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# Assessing plant pathogen infection rates in natural soils using R

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The potential of soils to naturally suppress inherent plant pathogens is an important ecosystem function. Usually, pathogen infection assays are used for estimating the suppressive potential of soils. In natural soils, however, co-occurring pathogens might simultaneously infect plants complicating the estimation of a focal pathogen's infection rate as a measure of soil suppressiveness. Here, we present a method in **R** correcting for these unwanted effects by developing a two pathogen mono-molecular infection model. We fit the two pathogen mono-molecular infection model to data by using an integrative approach combining a numerical simulation of the model with an iterative maximum likelihood fit. We show that in presence of co-occurring pathogens uncorrected data critically under- respectively overestimate soil suppressiveness measures. In contrast, our new approach enables to precisely estimate soil suppressiveness measures such as plant infection parameters that is necessary in case different pathogens are present. We propose our method to be particularly useful for exploring soil suppressiveness of natural soils from different sites (e.g., in biodiversity experiments).

### Assessing Plant Pathogen Infection Rates in Natural Soils using R

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#### **ABSTRACT**

The potential of soils to naturally suppress inherent plant pathogens is an important ecosystem function. 12 Usually, pathogen infection assays are used for estimating the suppressive potential of soils. In natural 13 soils, however, co-occurring pathogens might simultaneously infect plants complicating the estimation of 14 a focal pathogen's infection rate as a measure of soil suppressiveness. Here, we present a method in R 15 correcting for these unwanted effects by developing a two pathogen mono-molecular infection model. We 16 fit the two pathogen mono-molecular infection model to data by using an integrative approach combining 17 a numerical simulation of the model with an iterative maximum likelihood fit. We show that in presence of 18 co-occurring pathogens uncorrected data critically under- respectively overestimate soil suppressiveness 19 measures. In contrast, our new approach enables to precisely estimate soil suppressiveness measures 20 21 such as plant infection rate and plant resistance time. Our method allows a correction of measured infection parameters that is necessary in case different pathogens are present. We propose our method 22 to be particularly useful for exploring soil suppressiveness of natural soils from different sites (e.g., in 23 biodiversity experiments). 24

Keywords: infected control treatments, maximum likelihood estimation, ordinary differential equation,
 numerical simulation, biodiversity, soil resistance, bioassay, R, bbmle, deSolve, Manual

#### <sup>20</sup> numerical simulation, biodiversity, son resistance, bioassay, R, bonne, desorv

#### 27 INTRODUCTION

Pathogen infection assays are a standard method for estimating plant resistance to pathogens, induced 28 systemic resistance in plants, the effect of artificial or natural plant protectants (e.g. plant beneficial 29 bacteria), and a soil's suppressive potential. Such bioassays compose of a soil or substrate inoculated with 30 a pathogen and a pathogen sensitive plant, and data is collected at just a single point in time (Maurhofer 31 et al., 1994; Pierson and Weller, 1994; Postma et al., 2008) or at multiple points in time (e.g. Postma et al., 32 2008; Hanse et al., 2011; Latz et al., 2012, 2016). Remarkably, in the latter case often only one single 33 point in time is chosen for evaluation (e.g. Postma et al., 2008; Hanse et al., 2011; Latz et al., 2012), or 34 the increase from one to the next point in time is evaluated (Kushalappa and Ludwig, 1982). However, 35 disease progression is more precisely described by classical growth curve models (Neher and Campbell, 36 1992). Out of the plethora of growth models (Paine et al., 2012), the mono-molecular model has often 37 been used to describe bioassays with soil-borne pathogens (Stanghellini et al., 2004; Wilson et al., 2008). 38 The mono-molecular infection model describes the disease progression (the change of infections over 39 time) with an initial linear increase of infections (the infection rate), followed by a saturation (given by 40 the maximum number of infectable plants, also known as carrying capacity or asymptotic growth). 41 The infection rate was suggested to be the most important parameter for determining pathogenicity 42 (Raaijmakers et al., 2009). However, when estimating a soil's suppressive potential, the time until 43

- infections occur (resistance time) might be even more important since pathogen inhibition occurs largely
- <sup>44</sup> Infections occur (resistance time) high be even more important since pathogen minimum occurs largery <sup>45</sup> during pathogen growth. Actually, only a few experimental setups allow the investigation of both, infection
- <sup>46</sup> rate and resistance time. To measure an infection rate it is necessary to use a system with multiple plant



**Figure 1.** Two different possible setups for infection treatments. The circular setup with a centered pathogen surrounded by plants (a) may lead to a steep linear infection scenario as all plants are probably infected by the source pathogen at more or less the same time. Only the linear spatial assembly (b) allows for a consecutive infection of plants resulting in a linear increase that can be modeled by the mono-molecular infection model.

47 individuals (Figure 1) where plants can be infected one after another (i.e. measuring a time-series). In such experiments, the pathogen inoculant can be applied in different ways: (i) equally distributed 48 application, i.e. homogeneously mixed in the soil or growth-substrate, or (ii) single point application 49 (where pathogen spread can be assessed; Figure 1). If a pathogen is homogeneously distributed in the 50 plant growth substrate, it is possible to measure the number of infected plants over time. The measured 51 infection rate, however, would not represent the infection rate per se but rather the resistance variance of 52 the plant community to the pathogen. The same problem occurs if a pathogen is applied to one location in 53 the substrate and plants are planted at equal distances around the inoculum (Figure 1A). Linear spatial 54 designs (Figure 1B) have the potential to estimate the correct infection rate in addition to the resistance 55 time, whereas the further mentioned approaches solely allow to estimate the resistance time. Hence, it is 56 important to keep in mind that the design determines the hypothesis that can be tested. Another difficulty 57 in performing pathogen infection assays occurs if natural field soils are used as substrate (e.g. Mendes 58 et al., 2011; Latz et al., 2012, 2016). Here, in addition to the applied pathogen, other unknown pathogens 59 may already exist in the soil and may increase infection in the plants. To cope with this problem, control 60 treatments may be used to reveal the occurrence of natural soil inhabiting pathogens. If controls show 61 infections, (i) these infections might be ignored if they are evaluated as statistically not relevant (Fig. 2A), 62 (ii) the treatments where the corresponding controls showed infections may be excluded from further 63 64 analyses (Fig. 2B), (iii) the treatments may be linearly corrected by simply subtracting the total amount of infectable plants by the infections that occurred in the control (Fig. 2C). The third approach may 65 lead to erroneous results in non-linear analyses as shown for functional response models (McCoy et al., 66 2012). However, none of these approaches are desirable as they may lead to a bias in single infection rate 67 measures (due to ignoring or wrongly correcting infections of a naturally occurring pathogen) and the 68 loss of data (exclusion of treatments where the corresponding control was infected). 69

Here, we present an alternative approach that incorporates infections caused by any additional 70 pathogens in the system by using a two pathogen mono-molecular infection model inspired by the 71 competition model for logistic growth (Lotka, 1925; Volterra, 1926). This two pathogen mono-molecular 72 model is an ordinary differential equation system with two equations. Systems with two equations are 73 hardly analytically integrable to a single equation describing the progress of infections over time, therewith 74 75 preventing the use of standard linear or non-linear fitting algorithms. To overcome this limitation, we applied a numerical integration routine (Soetaert et al., 2010) combined with a maximum likelihood 76 optimizer (Bolker and Team, 2016) to fit our model to data. Our method allows for the use of natural soils 77 (i) already contaminated with naturally occurring pathogens, and (ii) from different origins and habitats, 78 while allowing for accurate evaluation of pathogenicity and plant resistance patterns in the field. 79



**Figure 2.** Known practices to deal with infections observed in control treatments. Number of infected plants at each time-point in a single, independent pot. Grey dots: control pots (without having added the pathogen); black dots: treatment pots (with having added the pathogen). (a) Infected controls are ignored and treatment data remains uncorrected. (b) In case the control pot showed an infection the respective treatment data is excluded (red crosses). (c) The treatment data is "corrected" by subtracting the number of infections in the control from the number of infections in the treatment (red dots, note that may lead to some negative infections).

#### 80 METHODS

#### 81 Simulations

We solved the differential equation systems (eqn. (2) & (3)) using the lsoda()-function (version 82 1.13; references: Soetaert and Herman, 2008; Soetaert et al., 2010) in R (R Core Team, 2016). The 83 time-series length was set to 30 days with a temporal resolution of 0.01 days.  $I_{max}$  was fixed to 10 84 plants. We simulated two different scenarios; scenario 1: the natural pathogen has lower infection 85 rates  $(0.001 \le r_{control} \le 0.1; 0.1 \le r_{treatment} \le 0.5)$  and occurs earlier in the time series as the treatment 86 pathogen ( $1 \le t_{0_{control}} \le 5$ ;  $5 \le t_{0_{treatment}} \le 10$ ); and scenario 2: the natural pathogen has comparable 87 infection rates  $(0.01 \le r_{control} = 0.1; 0.01 \le r_{treatment} \le 0.1)$  to the experimentally added pathogen and 88 occurs later in the time series ( $5 \le t_{0_{control}} \le 10$ ;  $1 \le t_{0_{treatment}} \le 5$ ). We draw all infection rates, r, and time 89 of first infections,  $t_0$ , from uniform distributions. 90

After simulating the time series, we sampled randomly four data-points for each full time-point (i.e. 91 t = 1, 2, ..., 30) assuming a binomial distribution with a size of  $I_{max}$  and a probability of the simulated 92 number of infections at time t divided by  $I_{max}$  resulting in 120 independent data points for each simulated 93 infection assay. Additionally simulated one or four consecutive time series resulting in 30 data points 94 of one experimental unit (temporal autocorrelated) and 120 of four experimental units (each time series 95 contains 30 temporal autocorrelated data points). We repeated this simulation of data 1100 times for each 96 scenario. We excluded model fits for both, the one-pathogen model and the two-pathogen model, if the 97 fitting of one or the other failed and used the first 1000 results of the cleaned data set. 98

#### 99 Statistical analyses

We analyzed the simulated data using an iterative maximum likelihood algorithm (function mle2 () from the package **bbmle** version 1.0.18; references: Bolker, 2008; Bolker and Team, 2016) to fit equations (2) & (3) to the data using R (R version 3.3.0; reference: R Core Team, 2016). See the supplemental manual for an in-depth description of the methodology.

We saved all results for the one-pathogen model fitting and the two-pathogen model fitting for each scenario and each setting (independent, one time series and four time series). Subsequently, we analyzed the  $log_{10}$ -ratio of the fitted parameters to the initially simulated values. The starting values for infection rates where set to 50% of the simulated value and for resistance time to 75% of the simulated value. In scenario 2, the starting values for infection rates where set to 50% of the simulated value and for resistance time to 0.5 days of the simulated value.

#### 110 Additional Files

#### 111 Additional file 1 — MainSources.zip

<sup>112</sup> This compressed folder contains the sub-folders "data", "script" and "source". The folder "data" contains

the data to reproduce figure 4, "data\_scenario01.csv" & "data\_scenario02.csv" shown in figure 4. The

<sup>114</sup> folder "script" contains the script files "scenario01.r" to "scenario06.r" that allow for reproducing the

data shown in figure 4. The folder "source" contains the R-source files "infections.models.r" and

<sup>116</sup> "infection.nll.r" that are required to run the script files.

#### 117 Additional file 2 — manual-assessing-plant.pdf

- <sup>118</sup> This document includes an in-depth description on how to apply the method presented in this study in R.
- <sup>119</sup> Including how to create regression lines, trouble shooting, how to use the functions if there are different
- <sup>120</sup> Imax, and an in-depth description of the source files.

#### 121 Additional file 3 — ManualSources.zip

- <sup>122</sup> This compressed folder includes all necessary data, scripts and source files to reproduce the statistics and
- 123 plots from the manual.

#### 124 **RESULTS AND DISCUSSION**

#### 125 The Model

The mono-molecular infection model (Raaijmakers et al., 2009; Paine et al., 2012) describes the increase of infections in a (plant) community over time,  $dIdt^{-1}$ , by:

$$\frac{dI}{dt} = r\left(I_{max} - I\right) \tag{1}$$

with r [time<sup>-1</sup>] being the infection rate and  $I_{max}$  [Infected (Plants) Area<sup>-1</sup>] being the maximum number of potentially infectable plants.

The infection of the first plant is not necessarily instantaneous, but depends on the resistance of the soil and the plants to the pathogen, leading to a lag phase at the beginning of the experiment. To account for this mechanism, we extend the mono-molecular infection model by the resistance time,  $t_0$ :

$$\frac{dI}{dt} = \begin{cases} 0 & \text{if } t < t_0 \\ r(I_{max} - I) & \text{if } t \ge t_0. \end{cases}$$

$$\tag{2}$$

Below  $t_0$  new infections are zero whilst above, the occurrence of new infections follows the monomolecular infection model. We will refer to this model as one-pathogen model (Fig. 3A, B).

In experiments using natural soils, natural occurring pathogens may be responsible for additional infections during the experimental trial. To correct for those infections, we extend the one-pathogen model to a two-species mono-molecular infection model, inspired by the two-species competition growth model (Lotka, 1925; Volterra, 1926):

$$\frac{dI_p}{dt} = \begin{cases} 0 & \text{if } t < t_{0_p} \\ r_p \left( I_{max} - (I_p + I_c) \right) & \text{if } t \ge t_{0_p}, \end{cases}$$

$$\frac{dI_c}{dt} = \begin{cases} 0 & \text{if } t < t_{0_c} \\ r_c \left( I_{max} - (I_p + I_c) \right) & \text{if } t \ge t_{0_c}, \end{cases}$$
(3)

where  $I_p$  is the number of infected plants due to the pathogen,  $I_c$  is the number of infected plants in the control;  $r_p$  and  $r_c$  are the infection rates of the pathogen and the control treatment, respectively; and  $t_{0_p}$  and  $t_{0_c}$  are the resistance times of the pathogen and the control treatment, respectively. We will refer to this model as two-pathogen model.

Below, we will give two examples of different model-parameter combinations, based on two different biological examples that might lead to two different misleading fitting results if the one-pathogen model is used in case of contaminated pots.

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**Figure 3.** Different model configurations. (a) The one-pathogen model with 2 different settings of parameter values (light red line:  $r = 0.01 \& t_0 = 1$ ; blue line:  $r = 0.1 \& t_0 = 8$ ). (b) The two-pathogen model (black line) incorporates the parameter values of (a) and lies slightly above the one-pathogen model (blue line in (a)); hypothetically, using a one-pathogen model to fit the black line will result in a different parameter estimation (dashed orange line). (c) The one-pathogen model with 2 different settings of parameter values (light red line:  $r = 0.05 \& t_0 = 8$ ; blue line:  $r = 0.05 \& t_0 = 2$ ). (d) The two-pathogen model (black line) incorporates the parameter values of (c) and lies only above the one-pathogen model (blue line in (c)) at a late stage of the experiment, hypothetically using a one-pathogen model to fit may result in the orange model fit.

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First, we assume a high infection rate r, and an experimental pot showing a high resistance time, 137 This will result in a first half of the experiment without any infections while in the second half of  $t_0$ . 138 the experiment the plants will become infected rapidly (Fig. 3A, blue line). We interpret in this case 139 an experimentally added pathogen (treatment pathogen) being inoculated in a defined distance to the 140 seedlings, a soil showing high suppressivess and/or highly resistant plants (high resistance time), but the 141 pathogen being highly abundant and able to infect plants rapidly after the first infection (high infection 142 rate). However, this scenario presumes sterile soil previous to having added a treatment pathogen, whereas 143 natural soils might be contaminated by already naturally occurring pathogens. A contaminated control pot 144 without an experimentally added pathogen may then show early infections followed by a shallow increase 145 146 of infections over time (Fig. 3A, light red line). The combined progression of the infections over time in a contaminated treatment pot is more complex than that of assuming only a treatment pathogen being 147 present, with showing a shallow increase of infections at low densities and a steep increase of infections 148 in the second half of the experiment (Fig. 3B, black line). Applying the one-pathogen model to estimate 149 the resistance time and infection rate would lead to a misleading fit (Fig. 3B, dashed orange line). 150

Second, we assume the plants having a rather small resistance time,  $t_0$ , and the pathogen being less 151 aggressive (low infection rate, r; Fig. 3C, blue line. Here, we assume a perfectly sterile experiment for 152 both, the treatment and the control. In this example, the control treatments should not show any infections 153 over time. However, pathogens could also disperse into the experimental pots during the experimental 154 trial, leading to late infections of the control (Fig. 3C, red line). This might be the case when experimental 155 pots can not be isolated from the environment, e.g. partially open mesocosms, resulting in more than the 156 treatment pathogen being responsible for infections (Fig. 3D, black line vs. Fig. 3C, blue line). Applying 157 here the one-pathogen model to estimate the infection parameters may lead to the correct estimation of the 158 resistance time but to a underestimation of the infection rate of the treatment pathogen (Fig. 3D, dashed 159 orange line). 160

In both scenarios, the use of the one-pathogen model would lead to misleading parameter estimations. To overcome this issue the two-pathogen model should be fitted to the data.

#### 163 Statistical model evaluation

#### 164 Independent data

We tested our model framework by simulating two separate scenarios (subsequently called scenario 1 165 and scenario 2). In scenario 1, naturally occurring pathogens infect seedlings earlier than the treatment 166 pathogen, but the naturally occurring pathogens are less infectious (i.e. a lower infection rate, r; Fig. 167 3A,B). In scenario 2, the naturally occurring pathogens infect the seedlings later than the treatment 168 pathogen but are similar infectious (Fig. 3C,D). We simulated 1000 data sets where each simulated data 169 point represents an independent measure (i.e. the end point of a single time series) for each scenario and 170 fitted (i) the one-pathogen model to each data set (equation (2)) and (ii) the two-pathogen model (equation 171 (3) to each data set. We compared the fitted parameter values (i.e. the infection rate, r, and the time of 172 173 first infection,  $t_0$ ) by taking the log-ratio. See methods for a detailed description of the procedure.

Using the one-pathogen model leads to a systematic underestimation of infection rates, r, (Fig. 4A, lower row) whereas the two-pathogen model performs well (Fig. 4A, upper row). Also, the resistance times,  $t_0$ , are underestimated by the one-pathogen model (Fig. 4B, lower row) whereas the two-pathogen model predicts the resistance time very precisely (Fig. 4B, upper row).

The underestimation of both the resistance times and the infection rates nicely reflect our assumptions 178 when fitting the one-pathogen model to the treatment (Fig. 3C). The real increase in infection is 179 rather strong, and coupled to a late first occurrence of infections (Fig. 5A, gray dashed line). But 180 the one-pathogen model estimates a mixed increase of both, the infections caused by the control and the 181 treatment pathogens. This means that the resistance time is driven by the control pathogen leading to 182 183 an underestimation of infection rates (Fig. 5A, orange dashed line). The two-pathogen model, however, resolves the strong non-linear interactions between the model paraemters and leads to a infection curve 184 with the correct infection rates and resistance times (Fig. 5A, blue line) that lies slightly beneath the total 185 infection (Fig. 5A, black line). 186

In the second scenario (higher resistance time for the control pathogen with similar infection rates
 for both) the one-pathogen model overestimates the infection rates systematically (Fig. 4C, lower row).
 Surprisingly, also the resistance times are overestimated (Fig. 4D, lower row) contrasting our expectations.

<sup>190</sup> In contrast, the two-pathogen model predicts the simulated parameter values precisely and outperforms



**Figure 4.** Results of the model evaluation of the one pathogen model versus the two pathogen model. The results of scenario 1 (a & b) and scenario 2 (c & d) for infections rate,  $r_p$ , (a & c) and resistance time,  $t_{0_p}$  (b & d). The  $log_{10}$ -ratio of the parameter fit to the real parameter used for simulating is given on the x-axis. If zero, the fit is perfectly reflecting the simulation, if larger than zero, the fit overestimates the real value, if smaller, the fit underestimates the real value.

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**Figure 5.** Mechanics of fitted results. (a) The two-pathogen model shows a steep increase (dashed gray line) in infections (black line) when the treatment pathogen enters the system (blue line). The controls, however, lead to infections earlier (red line) leading to a decreased increase in infections using a one-pathogen model for fitting to contaminated data (dashed orange line). (b) The infection rates are overestimate by using a one-pathogen model (orange line) to a treatment with two pathogens (black line). The real treatment infections must be lower (dashed cyan line) as only a part of the infections are caused by the treatment pathogen (blue line), the rest is caused by the control pathogen (red line). (c) The resistance time is mainly inferred by the fit using knowledge on the correct infection rates (dashed cyan line), if the infection rate is overestimated due to additional late occurring control infections (black dot) the resistance time is also overestimated (dashed orange line).

the one-pathogen model dramatically (Fig. 4C,D, upper rows). The overestimation of the infection rates 191 by the one-pathogen model can be explained by an additional boost of infections later in the experiment 192 (Fig. 5B, black line) by additional infections of the control pathogen (Fig. 5B, red line) additionally to the 193 infections of the treatment pathogen (Fig. 5B, blue line). This additional infections lead to an increase in 194 estimated infection rates (Fig. 5B, orange line) compared to the prediction of the isolated infections of the 195 treatment pathogen (Fig. 5B, cyan line). Interestingly, the resistance times are also overestimated. This is 196 a rather small effect and may caused by the fact that, if the correct resistance time lies between two time 197 steps (e.g.  $t_0 = 2.1$ ), the next full time step (e.g.  $t_0 = 3$ ) may show the first infection and the third time 198 step the second infection we expect a rather linear increase from zero to two from time step 2 to 4 (Fig. 199 5C, cyan line). If a control pathogen also causes an infection at the third time step, the fitting algorithm 200 will estimate a steeper increase to the cost of a higher estimates resistance time (that must still be below 3 201 in this example, Fig. 5C, orange line). 202

#### 203 Consecutive time-series data

For the above described model comparison we used data that consisted of independent measures. This 204 means each data point was derived from a single experimental pot that has been destructively sampled. If 205 applying this approach to an experiment running 30 days with a resolution of one measurement per day and 206 4 replicates the total amount of pots that must be maintained is 120 (as in our above described analyses). 207 Applying an additional gradient (e.g. biodiversity) would lead to a not feasible amount of experimental 208 units. To avoid such a laborious approach, most studies measure consecutive time series where data for 209 each temporal replicate originates from the same experimental unit. To test if our model approach is also 210 able to fit such data adequately we simulated: (1) data of a single time series resulting in 30 measures 211 from one experimental pot; (2) data of four time series resulting in 120 measures from four experimental 212 pots. We only applied the two-pathogen model to the simulated data. Subsequently, we compared the 213 deviations of the model fits to the original simulated parameter values and we cross-compared the quality 214 of the fits using independent data (120 measures from 120 experimental pots). 215

Fitting the model to data from a single time series in scenario 1 (Fig. 6A,B, topmost rows, naturally occurring pathogens infect the plants earlier but less strong) leads to a slight overestimation of infection rates but in average correctly estimated resistance times. Using data from four consecutive time series (Fig. 6A,B, middle rows) results in a very precise fit that is not distinguishable from the fit using independent data (Fig. 6A,B, lowermost rows). In scenario 2 (Fig. 6C,D, topmost rows, naturally occurring pathogens



**Figure 6.** Results of the model evaluation comparing independent measures and consecutive time-series data. The results of scenario 1 (a & b) and scenario 2 (c & d) for infections rate,  $r_p$ , (a & c) and resistance time,  $t_{0_p}$  (b & d). The  $log_{10}$ -ratio of the parameter fit to the real parameter used for simulating is given on the x-axis. If zero, the fit is perfectly reflecting the simulation, if larger than zero, the fit overestimates the real value, if smaller, the fit underestimates the real value.

infect the plants later but equally strong) both, the infection rate and the resistance time, are systematically 221 overestimated. Using data from four time series to estimate the parameter values statistically increases the 222 preciseness of the fit dramatically and the results do not differ significantly from the expected simulated 223 values (Fig. 6C,D, middle rows) and are only marginally worse than the results from the fit using 224 independent data (Fig. 6C,D, lowermost rows). The systematic overestimation of infection rates in both, 225 scenario 1 and scenario 2, might be reasoned by the fact that in consecutive time series the number of 226 infected plants can only increase opposing the independent measures where infection can also decrease as 227 they are results from independent time series (e.g. Fig. 2A). 228

#### 229 General discussion

In both scenarios, the two-pathogen model outclasses the one-pathogen model in predicting both, re-230 sistance time and infection rates. Moreover, our approach allows to use data from just a few (in our 231 case: four) consecutive time series reducing the number of pots to be maintained dramatically (in our 232 example 4 versus 120 pots). This reduction of experimental units also allows to investigate the suppres-233 sive potential of soils in dependence of other independent variables such as biodiversity, environmental 234 changes (e.g. a nutrient or temperature gradient), diversity and abundance of plant beneficial bacteria or 235 pesticides (see reference (Latz et al., 2016) as an example). To provide a relatively simple entry into our 236 statistical method, we provide the R-code to reproduce all data and statistics presented above. Moreover 237 we provide an in-depth manual as additional online file (see section additional files below for further 238 information). Our model approach should be easily extendable to other kinds of growth or infection 239 models (find other growth models in reference Paine et al., 2012) to e.g. describe pathogen dispersion 240 in larger plant communities or to include more than one treatment pathogen to estimate the competition 241 ability of different pathogens when used together. The statistical method presented here is also superior to 242 classical analytic approaches such as the linearization of the growth model (Neher and Campbell, 1992), 243 the estimation of infection rates by analyzing the initial increase in infections (Kushalappa and Ludwig, 244 1982), or the arbitrary selection of a single point in time (Maurhofer et al., 1994; Pierson and Weller, 245 1994; Postma et al., 2008; Hanse et al., 2011; Latz et al., 2012) as it allows (1) to analyze the complete 246 disease progression over time and (2) it allows to correct for naturally occurring pathogens. 247

#### 248 CONCLUSIONS

Keystone plants as well as diverse plant communities have shown to increase the pathogen suppressive 249 potential of soils, an effect that would vanish if soils would be sterilized. However, if standard approaches 250 or the one-pathogen infection model is applied, a sterile soil is required to prevent infections by non-251 treatment pathogens and non-sterile soils consequently prevent the correct estimation of the pathogen 252 suppressive potential of natural soils. This problem can be overcome by using the two-pathogen model 253 presented in this study as it allows for the correct estimation of infection rates and resistance times using 254 natural soils. Our method will thus enable to estimate the natural suppressive potential of soils allowing 255 to investigate how e.g. keystone plants or specifically mixed plant communities naturally contribute to a 256 soil resistance against pathogens. 257

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