameters - a common problem when using simple Metropolis-Hastings type samplers 368 (Foreman-Mackey et al., 2013). INoDS uses twice the number of walkers as the total model pa-369 rameters, and the temperature is set to T = 15 to maximize the sampling of the parameter space. 370 The values of The number of sampling steps and burn-in is specified by the user. Convergence is 371 assessed using an autocorrelation plot of few randomly selected walkers. From the joint posterior 372 estimates of β and ϵ , we report the parameter combination with the highest maximum likelihood 373 value. 374

375 Statistical significance of infection transmission parameter

The statistical significance of parameter β is determined by comparing the force of infection explained by edge connections (= $\beta w_i(t-1)$) at each infection event to the error parameter ϵ . The *p*-value is calculated as the proportion of transmission events where the force of infection is greater than the error estimate. The per contact transmission rate β is considered to be statistically significant when its calculated *p*-value is less than 0.05.

381 Interpretation of the error parameter

In principle, inclusion of the error parameter in eq. 1 is similar to the asocial learning rate used in 382 the network based diffusion analysis approach in the behavior learning literature (Franz and Nunn, 383 2009: Aplin et al., 2013). However, in contrast to the asocial learning rate which quantifies the 384 rate of spontaneous learning, ϵ in INoDS formulation serves to improve the estimation of the per 385 contact transmission rate, β , when either the contact network or infection spread is not completely 386 sampled (Appendix Figure 1 and 2). The magnitude of ϵ can also be used to (approximately) 387 assess the magnitude of missing data (Appendix Figure 3). The percentage transmission events 388 where ϵ is greater than the force of infection explained by edge connections (= $\beta w_{i}(t-1)$) increases 389 proportionately with increasing amount of missing network data. The relative differences between 390 social force of infection and unexplained transmission events, however is less sensitive to missing 391 data on infection cases. 392

³⁹³ Predictive power of a contact network hypothesis

We assess the predictive power of a contact network hypothesis by performing comparisons 394 with an ensemble of randomized networks with same number of nodes and edge connectivity. 395 Specifically, the likelihood of the infection data given the network and estimated model parameters 396 (i.e., $\mathcal{L}(D|H, \Theta)$) is compared to a distribution of likelihoods of infection data (given the estimated 307 model parameters) obtained from the null networks (i.e., $\mathcal{L}(D|H_{\Omega_1},\Theta)$, $\mathcal{L}(D|H_{\Omega_2},\Theta)$,..., $\mathcal{L}(D|H_{\Omega_2},\Theta)$; 398 n = 500). Null networks are generated by randomizing edge connections of the contact network 300 hypothesis, which preserves the edge density in the permuted networks. Next, a p-value is calculated 400 as the proportion of randomizations which generate a likelihood greater than the likelihood of 401 the empirical network hypothesis. The empirical contact network is considered to have a higher 407 predictive power than the null expectation when its calculated *p*-value is less than 0.05. 403

404 Model selection of competing network hypotheses

To facilitate model selection in cases where there are more than one network hypothesis, we compute the marginal likelihood of the infection data given each contact network model. The marginal likelihood, also called the Bayesian evidence, measures the overall model fit, i.e., to what extent the infection time-series data can be simulated by a network hypothesis (H_A). Bayesian evidence is based on the average model fit, and calculated by integrating the model fit over the entire parameter space:

$$P(D|H) = \int \mathcal{P}(\Theta|H)\mathcal{L}(D|H,\Theta)d\Theta$$
(4)

Since it is difficult to integrate Eq.4 numerically, we estimate the marginal likelihood of network models using thermodynamic integration, or path sampling (*Lartillot and Philippe, 2006*) method implemented in *emcee* package in Python. Model selection can be then performed by computing pair-wise Bayes factor, i.e. the ratio of the marginal likelihoods of two network hypotheses. The log Bayes factor to assess the performance of network hypothesis H_A over network hypothesis H_B , is expressed as:

$$log(B_{BA}) = log(P(D|H_B)) - log(P(D|H_A))$$
(5)

The contact network with a higher marginal likelihood is considered to be more plausible, and a log Bayes' factor of more than 3 is considered to be a strong support in favor of the alternative network model (H_B) (*Kass and Raftery*, **1995**).

420 Evaluating INoDS performance

We evaluated the accuracy of the in estimating the unknown transmission parameter β , and its 421 robustness to missing data was evaluated. To do so we first constructed a dynamic synthetic 422 network using the following procedure. At time-step t = 0, a static network of 100 nodes, mean 423 degree 4, and Poisson degree distribution was generated using the configuration model (*Mollov* 424 and Reed, 1995). At each subsequent time-step, 10% of edge-connections present in the previous 425 time-step were permuted, for a total of 100 time-steps. Next, through the synthetic dynamic 426 network, we performed 10 independent SI disease simulations with per contact rate of infection 427 transmission 0.01 to 0.1. Model accuracy was determined by comparing the estimated transmission 428 parameter, β , with the true transmission rate β^* that was used to perform disease simulations. 429 Since the synthetic network dataset did not contain any missing data, model accuracy was also 430 tested by evaluating the deviation of the estimated error parameter ϵ , from the expected value of 431 432 zero.

We also tested robustness of the tool in establishing the epidemiological relevance of a hypoth-433 esized contact network against three potential sources of error in data-collection: missing nodes. 434 missing edges, and missing cases. The three scenarios of missing data were created by randomly 435 removing 10-95% of nodes, edges or infection cases from the simulated dataset described above 436 True positive rate was calculated as the proportion of times the hypothesized contact network 437 model with missing data was correctly distinguished as statistically significant from an ensemble 438 of null networks generated by randomizing its edge connections. We calculated the true negative 439 rate as the proportion of times a network with the same degree distribution as the contact network 440 hypothesis, but randomized edge connections, was correctly classified as statistically insignificant. 441 Next, we compared INoDS with two previous approaches (k-test and network position test) that 442 have been used to establish an association between infection spread and contact network in a 443 host population. The k-test procedure involves estimating the mean infected degree (i.e., number 444 of direct infected contacts) of each infected individual in the network, called the k-statistic. The 445 *p*-value in the *k*-test is calculated by comparing the observed *k*-statistic to a distribution of null 446 k-statistics which is generated by randomizing the node-labels of infection cases in the network 447 (VanderWaal et al., 2016). Network position test compares the degree of infected individuals to that 118

of uninfected individuals (Godfrey et al., 2009, 2010; MacIntosh et al., 2012). The observed network

450 is considered to be epidemiologically relevant when the difference in average degree between

⁴⁵¹ infected and uninfected individuals exceeds (at 5% significance level) the degree difference in an

ensemble of randomized networks.

453 Applications to empirical data-sets

We demonstrate the applications of our approach using two datasets from the empirical literature. 454 The first dataset comprises of dynamic networks of bee colonies (N = 5-7 individuals), where edges 455 represent direct physical contacts that were recorded using a color-based video tracking software. 456 A bumble bee colony consists of a single queen bee and infertile workers. Here, we focus on the 457 infection experiments in two colonies where infection was artificially introduced through a randomly 458 selected forager (colony UN1 and UN2). Infection progression through the colonies was tracked by 459 daily screening of individual feces, and the infection timing was determined using the knowledge of 460 the rate of replication of *C. bombi* within its host intestine. 461

The second dataset monitors the spread of the commensal bacterium Salmonella enterica in two 462 separate wild populations of the Australian sleepy lizard *Tiligua rugosa*. The two sites consisted of 43 463 and 44 individuals respectively, and these represented the vast majority of all resident individuals at 16/ the two sites (i.e., no other individuals were encountered during the study period). Individuals were 465 fitted with GPS loggers and their locations were recorded every 10 minutes for 70 days. Salmonella 466 infections were monitored using cloacal swabs on each animal once every 14 days. Consequently, 467 the disease data in this system do not identify the onset of each individual's infection. We used 468 a SIS (susceptible-infected-susceptible) disease model to reflect the fact that sleepy lizards can 469 be reinfected with salmonella infections. Proximity networks were constructed by assuming a 470 contact between individuals whenever the location of two lizards was recorded to be within 14m 471 distance of each other (Leu et al., 2010). The dynamic networks at both sites consisted of 70 472 static snapshots, with each snapshot summarizing a day of interactions between the lizards. We 473 constructed two contact network hypotheses to explain the spread of salmonella. The first contact 474 network hypothesis placed binary edges between lizards if they were ever within 14m distance 475 from each other during a day. The second contact network assigned edge weights proportional to 476 the number of times two lizards were recorded within 14m distance of each other during a day. 477 Specifically, edge weights between two lizards were equal to their frequency of contacts during a 478

day normalized by the maximum edge weight observed in the dynamic network.

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- R86 Competing interests
- ⁴⁸⁷ The authors declare that no competing interests exist.

488 **References**

Aiello CM, Nussear KE, Esque TC, Emblidge PG, Sah P, Bansal S, et al. Host contact and shedding patterns
 clarify variation in pathogen exposure and transmission in threatened tortoise <i>Gopherus agassizii</i>
 implications for disease modeling and management. Journal of Animal Ecology. 2016; p. n/a–n/a. http:

492 //doi.wiley.com/10.1111/1365-2656.12511, doi: 10.1111/1365-2656.12511.

Antonovics J, Wilson AJ, Forbes MR, Hauffe HC, Kallio ER, Leggett HC, et al. The evolution of transmission
 mode. Philosophical Transactions of the Royal Society B: Biological Sciences. 2017; 372:20160083. doi:
 10.1098/rstb.2016.0083.

- Aplin LM, Farine DR, Morand-Ferron J, Cole EF, Cockburn A, Sheldon BC. Individual personalities predict
 social behaviour in wild networks of great tits (Parus major). Ecology Letters. 2013; 16(11):1365–1372. doi:
 10.1111/ele.12181.
- Bansal S, Grenfell BT, Meyers LA. When individual behaviour matters: homogeneous and network models
 in epidemiology. Journal of the Royal Society, Interface / the Royal Society. 2007 oct; 4(16):879–91. doi:
 10.1098/rsif.2007.1100.
- Bansal S, Read J, Pourbohloul B, Meyers LA. The dynamic nature of contact networks in infectious disease
 epidemiology. Journal of Biological Dynamics. 2010; 4(5):478–489. doi: 10.1080/17513758.2010.503376.
- Bull CM, Godfrey SS, Gordon DM. Social networks and the spread of Salmonella in a sleepy lizard population.
 Molecular Ecology. 2012; 21(17):4386–4392. doi: 10.1111/j.1365-294X.2012.05653.x.
- Chen S, White BJ, Sanderson MW, Amrine DE, Ilany A, Lanzas C. Highly dynamic animal contact network and
 implications on disease transmission. Scientific reports. 2014; 4:4472. doi: 10.1038/srep04472.
- Craft ME. Infectious disease transmission and contact networks in wildlife and livestock. Philosophical
 Transactions of the Royal Society of London Series B, Biological Xciences. 2015; 370(1669):1–12. doi:
 10.1098/rstb.2014.0107.
- Craft ME, Caillaud D. Network models: An underutilized tool in wildlife epidemiology? Interdisciplinary
 Perspectives on Infectious Diseases. 2011 jan; 2011:676949. doi: 10.1155/2011/676949.
- Cross PC, Creech TG, Ebinger MR, Heisey DM, Irvine KM, Creel S. Wildlife contact analysis: emerging methods,
 questions, and challenges. Behavioral Ecology and Sociobiology. 2012 jul; 66(10):1437–1447. http://link.
 springer.com/10.1007/s00265-012-1376-6, doi: 10.1007/s00265-012-1376-6.
- Danon L, Ford AP, House T, Jewell CP, Keeling MJ, Roberts GO, et al. Networks and the epidemiology of
 infectious disease. Interdisciplinary perspectives on infectious diseases. 2011 jan; 2011:284909. doi:
 10.1155/2011/284909.
- Eames K, Bansal S, Frost S, Riley S. Six challenges in measuring contact networks for use in modelling. Epidemics.
 2015; 10:72–77. doi: 10.1016/j.epidem.2014.08.006.
- Farine D. The dynamics of transmission and the dynamics of networks. Journal of Animal Ecology. 2017; 86(3):415–418. doi: 10.1111/1365-2656.12659.
- Farine DR, Whitehead H. Constructing, conducting, and interpreting animal social network analysis. The Journal
 of animal ecology. 2015; (July):1144–1163. doi: 10.1111/1365-2656.12418.
- Fefferman NH, Ng KL. How disease models in static networks can fail to approximate disease in dynamic net works. Physical Review E Statistical, Nonlinear, and Soft Matter Physics. 2007; 76(3):1–11. doi: 10.1103/Phys RevE.76.031919.
- Fenner AL, Godfrey SS, Michael Bull C. Using social networks to deduce whether residents or dispersers
 spread parasites in a lizard population. Journal of Animal Ecology. 2011; 80(4):835–843. doi: 10.1111/j.1365-2656.2011.01825.x.
- Foreman-Mackey D, Hogg DW, Lang D, Goodman J. emcee: The MCMC Hammer. Publications of the Astronom ical Society of the Pacific. 2013; 125(925):306–312. doi: 10.1086/670067.
- Franz M, Nunn CL. Network-based diffusion analysis: a new method for detecting social learning. Proc Biol Sci.
 2009; 276(1663):1829–1836. doi: 10.1098/rspb.2008.1824.
- **Godfrey SS.** Networks and the ecology of parasite transmission: A framework for wildlife parasitology. International journal for parasitology Parasites and wildlife. 2013 dec; 2(1):235–245. doi: 10.1016/j.ijppaw.2013.09.001.
- Godfrey SS, Bull CM, James R, Murray K. Network structure and parasite transmission in a group living
 lizard, the gidgee skink, Egernia stokesii. Behavioral Ecology and Sociobiology. 2009; 63(7):1045–1056. doi:
 10.1007/s00265-009-0730-9.
- Godfrey SS, Moore JA, Nelson NJ, Bull CM. Social network structure and parasite infection patterns in a territorial
 reptile, the tuatara (Sphenodon punctatus). International Journal for Parasitology. 2010; 40(13):1575–1585.
 doi: 10.1016/j.ijpara.2010.06.002.

- 544 Groendyke C, Welch D, Hunter DR. Bayesian inference for contact networks given epidemic data. Scandinavian
 545 Journal of Statistics. 2011; 38(3):600–616.
- Hamede RK, Bashford J, McCallum H, Jones M. Contact networks in a wild Tasmanian devil (Sarcophilus
- harrisii) population: using social network analysis to reveal seasonal variability in social behaviour and its
 implications for transmission of devil facial tumour disease. Ecology letters. 2009 nov; 12(11):1147-57. doi:
 10.1111/i.1461-0248.2009.01370.x.
- Jennions M, Møller A. A survey of the statistical power of research in behavioral ecology and animal behavior.
 Behavioral Ecology. 2003; 14(3):438–445. doi: 10.1093/beheco/14.3.438.
- Johnson PCD, Barry SJE, Ferguson HM, Müller P. Power analysis for generalized linear mixed models in ecology and evolution. Methods in Ecology and Evolution. 2015; 6(2):133–142. doi: 10.1111/2041-210X.12306.
- Kass R, Raftery A. Bayes Factors. Journal of the American Statistical Association. 1995; 90(430):773–795. doi:
 10.1080/01621459.1995.10476572.
- Krause J, Krause S, Arlinghaus R, Psorakis I, Roberts S, Rutz C. Reality mining of animal social systems. Trends in
 ecology & evolution. 2013 sep; 28(9):541–51. doi: 10.1016/j.tree.2013.06.002.
- Lartillot N, Philippe H. Computing Bayes Factors Using Thermodynamic Integration. Systematic Biology. 2006;
 55(2):195–207. doi: 10.1080/10635150500433722.
- Leu ST, Kappeler PM, Bull CM. Refuge sharing network predicts ectoparasite load in a lizard. Behavioral ecology
 and sociobiology. 2010 sep; 64(9):1495–1503. doi: 10.1007/s00265-010-0964-6.
- MacIntosh AJJ, Jacobs A, Garcia C, Shimizu K, Mouri K, Huffman Ma, et al. Monkeys in the middle: para site transmission through the social network of a wild primate. PloS one. 2012 jan; 7(12):e51144. doi:
 10.1371/journal.pone.0051144.
- 565 Manlove KR. Cassirer EF. Plowright RK. Cross PC. Hudson Pl. Contact and contagion: Probability of transmission
- given contact varies with demographic state in bighorn sheep. Journal of Animal Ecology. 2017; 86(4):908–920.
 doi: 10.1111/1365-2656.12664.
- Molloy M, Reed B. A Critical Point for Random Graphs With a Given Degree Sequence. Random Structures and
 Algorithms. 1995; 6(2-3):161–180.
- Newman MEJ. Spread of epidemic disease on networks. Physical Review E Statistical, Nonlinear, and Soft
 Matter Physics. 2002; 66(1):1–11. doi: 10.1103/PhysRevE.66.016128.
- Otterstatter MC, Thomson JD. Contact networks and transmission of an intestinal pathogen in bumble bee
 (Bombus impatiens) colonies. Oecologia. 2007 nov; 154(2):411–21. doi: 10.1007/s00442-007-0834-8.
- Pellis L, Ball F, Bansal S, Eames K, House T, Isham V, et al. Eight challenges for network epidemic models.
 Epidemics. 2014; 10:58–62. doi: 10.1016/j.epidem.2014.07.003.
- **Robert C**, Casella G. Monte Carlo Statistical Methods. Springer Texts in Statistics; 2004.
- Rohani P, Zhong X, King AA. Contact Network Structure Explains the Changing Epidemiology of Pertussis.
 Science. 2010; 330(6006):982–985. doi: 10.1126/science.1194134.
- Sah P, Leu ST, Cross PC, Hudson PJ, Bansal S. Unraveling the disease consequences and mechanisms of modular
 structure in animal social networks. Proceedings of the National Academy of Sciences of the United States of
 America. 2017 apr; 114(16):4165–4170. doi: 10.1073/pnas.1613616114.
- Sah P, Mann J, Bansal S. Disease implications of animal social network structure: a synthesis across social
 systems. Journal of Animal Ecology. 2017; .
- Sah P, Nussear KE, Esque TC, Aiello CM, Hudson PJ, Bansal S. Inferring social structure and its drivers from
 refuge use in the desert tortoise, a relatively solitary species. Behavioral Ecology and Sociobiology. 2016; p.
 1–13. doi: 10.1007/s00265-016-2136-9.
- Shirley MDF, Rushton SP. The impacts of network topology on disease spread. Ecological Complexity. 2005;
 2(3):287–299. doi: 10.1016/j.ecocom.2005.04.005.
- Silk MJ, Jackson AL, Croft DP, Colhoun K, Bearhop S. The consequences of unidentifiable individuals for the
 analysis of an animal social network. Animal Behaviour. 2015; 104:1–11. doi: 10.1016/j.anbehav.2015.03.005.

- Stack JC, Bansal S, Anil Kumar VS, Grenfell B. Inferring population-level contact heterogeneity from common
 epidemic data. Journal of the Royal Society, Interface / the Royal Society. 2013; 10(78):20120578. doi:
- ⁵⁹³ 10.1098/rsif.2012.0578.
- Tanner MA, Wong WH. The Calculation of Posterior Distributions by Data Augmentation: Rejoinder. Journal of
 the American Statistical Association. 1987; 82(398):548–550. doi: 10.2307/2289463.
- VanderWaal K, Enns EA, Picasso C, Packer C, Craft ME. Evaluating empirical contact networks as potential
 transmission pathways for infectious diseases. Journal of The Royal Society Interface. 2016; 13(121):20160166.
- ⁵⁹⁸ doi: 10.1098/rsif.2016.0166.
- Velthuis aGJ, Bouma A, Katsma WEa, Nodelijk G, De Jong MCM. Design and analysis of small-scale
 transmission experiments with animals. Epidemiology and infection. 2007; 135(2):202–217. doi:
 10.1017/S095026880600673X.
- Volz E, Meyers LA. Susceptible-infected-recovered epidemics in dynamic contact networks. Proceedings of the
 Royal Society B: Biological Sciences. 2007; 274(1628):2925–2934. doi: 10.1098/rspb.2007.1159.
- Welch D, Bansal S, Hunter DR. Statistical inference to advance network models in epidemiology. Epidemics.
 2011; 3(1):38–45. doi: 10.1016/j.epidem.2011.01.002.
- White LA, Forester JD, Craft ME. Using contact networks to explore mechanisms of parasite transmission in
 wildlife. Biological Reviews, 2015: http://doi.wilev.com/10.1111/brv.12236, doi: 10.1111/brv.12236.

608 Appendix 1



Appendix 1 Figure 1. Relative error in the estimations of parameter β under missing data conditions with and without the inclusion of the error parameter (ϵ) in the INoDS formulation. The simulated infectious disease spread (SI model, per contact transmission rate pathogen = 0.03) was performed on static network with 100 nodes, Poisson degree distribution, and a average degree of 3. Relative error was calculated as $\frac{\beta - \beta'}{\beta}$, where β is the per contact transmission rate of the simulated pathogen (=0.03) and β' is the value of social transmission parameter estimated



Appendix 1 Figure 2. Estimation of per contact transmission rate (β) and the error parameter (ϵ) by INoDS under three forms of missing data conditions - (A) missing nodes, (B) missing edges and (C) missing infection cases. Simulations of susceptible-infected (SI) model of infectious disease spread were performed on static network with 100 nodes, Poisson degree distribution, and an average degree of 3. Each boxplot summarizes the results of 10 independent disease simulations; the horizontal line in the middle is the mean of estimated parameter values, the top and the bottom horizontal line is the standard deviation, and the tip of the vertical line represents the maximum/minimum value. The solid red line represents the true value of β used in the disease simulations. Since the simulations were performed on a known synthetic network, the expected value of error parameter is zero.



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Appendix 1 Figure 3. Relationship between error ϵ and force of infection (= $\beta w_i(t - 1)$) with increasing percentage of missing data. Each boxplot summarizes the results of 10 independent disease simulations (indicated by points); the horizontal line in the middle is the mean percent transmission events where the asocial force is greater than the infection force contributed by the social connections. The top and the bottom horizontal line is the standard deviation, and the tip of the vertical line represents the maximum/minimum value.



Appendix 1 Figure 4. Plot of sensitivity and specificity of (A) INoDS, (B) *k*-test and (C) network position test to three common forms of missing data - missing nodes, missing edges and missing infection cases. The observed network in this case is a static network with 100 nodes, Poisson degree distribution and a mean network degree of 3. Simulations of pathogen spread with per contact transmission rate of 0.03 were performed through the observed static network. Null expectation in INoDS and network position test was generated by permuting the edge connections of the observed networks, creating an ensemble of null networks. In *k*-test, the location of infection cases within the observed network are permuted, creating a permuted distribution of *k*-statistic (*VanderWaal et al., 2016*)



Appendix 1 Figure 5. Identifying the contact network model of *Crithidia* spread in bumble bee colony (colony UN2). Edges in the contact network models represent physical interaction between the bees. Since the networks were fully connected, a series of filtered contact networks were constructed by removing weak weighted edges in the network. The x-axis represents the edge weight threshold that was used to remove weak edges in the network. Two types of edge weights were tested - duration and frequency of contacts. In addition, across all ranges of edge weight threshold, the weighted networks were converted to binary networks. The results shown are estimated values of the per contact rate of infection transmission β , and estimated values of error ϵ , for the (A-B) two types of binary network, (C) contact duration weighted network, (D) contact frequency weighted network. The faded bars correspond to networks where β parameter is statistically insignificant. Numbers above bars indicate the log Bayesian (marginal) evidence of the networks that were detected to have statistically significant higher predictive power as compared to an ensemble of null networks (P < 0.05, corrected for multiple comparisons). We note that frequency networks with more than 25% weak edge removed failed to converge in (C) and (D), and therefore the transmission parameter associated with these contact networks were not estimated.



Figure 3-Figure supplement 1. Robustness plot of (A) INoDS, (B) *k*-test and (C) network position test to three common forms of missing data - missing nodes, missing edges and missing infection cases. The null expectation in INoDS and the network position test was generated by permuting network edges, creating an ensemble of null networks. In the *k*-test, the location of infection cases within the observed network are permuted, creating a permuted distribution of the *k*-statistic (*VanderWaal et al., 2016*).