The Basic Reproductive Number for Disease Systems with Multiple Coupled Heterogeneities

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Abstract

In mathematical epidemiology, a well-known formula describes the impact of heterogeneity on the basic reproductive number for situations in which transmission is separable and for which there is one source of variation in susceptibility and one source of variation in infectiousness. This formula is written in terms of the magnitudes of the heterogeneities, as quantified by their coefficients of variation, and the correlation between them. A natural question to ask is whether analogous results apply when there are multiple sources of variation in susceptibility and/or infectiousness. In this paper we demonstrate that under three or more coupled heterogeneities, the basic reproductive number depends on details of the distribution of the heterogeneities in a way that is not seen in the well-known simpler situation. We provide explicit results for the cases of multivariate normal and multivariate log-normal distributions, showing that the basic reproductive number can again be expressed in terms of the magnitudes of the heterogeneities and the pairwise correlations between them. The results, however, differ between the two multivariate distributions, demonstrating that no formula of this type applies generally when there are three or more

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coupled heterogeneities. We see that the results are approximately equal when heterogeneities are relatively small and show that an earlier result in the literature (Koella, 1991) should be viewed in this light. We provide numerical illustrations of our results.

1 1. Introduction

The basic reproductive number, R_0 , plays a crucial role in determining both whether a pathogen is able to spread and the strength of control measures needed to halt its spread. The simplest descriptions of R_0 assume simple transmission scenarios, such as perfect mixing of a population and homogeneity of the individuals in the population, e.g. in terms of their susceptibility and infectiousness. The inadequacies of such descriptions have long been realized and much attention has been directed towards understanding the impact of heterogeneities in transmission on the basic reproductive number. Early efforts included accounting for differing activity levels amongst the population and various mixing 10 patterns of the population (e.g. proportionate/random mixing, assortative and 11 disassortative mixing). Much of this work was prompted by the heterogeneities 12 known to exist for the spread of sexually transmitted infections, notably gonor-13 rhea and HIV (Nold (1980); Hethcote and Yorke (1984); Anderson et al. (1986); 14 May and Anderson (1987); Jacquez et al. (1988); Gupta et al. (1989)). In the 15 context of vector-borne diseases, it has long been realized that vectors' bites 16 are not distributed uniformly across hosts; instead, there is a heterogeneity in 17 hosts' attractiveness to vectors, with some individuals being disproportionately 18 favored to receive bites (Carnevale et al. (1978); Dye and Hasibeder (1986); De 19 Benedicitis et al. (2003); Liebman et al. (2014)). 20

A now standard argument shows that the basic reproductive number for a multi-type transmission system can be calculated as the dominant eigenvalue of the next generation matrix (Diekmann and Heesterbeek (2000)). For an *n*-type setting, the next generation matrix is an *n* by *n* non-negative matrix whose (i, j)'th entry gives the average number of secondary infections of type *i* caused

by a type j individual in an otherwise entirely susceptible population. Conse-26 quently, much attention has been directed towards those special cases of het-27 erogeneous transmission that lead to next generation matrices whose dominant 28 eigenvalue is analytically tractable and hence for which the basic reproductive 29 number can be calculated explicitly. In the context of spatial heterogeneity, 30 these include symmetric spatial configurations such as equally-sized patches 31 with all-to-all or nearest neighbor contacts (see, for example Lloyd and May 32 (1996)).33

More generally, a commonly-studied situation involves separable transmission (Diekmann and Heesterbeek (2000)), where each group has a susceptibility, a_i and an infectiousness, b_i . In this case, the next generation matrix is of rank one, and, taking groups to be of equal sizes, has entries $a_i b_j/n$, and dominant eigenvalue

$$R_0 = \frac{1}{n} \sum_{i=1}^n a_i b_i.$$
 (1)

³⁹ Using the result

$$E(XY) = E(X)E(Y) + Cov(X,Y)$$
(2)

for the expectation of a product of random variables, eqn (1) can be rearranged into the following well-known formula (Dietz (1980); Dye and Hasibeder (1986)) that sheds insight into the impact of heterogeneity on R_0 in this separable setting:

$$R_{0} = \bar{a}\bar{b}\left(1 + r_{ab}\mathrm{CV}_{a}\mathrm{CV}_{b}\right)$$
$$= R_{0}^{\mathrm{hom}}\left(1 + r_{ab}\mathrm{CV}_{a}\mathrm{CV}_{b}\right). \tag{3}$$

Here, \bar{a} and b denote the average values of a_i and b_i , r_{ab} denotes the Pearson product-moment correlation coefficient between the a_i and b_i , CV_a and CV_b denote the coefficients of variation (i.e. standard deviation divided by the mean) of a_i and b_i and R_0^{hom} denotes the value of R_0 that would be predicted if the heterogeneity was ignored, i.e. the average values of a_i and b_i were used. We ⁴⁹ emphasize that these results are exact, holding for arbitrary distributions of the ⁵⁰ a_i and b_i .

Heterogeneity can inflate or deflate the value of R_0 , depending on whether there is positive or negative correlation between susceptibility and infectivity across the groups (Dietz, 1980). In the special case where susceptibility and infectivity are proportional, e.g. for a situation such as differing activity levels or mosquito biting preferences where the heterogeneity impacts both susceptibility and infectiousness in the same way, the formula reduces to

$$R_0 = R_0^{\text{hom}} \left(1 + \text{CV}_a^2 \right)$$
$$= R_0^{\text{hom}} \left(1 + \frac{\text{Var}(a)}{\bar{a}^2} \right).$$
(4)

This formula has appeared in the literature numerous times in a number of
different settings and guises (Dietz, 1980; Dye and Hasibeder, 1986; May and
Anderson, 1987).

Particularly with the increasing realization that many systems are subject 60 to multiple, often coupled, heterogeneities (Paull et al., 2012; Vazquez-Prokopec 61 et al., 2016), an important question is whether results such as eqns. (3) and (4) 62 generalize to situations in which there are more than two heterogeneities. In 63 this paper, we show that the answer to this question is no: the effect of multiple 64 interacting heterogeneities on the basic reproductive number depends on the 65 details of the distributions of the heterogeneities, in contrast to what occurs in 66 the two-heterogeneity setting. We provide results for both multivariate normal 67 and multivariate log-normal distributions of heterogeneities and demonstrate 68 that the two settings can give markedly different results. 69

70 2. Results

⁷¹ We assume that there are n types of individuals, resulting from N different ⁷² heterogeneities, N_1 of which impact susceptibility and N_2 of which impact in-⁷³ fectiousness. We further assume that the susceptibility of a type i individual

> ⁷⁴ can be written as the product $x_i^1 x_i^2 \cdots x^{N_1}$, taken over the heterogeneities that ⁷⁵ impact susceptibility, and that the infectivity of a type j individual can sim-⁷⁶ ilarly be written as $x_j^1 \cdots x_j^{N_2}$. Taking group sizes to be equal and assuming ⁷⁷ separable transmission, the entries of the next generation matrix will have the ⁷⁸ form $x_i^1 x_i^2 \dots x_i^{N_1} x_j^1 \cdots x_j^{N_2}/n$, where $N_1 + N_2 = N$. This matrix is of rank one ⁷⁹ and has dominant eigenvalue given by

$$R_0 = \frac{1}{n} \sum_{i=1}^n x_i^1 x_i^2 \cdots x_i^N.$$
 (5)

As explained above, the well-known result arises from the ability to express the expectation of the product of a pair of random variables in terms of their two expectations and their covariance. Extension of the result requires corresponding manipulations of expectations of products of three or more random variables—the so-called product moments of the joint distribution.

85 2.1. Analytic Results

In the case of a set of random variables whose joint distribution is multivariate normal, numerous authors have obtained results for product moments (see, for example, Isserlis (1918), Bendat and Piersol (1966), Bär and Dittrich (1971) and Song and Lee (2015)). For instance, in the four dimensional case we have (Bendat and Piersol (1966) and Bär and Dittrich (1971))

$$E(X_1X_2X_3X_4) = E(X_1X_2)E(X_3X_4) + E(X_1X_3)E(X_2X_4) + E(X_1X_4)E(X_2X_3) - 2E(X_1)E(X_2)E(X_3)E(X_4).$$
(6)

⁹¹ The expectations of pairwise products can be rewritten in the way described
⁹² above to give

$$E(X_{1}X_{2}X_{3}X_{4}) = E(X_{1})E(X_{2})E(X_{3})E(X_{4}) \times \left\{ (1 + r_{X_{1},X_{2}}CV_{X_{1}}CV_{X_{2}}) (1 + r_{X_{3},X_{4}}CV_{X_{3}}CV_{X_{4}}) + (1 + r_{X_{1},X_{3}}CV_{X_{1}}CV_{X_{3}}) (1 + r_{X_{2},X_{4}}CV_{X_{2}}CV_{X_{4}}) + (1 + r_{X_{1},X_{4}}CV_{X_{1}}CV_{X_{4}}) (1 + r_{X_{2},X_{3}}CV_{X_{2}}CV_{X_{3}}) - 2 \right\}.$$
(7)

- $_{\rm 93}$ We remark that the case of three random variables can be obtained by setting
- 94 $X_4 = 1.$
- For a set of N multivariate lognormally distributed random variables, prod-
- ⁹⁶ uct moments are given by the formula (Kotz et al. (2000))

$$\operatorname{E}\left(\prod_{j=1}^{N} X_{j}^{r_{j}}\right) = \exp\left(\boldsymbol{r}^{\mathrm{T}}\boldsymbol{\xi} + \frac{1}{2}\boldsymbol{r}^{\mathrm{T}}V\boldsymbol{r}\right),\tag{8}$$

where $\boldsymbol{\xi}$ and V are the mean and variance of the corresponding multivariate normal distribution. Some simple manipulation leads to

$$E(X_{1}X_{2}X_{3}X_{4}) = E(X_{1})E(X_{2})E(X_{3})E(X_{4}) \times (1 + r_{X_{1},X_{2}}CV_{X_{1}}CV_{X_{2}})(1 + r_{X_{1},X_{3}}CV_{X_{1}}CV_{X_{3}}) \times (1 + r_{X_{1},X_{4}}CV_{X_{1}}CV_{X_{4}})(1 + r_{X_{2},X_{3}}CV_{X_{2}}CV_{X_{3}}) \times (1 + r_{X_{2},X_{4}}CV_{X_{2}}CV_{X_{4}})(1 + r_{X_{3},X_{4}}CV_{X_{3}}CV_{X_{4}}).$$
(9)

Given that eqns (7) and (9) differ, and that their reduced forms when $X_4 = 1$ also differ, we have shown that there is no general formula of this type for the basic reproductive number when there are three or more coupled heterogeneities. We do notice, however, that the two formulae give approximately equal results in the limit of small coefficients of variation, *i.e.* when one can ignore products involving two or more pairs of coefficients of variation.

The majority of papers in the literature that provide analytic results for the basic reproductive number under heterogeneity focus on at most two coupled heterogeneities. One notable exception is the work of Koella (1991), which provides—without proof or qualification for its applicability—the following formula for a vector-borne pathogen subject to heterogeneities in mosquito biting rate, a, human susceptibility, b, and duration of human infection, ρ

$$R_0 = R_0^{\text{hom}} \left[1 + \frac{\text{Var}(a)}{\bar{a}^2} + 2\frac{\text{Cov}(a,b)}{\bar{a}\bar{b}} + 2\frac{\text{Cov}(a,\rho)}{\bar{a}\bar{\rho}} + \frac{\text{Cov}(\rho,b)}{\bar{\rho}\bar{b}} \right].$$
(10)

Note that the single biological heterogeneity in biting rate impacts both infectiousness and susceptibility, resulting in it being treated as two perfectly
correlated heterogeneities.

We note that equation (10) has no terms that involve products of pairs of 114 covariances (or, in the language of the earlier formulae, correlation coefficients). 115 As in the remark above comparing results between multivariate normal and 116 lognormal distributions, this formula should, in general, be seen as an approx-117 imation that is likely most accurate when coefficients of variation are small 118 (*i.e.* the heterogeneities are relatively minor). As a comment that is germane 119 to a numerical example shown below, we remark that the Koella formula does 120 agree with the result for the multivariate normal distribution, eqn (7), if the 121 coefficient of variation describing either human susceptibility or the duration of 122 human infection is equal to zero. 123

124 2.2. Numerical Results

We illustrate the above results using numerical simulation, allowing us to 125 explore the differences between predictions made using the formulae for the two 126 distributions and also using the formula in the small coefficient of variation limit. 127 For concreteness, we place these simulations within the vector-host setting de-128 scribed by Koella (1991), but for simplicity we hold one of the factors constant. 129 Specifically, hosts differ in their attractiveness to mosquitoes, impacting their 130 susceptibility and infectiousness (thus treated as two perfectly correlated het-131 erogeneities, X_1 and X_2 , within our framework), and also in their durations of 132 infection, X_3 . Setting $X_4 = 1$ and taking $X_2 = X_1$, we obtain the following two 133 formulae: 134

$$R_0 = \mathcal{E}(X_1)^2 \mathcal{E}(X_3) \left\{ 1 + \mathcal{C} \mathcal{V}_{X_1}^2 + 2r_{X_1, X_3} \mathcal{C} \mathcal{V}_{X_1} \mathcal{C} \mathcal{V}_{X_3} \right\}$$
(11)

¹³⁵ for bivariate normally distributed heterogeneities, and

$$R_0 = \mathcal{E}(X_1)^2 \mathcal{E}(X_3) \left\{ 1 + \mathcal{C} \mathcal{V}_{X_1}^2 \right\} \left\{ 1 + r_{X_1, X_3} \mathcal{C} \mathcal{V}_{X_1} \mathcal{C} \mathcal{V}_{X_3} \right\}^2$$
(12)

for bivariate lognormally distributed heterogeneities. We notice that in this reduced setting of $X_4 = 1$, the first of these formulae coincides with the small coefficient of variation limit of the two general formulae, and, as discussed above, also agrees with the Koella formula. Furthermore, we see that when the correlation coefficient, r_{X_1,X_3} , between the two heterogeneities is zero, the two

¹⁴¹ formulae are identical.

142 2.2.1. Bivariate Normal Distribution

Figure (1) illustrates the performance of eqn (11) when the group attributes 143 X_1 and X_3 are sampled from a bivariate normal distribution. Parameter values 144 (given in the figure caption) were chosen for illustrative purposes and are not 145 intended to represent a specific real-world infection. For each of a thousand 146 replicates, either ten (panel a) or a thousand (panel b) pairs of values of biting 147 rate and duration of human infection were sampled from a bivariate normal 148 distribution. The actual R_0 value for each replicate, calculated from eqn (5), is 149 plotted against the value of R_0 predicted for that replicate by the MVN formula, 150 eqn (11). In addition, we show (using a red square) the R_0 value that would be 151 predicted if there was no heterogeneity (*i.e.* X_1 and X_3 are set equal to their 152 respective average values) and the value predicted by the MVN formula for the 153 underlying MVN distribution (green star). 154

To aid comparison, the 45° diagonal line is shown on both plots: deviations 155 from these lines represent deviations from the values predicted by the MVN 156 formula. We quantitate these deviations by calculating the coefficient of deter-157 mination, R^2 , in a way that is familiar from regression theory. In both cases, 158 the R^2 value is high, and approximately equal to 0.99. The numerically calcu-159 lated values of R_0 fall in a cloud centered on the value predicted by the MVN 160 formula, with the size of the cloud being smaller for the panel resulting from the 161 larger number of groups. The deviations here result from sampling error, with 162 samples drawn from the bivariate normal not being perfectly representative of 163 the entire distribution. This effect is more pointed when there are fewer samples 164 (*i.e.* fewer groups), with the cloud of points shrinking as the number of groups 165 increases. In fact, since the basic reproductive number, eqn (5), is calculated 166 as the average of a sequence of independent, identically distributed quantities, 167 the central limit theorem can be used to quantify the variation seen about the 168

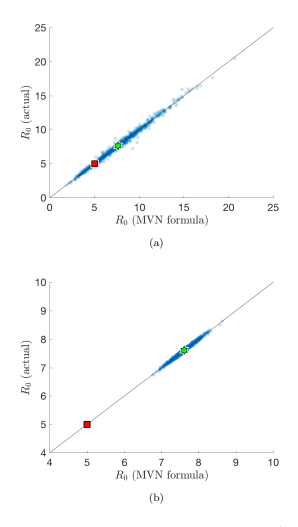


Figure 1: Comparison of R_0 values calculated numerically (using eqn 5) with those predicted by the MVN formula (eqn 11) for heterogeneities distributed according to an MVN distribution. Each blue circle represents the values of R_0 obtained for (panel a) a ten group model or (panel b) a thousand group model with pairs of values of biting rate, X_1 , and average duration of human infection X_3 drawn from a bivariate normal distribution with means 1 and 5, respectively, variances $Var(X_1) = 0.2$ and $Var(X_3) = 4$, and correlation $r_{X_1,X_2} = 0.9$. The value of the susceptibility parameter was fixed at 1 for each group. The red square denotes the value of R_0 if there was no heterogeneity, *i.e.* obtained using the average values, R_0^{hom} , while the green star denotes the value of R_0 calculated using the MVN formula using the means, variances and covariances of the underlying bivariate normal distribution. As described in the text, the predictive ability of the MVN formula is measured by the coefficient of determination, R^2 , and equals: (panel a) 0.991 and (panel b) 0.988. Note the different scales on the axes between the two panels.

central predicted value. We remark that a corresponding figure generated in the
case of two coupled heterogeneities would exhibit no deviation from the diagonal line as eqn (3) is exact: it does not rely on any distributional assumption of
heterogeneities across groups.

173 2.2.2. Bivariate Lognormal Distribution

Use of a multivariate lognormal distribution allows us to explore settings in which the components of transmission exhibit more severe heterogeneity and to assess the extent to which the impact of such heterogeneities are misrepresented by either the small coefficient of variation formula or by the formula that pertains in the multivariate normal case.

In figure (2), the biting rates and durations of infectiousness are drawn from 179 independent lognormal distributions, with means 1 and 3, and variances 1 and 180 8, respectively. We compare the performance of the MVN formula (panel a) 181 and MVLN formula (panel b) for a 1000-group setting (*i.e.* 1000 samples are 182 drawn from the distributions). In this case, because there is zero correlation 183 between the two heterogeneities, the population-level predictions of the MVN 184 and MVLN formulae (*i.e.* the values obtained using the moments of the un-185 derlying distribution) are identical (shown by a yellow diamond). We see that 186 the MVLN formula provides a better description $(R^2 = 0.748 \text{ using the MVN})$ 187 formula, while $R^2 = 0.852$ using the MVLN formula), which is to be expected 188 given that samples were drawn from an MVLN distribution. 189

Figure (3) explores a situation in which there is a positive correlation (r =190 0.6) between the two components, *i.e.* the biting rate and duration of infectious-191 ness, of the bivariate lognormal distribution. We see that the MVLN formula 192 performs well (panel b, $R^2 = 0.852$). The positive correlation leads to the MVN 193 formula underestimating R_0 compared to the MVLN formula, for example as 194 witnessed by the central estimates of R_0 (yellow diamond: MVLN, green star: 195 MVN). The MVN formula consistently provides a large underestimate of the 196 true value of R_0 (panel a), and its predictive ability is poor (the negative value 197 of R^2 , -0.596, indicates that the formula performs worse on these points than a 198

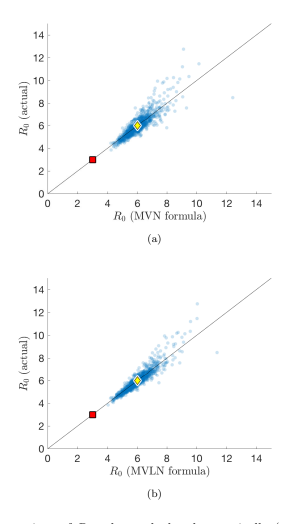


Figure 2: Comparison of R_0 values calculated numerically (using eqn 5) with those predicted by the formula derived from the MVN distribution (eqn 11) (panel a) and those predicted by the formula derived from the MVLN distribution (eqn 12) (panel b). Each blue circle represents the values of R_0 obtained for a thousand group model with values of biting rate, X_1 , and average duration of human infection, X_3 , drawn from independent lognormal distributions (means 1 and 3, variances 1 and 8, respectively). The value of the susceptibility parameter was fixed at 1 for each group. The red square denotes the value of R_0 obtained using the average values, R_0^{hom} , while the yellow diamond denotes the value of R_0 calculated using the MVLN formula using the means, variances and covariances of the bivariate lognormal distribution. Because the distributions of X_1 and X_3 are assumed to be independent in this figure, the populationlevel predictions of the MVLN and MVN formulae are identical. The predictive ability of the MVLN formula (panel b, $R^2 = 0.852$) is greater than that of the MVN formula (panel a, $R^2 = 0.748$), as should be expected given that draws were made from an MVLN distribution.

¹⁹⁹ constant predictor).

Finally, we consider a setting in which the two heterogeneities are negatively 200 correlated, with r = -0.2. Figure (4) shows that in this case, the MVLN 201 formula correctly predicts lower values of R_0 than does the MVN formula (e.g. 202 compare the locations of the yellow diamond and the green star, obtained from 203 MVLN and MVN formulae, respectively, using the moments of the underlying 204 distribution). The MVLN formula provides reasonable predictions ($R^2 = 0.735$), 205 whereas the MVN formula $(R^2 = -4.34)$ consistently overestimates the value 206 of R_0 , and often by a considerable amount. 207

208 3. Discussion

In this paper we have shown that the well-known result for the impact of two 209 coupled heterogeneities on the basic reproductive number of an epidemiological 210 system under separable transmission does not have a general counterpart when 211 there are three or more coupled heterogeneities. In the more general setting, 212 the formula for the basic reproductive number depends on details of the joint 213 distribution of the heterogeneities in a way that is quite different than in the 214 setting with two heterogeneities. We were able to derive formulae that related 215 the basic reproductive number to the magnitudes of the heterogeneities and 216 their pairwise correlations for the special cases of multivariate normal and mul-217 tivariate lognormal distributions of heterogeneities. Under particular limiting 218 cases (typically in the limit of low levels of heterogeneity), the two formulae give 219 similar predictions. We showed that an earlier result in the literature (Koella, 220 1991) should be viewed as an approximate result, although we noted that in ap-221 propriate limiting cases, the result agrees with our formula for the multivariate 222 normal distribution. 223

Given the reliance of the R_0 formulae obtained here on the joint distribution of the heterogeneities, our results are only exact as the number of groups in the multi-type model approaches infinity. For a finite number of groups, where the levels of the heterogeneities across groups are sampled from the underlying

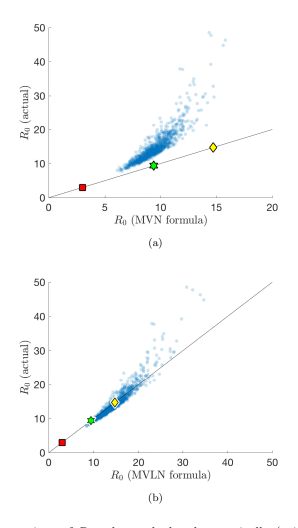


Figure 3: Comparison of R_0 values calculated numerically (using eqn 5) with those predicted by the formula derived from the MVN distribution (eqn 11) (panel a) and those predicted by the formula derived from the MVLN distribution (eqn 12) (panel b). Details are as in Figure (2), except that here there is a positive correlation, r = 0.6, between the two components of the bivariate lognormal distribution. In this case, the population-level predictions of the MVN and MVLN formulae (green star and yellow diamond, respectively) differ. The predictive ability of the MVLN formula (panel b, $R^2 = 0.828$) is greater than that of the MVN formula (panel a, $R^2 = -0.596$). The MVN formula performs worse than a constant predictor and consistently provides a large underestimate of the true value of R_0 .

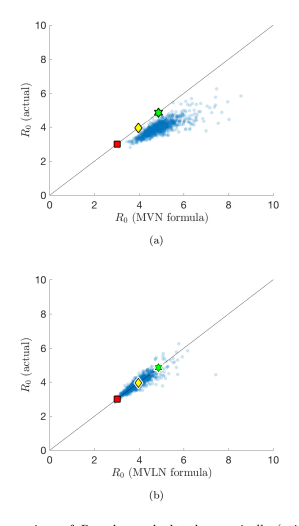


Figure 4: Comparison of R_0 values calculated numerically (using eqn 5) with those predicted by the formula derived from the MVN distribution (eqn 11) (panel a) and those predicted by the formula derived from the MVLN distribution (eqn 12) (panel b). Details are as in Figures (2) and (3), except that here there is a negative correlation, r = -0.2, between the two components of the bivariate lognormal distribution. Again, the predictive ability of the MVLN formula (panel b, $R^2 = 0.721$) is greater than that of the MVN formula (panel a, $R^2 = -4.91$), with the latter performing worse than a constant predictor and consistently providing a large overestimate of the true value of R_0 .

distribution, the predictions made by these formulae are not perfect. This again
is in contrast to the two heterogeneity setting, in which the well-known result
is exact.

Although theoretical attention has typically focused on the two heterogeneity 231 case, and this has provided much insight, heterogeneous transmission in the real 232 world typically involves more than two factors (Paull et al. (2012); Vazquez-233 Prokopec et al. (2016)). As such, it is important to gain understanding of 234 how multiple coupled heterogeneities impact transmission and the limitations of 235 general results that can be obtained in such more realistic settings. This paper 236 provides a theoretical step in that direction and aims to guide more detailed 237 studies that involve numerical exploration of specific situations, yielding further 238 insights into the epidemiological role of individual variability. 239

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