Fast fitting of neural ordinary differential equations by Bayesian neural gradient matching to infer ecological interactions from time series data

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Abstract

- 1. Inferring ecological interactions is hard because we often lack suitable parametric representations to portray them. Neural ordinary differential equations (NODEs) provide a way of estimating interactions nonparametrically from time series data. NODEs, however, are slow to fit, and inferred interactions have not been compared to the truth.
- 2. We provide a fast NODE fitting method, Bayesian neural gradient matching (BNGM), which relies on interpolating time series with neural networks, and fitting NODEs to the interpolated dynamics with Bayesian regularisation. We test the accuracy of the approach by inferring ecological interactions in time series generated by an ODE model with known interactions. We compare these results against three existing approaches for estimating ecological interactions, standard NODEs, ODE models, and convergent cross mapping (CCM). We also infer interactions in experimentally replicated time series of a microcosm featuring an algae, flagellate, and rotifer population, in the hare and lynx system, and the Maizuru bay community featuring 11 species.
- 3. Our BNGM approach allows us to cut down the fitting time of NODE systems to only a few seconds and provides accurate estimates of ecological interactions in the artificial system, as true ecological interactions are estimated with minimal error. Our benchmark analysis reveals that our approach is both faster and more accurate than standard NODEs and parametric ODEs, while CCM was found to be faster but less accurate. The analysis of the replicated time series reveals that only strongest interactions are consistent across replicates, while the analysis of the Maizuru community shows the strong negative impact of the chameleon goby on most species of the community, and a potential indirect negative effect of temperature by favouring goby population growth.
- 4. Overall, NODEs alleviate the need for a mechanistic understanding of interactions, and BNGM alleviates the heavy computational cost. This is a crucial step availing quick NODE fitting, cross-validation, and uncertainty quantification, as well as more objective estimation of interactions, and complex context dependence, than parametric models.

1 Introduction

The concept of population is central in ecology (Berryman 2002). Ecologists have had a longstanding interest in finding laws that govern population dynamics, namely changes in the number of individuals in the populations (Lawton 1999; Turchin 1999). Population dynamics can be characterised by a logistic growth, or similar forms, limited by ecological interactions with other organisms, and by the state of the environment (Turchin 2001; Berryman 2003). Intra-specific interactions correspond to interactions between individuals of different sex, age or size classes, belonging to the same species (Turchin 2001). Inter-specific interactions are interactions between individuals from different species, be it competitors, preys, predators, or pathogens (Turchin 2001; Berryman 2003). These interactions can cause populations to have lagged effects impacting their own growth, often called feedback effects, mediated by their impact on the other populations they 11 interact with (Berryman and Turchin 1997). 12 Characterising these interactions has been a longtime challenge. Ecologists started analysing time series data with parametric models (Royama 1984; Kendall et al. 1999; Ives et al. 2003; Gross, Ives, and Nordheim 2005), as time series of population counts are the most commonly collected longterm data in biology (Kendall et al. 1999). Initial analysis involved fitting simple auto-regressive 16 linear models to time series of a single species, leading to contentious interpretations of interactions 17 thereby inferred (e.g. Berryman and Turchin 1997). For instance, Royama et al. interpreted higher 18 order lags as evidence of species interactions (Royama 1984), while Lande et al. interpreted them as age-structure signatures (Lande et al. 2002). Coulson et al. showed they can even be caused by

interactions between the sexes (Mysterud, Coulson, and Stenseth 2002). Jonzen et al. added doubt
over interpreting lags by demonstrating that autocorrelation in environmental noise could prevent
altogether the reliable estimation of lag effects in single species time series data (Jonzén et al.
2002). More recent work has investigated time series of multiple species, environmental factors,
and has mechanistically modelled various ecological interactions (e.g. Bruijning, Jongejans, and
Turcotte 2019; Rosenbaum et al. 2019; Adams et al. 2020). In these models, ecological interactions
are quantified explicitly by specific parameters, rather than phenomenologically with lags. This
allowed for a more thorough quantification of interactions and comparison of alternative ecological
interactions architectures.

However, ecologists still face two main obstacles when estimating ecological interactions from time
series data. The first is that interactions are highly context-dependent, so that they change in time
with the state of the ecosystem and of the environment (Song et al. 2020). Ecological interactions
were traditionally considered linear or fixed, yet there is substantial evidence that this is not the
case in nature (e.g. Bonsall, Meijden, and Crawley 2003; Gross, Ives, and Nordheim 2005; Kendall
et al. 2005; Ushio et al. 2018; Bruijning, Jongejans, and Turcotte 2019; Rosenbaum et al. 2019;
Bonnaffé et al. 2021). The effect of the population on itself depends on the density of individuals
(e.g. Lingjaerde et al. 2001; Moe et al. 2005; Brook and Bradshaw 2006); while predation rates can
depend on the density of the predator (Jost and Ellner 2000; Yoshida et al. 2003). Many vital rates
underpinning ecological interactions are age- and size-dependent (Bonnaffé et al. 2018; Bonnaffé
et al. 2021), and governed by environmental variables, such as temperature (Brown et al. 2004).

- Interactions also change following evolution of the traits that underpin them (Turchin et al. 2003;
- 42 Yoshida et al. 2003). This makes it virtually impossible to model the full complexity of ecological
- interactions (Lawton 1999; Kendall et al. 1999).
- 44 This leads to the second obstacle, known as structural sensitivity, namely sensitivity of the results
- to the structure of the model (Wood 2001; Adamson and Morozov 2013). Because of the com-
- 46 plexity of the interactions, we often lack suitable mathematical representations to portray them
- 47 (Jost and Ellner 2000; Wood 2001; Ellner, Seifu, and Smith 2002; Wu, Fukuhara, and Takeda
- ⁴⁸ 2005). Parametric representations of the interactions are assumed *a priori*, which means that any
- 49 interaction quantified is ultimately contingent on this arbitrary choice, and hence potentially bi-
- ₅₀ ased (Jost and Ellner 2000; Wood 2001; Ellner, Seifu, and Smith 2002; Wu, Fukuhara, and Takeda
- 51 2005). Parametric inference of ecological interactions from time series data therefore only provides
- qualitative evidence, requiring further experimental verification and quantification (Kendall et al.
- 53 1999).
- Nonparametric modelling provides a powerful alternative that can help solve these problems (e.g.
- Jost and Ellner 2000; Wood 2001; Ellner, Seifu, and Smith 2002; Wu, Fukuhara, and Takeda 2005;
- 56 Pasquali and Soresina 2018). Nonparametric forms give more freedom to researchers wishing
- 57 to model population dynamics, and allow a test of whether the linear or linearised assumption of
- standard models is warranted. Interactions are quantified as the sensitivity of the nonparametric ap-
- proximation of the dynamics with respect to other state variables (Sugihara et al. 2012; Ushio et al.
- 60 2018). Nonparametric models require minimal assumptions regarding the mathematical nature of

ecological interactions (Jost and Ellner 2000; Gross, Ives, and Nordheim 2005), and hence provide interaction estimates that are more robust to model structure (Wood 2001). In particular, artificial neural networks (ANNs) offer a promising, yet underused, nonparametric alternative to linear functional forms. In previous work, we introduced a powerful framework, relying on neural ordinary differential equations (NODEs, Chen et al. 2019) to approximate the dynamics of populations nonparametrically, from which we derive ecological interactions (Bonnaffé, Sheldon, and Coulson 2021). More specifically, the ANNs embedded in the ODEs learn nonparametrically the shape of the per capita growth rate of the populations and its dependence on the state variables of the system (Bonnaffé, Sheldon, and Coulson 2021). Combined with the Geber method (Hairston et al. 2005), we are able to estimate the direction, strength, and degree of nonlinearity of interactions. One limitation of the approach lies in the computational cost of fitting the NODEs (Chen et al. 2019; Bonnaffé, Sheldon, and Coulson 2021). This is due to the fact that NODEs, as with ODEs, need to be simulated over the entire range of the time series in order to compute the likelihood of the trajectories of the model. This can be avoided by using gradient matching, which requires interpolating the time series, and fitting the ODEs directly to the interpolated dynamics (Jost and Ellner 2000; Aarts and Veer 2001; Ellner, Seifu, and Smith 2002). Although a similar approach has been proposed (see Treven et al. 2021), there are no implementations of it to fitting NODEs, in spite of its great potential for cutting down computational costs. In addition, given the novelty of the framework, the accuracy and robustness of NODEs in estimating ecological interactions

remain largely unexplored. Most of the work to date is concerned with the accuracy of the fitted

trajectories and of the forecasts (Mai, Shattuck, and O'Hern 2016; Treven et al. 2021; Frank 2022),
while little attention has been given to the functional form of the processes that are producing
the dynamics approximated by NODEs (but see Hu et al. 2020 for a step in this direction). It
is important to understand to what extent the neural networks embedded within NODEs carry
meaningful biological information (Novak and Stouffer 2021).

In this manuscript, we first introduce a novel fitting technique for NODEs, Bayesian neural gradient matching (BNGM). The method extends gradient matching by using neural networks to interpolate the time series data instead of splines (Ellner, Seifu, and Smith 2002), and Bayesian regularisation to fit NODEs to the interpolated dynamics (Cawley and Talbot 2007). This cuts down the fitting time of NODEs to only a few seconds, compared to about 30 minutes in our previous work (Bonnaffé, Sheldon, and Coulson 2021), allowing for efficient cross-validation, and uncertainty quantification. We then demonstrate that NODEs are highly accurate in recovering ecological interactions in an artificial three-species prey-predator system where truth is known. Finally, we conclude the work by characterising ecological interactions in three replicates of an experimental three-species prey-predator system with an algae, flagellate, and rotifer (Hiltunen et al. 2013), as well as in the classic hare and lynx time series (Odum and Barrett 1972). We find that only main interactions, between the algae and the rotifer, are conserved across the three replicates, and not the interactions of the flagellate with the other species. We also find that in most cases linear interactions are sufficient to explain the dynamics apart from nonlinearity in the effect of the prey on the top predator in both the rotifer and lynx.

2 Material and Methods

2 2.1 Method overview

We provide a nonparametric method for estimating ecological interactions from time series data of species density. We do this by approximating the dynamics of each species with neural ordinary differential equations (NODEs, Bonnaffé, Sheldon, and Coulson 2021). We then compute ecological interactions as the sensitivity of these dynamics to a change in the respective species densities (Sugihara et al. 2012; Bonnaffé, Sheldon, and Coulson 2021). We provide a novel method, Bayesian neural gradient matching (BNGM), allowing us to fit NODE systems in a only a few seconds.

10 2.2 Neural ordinary differential equation

A NODE is a class of ordinary differential equation (ODE) that is partly or entirely defined as an ar-111 tificial neural network (ANN) (Chen et al. 2019). They are useful to infer dynamical processes non-112 parametrically from time series data (Bonnaffé, Sheldon, and Coulson 2021). We choose NODEs 113 over standard statistical approaches because they offer two advantages. The first is that NODEs 114 approximate the dynamics of populations nonparametrically. NODEs are therefore not subjected 115 to incorrect model specifications (Jost and Ellner 2000; Adamson and Morozov 2013). This provides a more objective estimation of the inter-dependences between state variables. The second 117 advantage is that it is a dynamical systems approach. So that the approach includes lag effects 118 through interacting state variables, not only direct effects between them.

We first consider a general NODE system,

$$\frac{dy_i}{dt} = f_p(y, \theta_i), \tag{1}$$

where dy_i/dt denotes the temporal change in the i^{th} variable of the system, y_i , as a function of the other state variables $y = \{y_1, y_2, ..., y_I\}$. The function f_p is a nonparametric function of the state variables and its shape is controlled by the parameter vector θ_i . In the context of NODEs, f_p is an ANN. The most common class of ANN used in NODEs are single-layer fully connected feedforward ANNs (e.g. Wu, Fukuhara, and Takeda 2005), also referred to by single layer perceptrons (SLPs, e.g. Bonnaffé, Sheldon, and Coulson 2021),

$$f_p(y, \theta_i) = f_{\lambda} \left(\theta_i^{(0)} + \sum_{j=1}^{J} \theta_{ij}^{(1)} f_{\sigma} \left(\theta_{ij}^{(2)} + \sum_{k=1}^{I} \theta_{ijk}^{(3)} y_k \right) \right), \tag{2}$$

which feature a single layer, containing J neurons, that maps the inputs, here the state variables y, to a single output, the dynamics of state variable i, dy_i/dt . The parameter vector θ_i contains the weights $\theta^{(l)}$ of the connections in the SLPs. SLPs can be viewed as weighted sums of activation functions f_{σ} , which are usually chosen to be sigmoid functions $f(x) = 1/(1 + \exp(-x))$. The link function f_{λ} allows to map the output of the network to a specific domain, for instance applying tanh will constrain the dynamics between -1 and 1, $dy_i/dt \in]-1,1[$. Multi-layer networks can also be used but are generally considered unnecessary since pioneering work established that a single layer is sufficient to approximate any continuous function to a desired level of error (Funahashi and Nakamura 1993).

This general form can be changed to represent biological constraints on the state variables. In particular for population dynamics, the state variables are strictly positive population densities, $y_i = N_i \in \mathcal{R}^+$. We could hence re-write equation (1) as, $dN_i/dt = f_p(N, \theta_i)N_i$, where the SLPs approximate the per-capita growth rate of the populations. More details regarding these models can be found in our previous work (Bonnaffé, Sheldon, and Coulson 2021).

2.3 Fitting NODEs by Bayesian neural gradient matching

In this section, we describe how to estimate the parameters θ of the NODE system given a set of time series. Fitting NODEs can be highly computationally intensive, which hinders uncertainty 143 quantification, cross-validation, and model selection (Bonnaffé, Sheldon, and Coulson 2021). We 144 solve this issue by introducing BNGM, a computationally efficient approach to fit NODEs. The ap-145 proach involves two steps (Fig. 1). First, we interpolate the state variables and their dynamics with 146 neural networks (Fig. 1, red boxes). Second, we train each NODE to satisfy the interpolated state 147 and dynamics (Fig. 1, blue boxes). This bypasses the costly numerical integration of the NODE 148 system and provides a fully mathematically tractable expression for the posterior distribution of the 149 parameter vector θ . We coin the term BNGM to emphasise two important refinements of the stan-150 dard gradient matching algorithm (Ellner, Seifu, and Smith 2002). The first is that we use neural 151 networks as interpolation functions, and the second is that we use Bayesian regularisation to limit 152 overfitting and estimate uncertainty around parameters (Cawley and Talbot 2007). 153

154 Interpolating the time series

The first step is to interpolate the time series and differentiate it with respect to time in order to approximate the state and dynamics of the variables. We perform the interpolation via nonparametric regression of the interpolating functions on the time series data,

$$Y_{it} = \tilde{y}_i(t, \boldsymbol{\omega}_i) + \boldsymbol{\varepsilon}_{it}^{(o)}, \tag{3}$$

where Y_{it} is observed value of the state variable i at time t, $\tilde{y}_i(t, \omega_i)$ is the value predicted by the interpolation function given the parameter vector ω_i , and $\varepsilon_{it}^{(o)}$ is the observation error between the observation and prediction. The interpolation function is chosen to be a neural network,

$$\tilde{y}_i(t, \boldsymbol{\omega}_i) = f_{\lambda} \left(\boldsymbol{\omega}_i^{(0)} + \sum_{j=1}^J \boldsymbol{\omega}_{ij}^{(1)} f_{\sigma} \left(\boldsymbol{\omega}_{ij}^{(2)} + \boldsymbol{\omega}_{ij}^{(3)} t \right) \right), \tag{4}$$

where the parameter vector ω_i contains the weights $\omega^{(l)}$ of the network. We can further differentiate this expression with respect to time to obtain an interpolation of the dynamics of the state variables (Fig. 1, red boxes),

$$\frac{\partial \tilde{y}_i}{\partial t}(t, \boldsymbol{\omega}_i) = \sum_{j=1}^J \boldsymbol{\omega}_{ij}^{(1)} \boldsymbol{\omega}_{ij}^{(3)} \frac{\partial f_{\boldsymbol{\sigma}}}{\partial t} \left(\boldsymbol{\omega}_{ij}^{(2)} + \boldsymbol{\omega}_{ij}^{(3)} t \right) \frac{\partial f_{\boldsymbol{\lambda}}}{\partial t} \left(\boldsymbol{\omega}_i^{(0)} + \sum_{k=1}^J \boldsymbol{\omega}_{ik}^{(1)} f_{\boldsymbol{\sigma}} \left(\boldsymbol{\omega}_{ik}^{(2)} + \boldsymbol{\omega}_{ik}^{(3)} t \right) \right). \tag{5}$$

Fitting NODEs to the interpolated time series

The second step is to train the NODE system (Eq. 1) to satisfy the interpolated dynamics. Thanks to the interpolation step, this simply amounts to performing a nonparametric regression of each NODE (Eq. 1) on the interpolated dynamics (Eq. 5),

$$\frac{\partial \tilde{y}_i}{\partial t}(t, \omega_i) = \frac{dy_i}{dt} (\tilde{y}, \theta_i) + \varepsilon_{it}^{(p)}, \tag{6}$$

where $\varepsilon_{it}^{(p)}$ is the process error, namely the difference between the interpolated dynamics, $\partial \tilde{y}_i/\partial t$ and the NODE, dy_i/dt , given the interpolated state variables $\tilde{y} = \{\tilde{y}_1, \tilde{y}_2, ..., \tilde{y}_I\}$ (Fig. 1, blue boxes).

171 Bayesian regularisation

In the context of standard gradient matching, defining the observation model (Eq. 3) and process model (Eq. 6) would be sufficient to fit the NODE system (Eq. 1) to the time series via optimisation 173 (Jost and Ellner 2000; Ellner, Seifu, and Smith 2002; Wu, Fukuhara, and Takeda 2005). We could 174 find the parameter vector ω_i and θ_i that minimise the sum of squared observation and process errors, 175 $\varepsilon_{it}^{(o)}$ and $\varepsilon_{it}^{(p)}$ (Eq. 3 and 6). However, this approach is prone to overfitting, and does not provide estimates of uncertainty around model predictions. To account for this, we introduce Bayesian 177 regularisation, which allows us to control for overfitting by constraining parameters with prior 178 distributions (Cawley and Talbot 2007), and to root our interpretation of uncertainty in a Bayesian 179 framework. 180

First, we define a simple Bayesian model to fit the interpolation functions (Eq. 3) to the time series data. We assume normal distributions for the observation error, $\varepsilon_{ij}^{(o)} \sim \mathcal{N}(0, \sigma_i)$, and for the parameters, $\omega_{ij} \sim \mathcal{N}(0, \gamma_{ij})$. Here, we are only interested in interpolating the time series accurately,

irrespective of the value of σ_i and γ_{ij} . Therefore, we use the approach developed by Cawley and Talbot to average out the value of the parameters σ_i and γ_{ij} in the full posterior distribution (Cawley and Talbot 2007), assuming gamma hyperpriors $p(\xi) \propto \frac{1}{\xi} \exp\{-\xi/2\}$ for both parameters. This yields the following expression for the log marginal posterior density of the parameters,

$$\log P(\omega_i \mid Y_i) \propto -\frac{N^{(o)}}{2} \log \left(1 + \sum_{t=1}^{N^{(o)}} \left(\varepsilon_{it}^{(o)} \right)^2 \right) - \frac{M^{(o)}}{2} \log \left(1 + \sum_{j=1}^{M^{(o)}} \omega_{ij}^2 \right)$$
(7)

where P is the marginal posterior density, $\omega_i = \{\omega_{i1}, \omega_{i2}, ..., \omega_{iM^{(o)}}\}$ is the observation parameter vector controlling the interpolation function, $Y_i = \{Y_{i1}, Y_{i2}, ..., Y_{iN^{(o)}}\}$ corresponds to the sequence of observations of state variable i at time step t, $N^{(o)}$ is the total number of time steps in the time series, $\varepsilon_{it}^{(o)}$ is the observation error at time step t between the interpolated and observed value of variable i, $M^{(o)}$ is the total number of parameters. More details on how to derive this expression can be found in a supplementary file (Supplementary A).

Then, we define a simple Bayesian model to fit the NODEs to the interpolated dynamics, given the interpolated states. We assume normal distributions for the observation error, $\varepsilon_{it}^{(p)} \sim \mathcal{N}(0, \sigma_i)$, and parameters, $\theta_{ij} \sim \mathcal{N}(0, \delta_{ij})$. This gives the following expression for the log posterior density of the parameters given the interpolations,

$$\log p(\theta_i \mid \boldsymbol{\omega}) \propto -\frac{1}{2} \sum_{t=1}^{N^{(p)}} \left(\frac{\boldsymbol{\varepsilon}_{it}^{(p)}}{\sigma_i}\right)^2 - \frac{1}{2} \sum_{j=1}^{M^{(p)}} \left(\frac{\theta_{ij}}{\delta_{ij}}\right)^2$$
(8)

where $\theta_i = \{\theta_{i1}, \theta_{i2}, ..., \theta_{iM^{(p)}}\}$ are the NODE parameters of the i^{th} variable, $\omega = \{\omega_1, \omega_2, ..., \omega_l\}$

are the interpolation parameters of each state variable, $\varepsilon_{it}^{(p)}$ is the process error of variable i at time 199 step t between the interpolated dynamics and NODE prediction, σ_i is the standard deviation of the likelihood, $M^{(p)}$ is the total number of parameters, δ_{ij} is the standard deviation of the prior 201 distribution of parameter θ_{ij} . 202 This approach allows us to limit overfitting by adjusting the constraint on the parameters, which 203 is controlled by the standard deviation of the parameter prior distributions, δ_{ij} (Cawley and Talbot 204 2007; Bonnaffé, Sheldon, and Coulson 2021). We could set small values of δ to limit the degree 205 of nonlinearity in the response, or to eliminate specific variables from the model by constraining 206 their parameters to be close to zero. We identify the appropriate degree of constraint δ_i on NODE 207 parameters via cross-validation. We split the interpolated data into a train, validation, and test set, 208 for instance, in three thirds. We train the NODE model on the train set and predict the validation 209 set. We repeat this process for increasing values of δ_i , until we find the value that maximises the 210 log likelihood of the validation data. We can perform multiple folds of validation by swapping the 211 train and validation set, or by varying the size of the train/validation split. 212

2.4 Inference and uncertainty quantification

Finally, we estimate uncertainty in parameter values by anchored ensembling, which produces approximate Bayesian estimates of the posterior distribution of the parameters (Pearce et al. 2018).

This involves sampling a parameter vector from the prior distributions, $\theta_i \sim \mathcal{N}(0, \delta_i)$, and then optimising the posterior distribution from this starting point, $\theta_i^* = argmax \log p(\theta_i \mid \omega)$. By re-

peatedly taking samples, the sampled distribution θ^* approaches the posterior distribution and provides estimates and error around the quantities that can be derived from the models. The expectation and uncertainty around derived quantities can then be obtained by computing the mean and variance of the approximated posterior distributions. The great strength of this approach is that it is unlikely to get stuck in local maxima hence providing a more thorough exploration of the parameter space.

2.4 2.5 Analysing NODEs

In this study we are mainly interested in two outcomes of NODEs, namely inferring the direction (or effect) and strength (or contribution) of interactions between the state variables (Bonnaffé, Sheldon, and Coulson 2021). We define the direction of the interaction between variable y_i and y_j as the derivative of the dynamics of y_i with respect to y_j , and vice versa (Sugihara et al. 2012),

$$e_{ijt} = \frac{\partial}{\partial y_j} \frac{dy_i}{dt}.$$
 (9)

Knowing the direction, however, is not sufficient to determine the importance of a variable for the dynamics of another. Given the same effects, a variable that fluctuates a lot will have a greater impact on the dynamics of a focal variable, compared to a variable that remains quasi-constant. For example, a predator can have a negative effect on the prey population, but its actual impact/contribution to the dynamics of the prey population depends on its own dynamics, that is if the predator population decreases, it has a positive contribution to the change in growth rate of

the prey population. We hence compute the strength of the interaction by multiplying the dynamics of a variable y_j by its effect on the focal variable y_i , also known as the Geber method (Eq. 3 in Hairston et al. 2005),

$$c_{ijt} = \frac{dy_j}{dt} \frac{\partial}{\partial y_i} \frac{dy_i}{dt}.$$
 (10)

To summarise results across the entire time series we can compute the mean effects e_{ij} by averaging e_{ijt} across all time steps, $e_{ij} = 1/N^{(p)} \sum_t e_{ijt}$, as well as the relative total contribution, c_{ij} , of a variable to the dynamics of another by computing the relative sum of square contributions, $c_{ij} = \left(\sum_{ijt} c_{ijt}^2\right)^{-1} \sum_t c_{ijt}^2$. By computing the direction and strength of interactions between all the variables in the system we can build dynamically informed ecological interaction networks (e.g. fig. 5). Other metrics can be computed by analysing the NODEs, such as equilibrium states, these are discussed in our previous work (Bonnaffé, Sheldon, and Coulson 2021).

5 3 Case studies

246 3.1 Case study 1: artificial tri-trophic prey-predator oscillations

In this first case study, we aim to demonstrate the accuracy of the NODE fitted by BNGM in inferring nonlinear per-capita growth rates in a system where truth is known. Hence, we simulate a set of time series from a tri-trophic ODE model with known equations and parameters, and we compare the fitted NODEs to the actual ODEs.

251 System

We consider a tri-trophic ODE system consisting of a prey, an intermediate predator, and a top
predator. The system is built on the real tri-trophic system featuring algae, flagellates, and rotifers,
considered in case study 2 (Hiltunen et al. 2013),

$$\frac{dG}{dt} = \left(\alpha \left(1 - \frac{G}{\kappa}\right) - \frac{\beta B}{1 + \delta G} - \frac{\gamma R}{1 + \delta G}\right) G$$

$$\frac{dB}{dt} = \left(\frac{\beta G}{1 + \delta G} - \phi R - \mu\right) B$$

$$\frac{dR}{dt} = \left(\frac{\gamma G}{1 + \delta G} + \phi B - \nu\right) R,$$
(11)

respectively, α is the prey intrinsic growth rate, limited by a carrying capacity κ , β and γ are the predation rates by the intermediate and top predator, δ is the saturation rate of prey predation, which 257 emulates the capacity of the algae to display predator defense at higher algal density (Hiltunen et 258 al. 2013), ϕ is the predation rate of the intermediate predator by the top predator, μ and ν are the 259 intrinsic mortality of the intermediate and top predator. We simulate a case of invasion, by introducing the top predator at a low density, with a set of 261 parameters that result in dampening prey-predator oscillations, namely $\alpha = 1$, $\beta = 2.5$, $\gamma = 1.5$, 262 $\kappa = 3$, $\delta = \phi = \mu = \nu = 1$. We focus on the middle section of the time series, $t \in [20, 50]$, as in 263 the initial section the top predator is rare, and in the later section populations have attained a fixed 264 equilibrium point. The resulting time series are presented in figure 2.

where G, B, and R, correspond to the prey, intermediate, and top predator population densities,

NODE model

In order to nonparametrically learn the per-capita growth rate of each species, and to derive ecological interactions, we define a three-species NODE system,

$$\frac{dR}{dt} = r_R(R, G, B, \beta_R)R$$

$$\frac{dG}{dt} = r_G(R, G, B, \beta_G)G$$

$$\frac{dB}{dt} = r_B(R, G, B, \beta_B)B,$$
(12)

where the per-capita growth rates r_R , r_G , and r_B are neural network functions of the density R, G, B of each species (function f_p , Eq. 2). We choose a combination of linear and exponential activation functions $f_{\sigma,j \leq J/2}(x) = x$, and $f_{\sigma,j > J/2}(x) = \exp(x)$. This allows us to progressively switch from a simple linear model to a nonlinear model by releasing the constraint on the parameters of the network during cross-validation. The number of units in the hidden layer J is chosen to be 10, as this is a commonly used number for systems of that size (e.g. Wu, Fukuhara, and Takeda 2005; Bonnaffé, Sheldon, and Coulson 2021).

276 Time series interpolation

We interpolate the time series using the neural network described in section 2.3 (Eq. 4). We set the number of neurons in the network to J=30. We use sinusoid activation functions, $f_{\sigma}(x)=$ sin(x), so that the weights $\omega_{ij}^{(1)}$, $\omega_{ij}^{(2)}$, and $\omega_{ij}^{(3)}$ control the amplitude, shift, and frequency of the oscillations in the time series, respectively. Given that the population densities are strictly positive R, G, $B \in \mathcal{R}^+$, we use an exponential link function, $f_{\lambda}(x) = \exp(x)$. We then approximate the marginal posterior distribution of the interpolation parameters, and thereby of interpolated states and dynamics, by taking 100 samples from the log marginal posterior distribution (Eq. 7) via anchored ensembling. In practice, the high number of parameters in the neural network equation may impede the fit of the time series, especially for short time series. We found that dividing the number of parameters $M^{(o)}$ (Eq. 7) by the number of neurons in the network J (Eq. 2) yields consistent fitting results. Interpolated states and dynamics are presented in figure 2.

Fitting NODEs to the interpolated time series

We fit the NODE system to the interpolated time series. In practice, we fit the NODE to the expec-289 tation of the interpolated state and dynamics, $E(\tilde{y}_i)$ and $E(d\tilde{y}_i/dt)$, by averaging over all sampled 290 interpolation parameters. An alternative approach could be to consider the interpolation that max-291 imises the log marginal posterior density, but this may decrease repeatability due to the difficulty of 292 reliably identifying a global maximum. Averaging across multiple interpolations ensures an overall 293 smoother and robust interpolation. In addition, we standardise the response and explanatory vari-294 ables with respect the their mean and standard deviation (i.e. $Z = (Y - \mu)/\sigma$). This is to facilitate the training of the NODE by equalizing the scale of the different parameters in the neural network. 296 Then, we identify the optimal regularisation parameter δ (Eq. 8) by cross-validation. To do that, 297 we split the data in three thirds, train NODEs on the first third, and calculate the log likelihood of 298 the validation set for increasing values of δ , from 0.01 (linear) to 0.3 (highly nonlinear), by increments of 0.025. This allows us to identify the maximum degree of nonlinearity, δ , in the per-capita 300 growth rate that ensures generalisability throughout the time series. Then, we approximate the

posterior distribution of the NODE parameters by taking 30 samples from the posterior distribution (Eq. 8). We ensure moderate temporal autocorrelation and normality by visualising the residuals of the models. We also ensure results repeatability by running the entire fitting process a second time.

Computing ecological interactions

Finally, we analyse the shape of the per-capita growth rates to recover the interaction between the 307 three species in the system. In particular, we look at the effect and contribution of each species to the dynamics of the others. The effect is computed as the sensitivity (i.e. the gradient) of the 309 per-capita growth rate of a given species with respect to the density of the other species (Sugihara 310 et al. 2012; Bonnaffé, Sheldon, and Coulson 2021). The contribution is computed following the 311 Geber method (Hairston et al. 2005), which consists in multiplying the dynamics of a variable by 312 its effects on the other variables. We further compute the importance of a species in driving the 313 dynamics of another by computing its relative total contribution compared to other species. More 314 details on how to compute these quantities can be found in section 2.5 and in our previous study 315 (Bonnaffé, Sheldon, and Coulson 2021).

7 Benchmark

In order to demonstrate the suitability of BNGM for fitting NODEs and inferring ecological interactions we compare our approach to three existing methods. For this purpose, we focus on the artificial time series, as this offers the possibility for comparing predictions to the truth, known from the equations that generated the time series.

We first consider a standard NODE model (Bonnaffé, Sheldon, and Coulson 2021), as our BNGM approach seeks to alleviate the computational cost of fitting NODEs. We define the per capita 323 growth rate as an ANN with a single layer, 3 inputs, 10 hidden nodes, and exponential activation functions. We use a Bayesian model, assuming log normal distributions for species den-325 sity $Y_i \sim \log \mathcal{N}(y_i, \theta_i)$, and uniform uninformative prior distributions for the network parameters 326 $\theta_i \sim \mathcal{U}(-10, 10)$, initial densities $y_{i0} \sim \mathcal{U}(0, 10)$, and variance $\sigma_i \sim \log \mathcal{N}(0.5, 0.5)$. Our imple-327 mentation of standard NODEs differs from our BNGM approach in three ways. First, the standard 328 NODE ANN has 3 outputs instead of one, as variables are fitted jointly. Second, computing the 329 posterior density of the parameters requires to solve the NODE system with a numerical ODE 330 solver (Runge-Kutta, package deSolve). Third, we do not constrain the parameters of the network 331 given that the prohibitive fitting times prevent the tuning of the regularisation parameters.

We also consider a parametric ODE model, as this is the closest parametric alternative to NODEs to infer ecological interactions. This model only differs from the standard NODE model in that the per-capita growth rate is approximated by second order polynomial functions $r_i(y, \theta_i) = \theta_i^{(0)} + \sum_j \theta_{ij}^{(1)} y_j + \sum_j \sum_k \theta_{ijk}^{(2)} y_{ij} y_{ik}$, instead of an ANN, which can handle simple nonlinearities.

To ensure the most meaningful comparison, we implemented the NODEBNGM, standard NODE, and parametric ODE models in base R, using BFGS for optimisation (function *optim*, R v4.2.0). We also followed a similar fitting procedure by independently training 30 models on the train/validation set (i.e. 2/3 of the time series) and predicting the test set (remaining third).

Finally, we implement convergent cross-mapping (CCM). This technique performs locally linear

approximations of the state space of the system to estimate the sensitivity of the dynamics of a variable to a change in other variables (Sugihara et al. 2012). For this we use the package rEDM 343 (v1.13.1, Sugihara et al. 2012), and adapt the example code provided for the three species system. We train the CCM model on the train set and predict outcomes on the test set. We then retrieve s-map coefficients (i.e. the interactions) and approximate the population dynamics and per-capita 346 growth using finite differences, given that the standard implementation of CCM does not provide 347 these estimates by default. 348 For all four methods, we compute the runtime as the average time required to train a single model. Using the best preforming model on the train set, we then predict the population dynamics, growth 350 rate, and ecological effects for the entire time series, including the test set. We compute the accu-351 racy of the predictions by computing the mean sum of squared error (MSE) of predictions versus 352

the truth for both the per capita growth rate and ecological effects, on the train and test set. We also

build the corresponding dynamical interaction networks, using the inferred mean effects and total

contributions, and compare them to the true network of interactions. Results are shown in figure 4

57 3.2 Case study 2: real tri-trophic prey-predator oscillations

and presented in detail in supplementary figures (Fig. S1-5).

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In this second case study, we want to assess the quality of the NODE analysis when performed on a real time series. We are further interested in comparing the direction and strength of uncovered ecological interactions across virtually identical replicated time series.

51 System

We consider a three-species laboratory microcosm consisting of an algal prey (Chlorella autroph-362 ica), a flagellate intermediate predator (Oxyrrhis marina), and a rotifer top predator (Brachionus 363 plicatilis). The algal prey is consumed by the intermediate and top predator, which also consumes 364 the intermediate predator (Arndt 1993). The dynamics of this system, here the daily change in the density of each species, were recorded in three replicated time series experiments performed 366 by Hiltunen and colleagues (Hiltunen et al. 2013). We use their time series because they describe 367 a simple yet biologically realistic ecosystem, and because the quality of the replication of their 368 microcosm reduces as much as possible observational and experimental error, and rules out envi-369 ronmental variation (Hiltunen et al. 2013). We digitised these time series by extracting by hand 370 the coordinates of every points in the referential of the axis of the graph of the original study, and 371 analysed them. 372

373 NODE analysis

We apply the same analysis as performed on the artificial tri-trophic prey-predator oscillations.

This allows us to recover a nonparametric approximation of the growth rate of each species, and then derive the direction and strength of the ecological interactions that underpin their dynamics.

We present detailed results of the analysis of the second time series (Fig. 5), and a summary comparison of the three time series (Fig. 6). Complementary results, including cross-validation plots, and detailed results for the other two replicates can be found in the supplementary material (Supplementary C-D).

3.3 Case study 3: real di-trophic prey-predator oscillations

We infer ecological interactions by NODE BNGM in the hare-lynx system (Odum and Barrett

³⁸³ 1972). This is to provide an example of a longer time series, and to offer a point of comparison

with previous and future implementations of NODEs, which commonly use this time series (e.g.

Bonnaffé, Sheldon, and Coulson 2021; Frank 2022).

386 System

The system is described in details in our previous work (Bonnaffé, Sheldon, and Coulson 2021).

The data consist in a 90-year long time series of counts of hare and lynx pelts collected by trappers

in the Hudson bay area in Canada (Odum and Barrett 1972). The time series displays characteristic

390 10-year long prey-predator oscillations.

NODE analysis

392 We apply the same analysis as previously described, to the exception that the NODE system only

features two variables, H and L, instead of 3. Results are presented in figure 7.

4 3.4 Case study 4: fish community from the Maizuru Bay

Finally, we demonstrate the capacity of NODEs to analyse the drivers of the dynamics of a larger

community by analysing the time series of the Maizuru bay community (Ushio et al. 2018).

397 System

The dataset for this system consists of 12-year long time series of fortnight abundance estimates of

three 200m long and 2m wide transects by underwater visual census conducted along the coast of the Maizuru fishery research station of Kyoto University from 2002 to 2014 (for more details see Ushio et al. 2018). Bottom sea temperature (at 10m depth) was also recorded on each census. The dataset contains 14 dominant species of fish and 1 genus of jellyfish. Only species with more than 1000 sightings were included in the final dataset.

We focussed our analysis on the species with the least sparse records. We discarded the follow-405 ing species from our analysis Engraulis japonicus, Plotosus lineatus, Chaenogobius gulosus, and 406 Siganus fuscescens. We also excluded periods which presented jellyfish blooms, as these were 407 isolated events which could cause numerical errors in the estimation of the dynamics of species 408 abundance. In total, we considered a time period of a hundred time steps from June 2004 to 409 August 2008, and 11 species out of 15, namely Aurelia sp, Sebastes inermis, Trachurus japoni-410 cus, Girella punctata, Pseudolabrus sieboldi, Halichoeres poecilopterus, Halichoeres tenuispinnis, Pterogobius zonoleucus, Tridentiger trigonocephalus, Sphyraena pinguis, and Rudarius ercodes. 412 We included the sea bottom temperature (in degrees celsius) as an additional environmental vari-413 able.

NODE analysis

We then analysed this dataset following the approach described in the method section. We split
the data into three thirds to create a training, validation, testing set (final third), and we followed
the same procedure as described before to tune the regularisation parameters. The NODE system

consisted of 11 NODEs, where the per-capita growth rate is determined by a single-layer ANN with
12 input nodes, 10 hidden nodes, exponential activation functions, and 1 output node. Due to the
high dimensionality of the effects obtained (11 by 12), we only present mean effects and relative
total contributions, obtained by taking the mean, and the relative mean squares, respectively, of
the effects and contributions across the entire time series (Fig. 8). The time series of effects
and contributions are presented in greater details in the supplementary material (Supplementary
G).

26 4 Results

27 4.1 Case study 1: artificial tri-trophic system

We present the results of fitting NODEs by BNGM to the artificial tri-trophic time series in figure 428 2 and 3. We find that both the interpolation of the state variables and dynamics are highly accurate 429 (Fig. 2), given that they closely match the ground truth, known from the equations of the ODE 430 model that we used to generate the time series (Eq. 11). Similarly, we find that the NODE approx-431 imation of the per-capita growth rate of each species also closely matches the ground truth (Fig. 3, a., d., g.). We find negative nonlinear effects of the two predators on the growth rate of the algae 433 (Fig. 3, b., blue and purple lines). This nonlinear pattern is mirrored by the effect of the algae on 434 the growth rate of the predators (Fig. 3, e. and h., red line). The interaction between the two preda-435 tors is also well-recovered (Fig. 3, e., blue line, and h., purple line), in spite of a slight tendency for overestimating the degree of nonlinearity of effects. The BNGM approach hence accurately recovers the dynamical characteristics of the artificial system.

4.2 Benchmark

Figure 4 shows the performance of fitting NODEs by BNGM, compared to standard NODEs, parametric ODEs, and convergent cross mapping models (CCM). We find that fitting NODEs by BNGM
provides the highest estimation accuracy for growth rates and ecological effects, both on the training and test set, as well as competitive runtimes. Standard NODEs provide similar estimation
accuracies, but take over 15 minutes to train. CCM is the fastest technique, as results are obtained
in under a second, but is relatively less accurate. Parametric ODEs are found to be both slow and
less accurate.

We present a detailed breakdown of the runtime of fitting NODEs by BNGM for each system in table 1. We find that it takes on average 5.35 minutes to fit NODEs by BNGM on the smaller systems with 3 or less species, and about 23 minutes to fit the larger system from the Maizuru bay, which features 12 variables. This includes performing $100 \times I$ and $30 \times I$ full optimisations of the posterior distribution of the interpolation and NODE parameters, respectively. This amounts to about 5.37 second to sample each variable of the NODE system once in the smaller systems, and about 28 seconds for the Maizuru bay community. This is over a 100 fold improvement over standard NODE models, which take on average 20 minutes (Fig. 4).

55 4.3 Case study 2: real tri-trophic prey-predator oscillations

We present an in-depth analysis of the drivers of the dynamics of the algae, flagellate, and rotifer 456 population in replicate B (Fig. 5). We find slightly positive nonlinear intra-specific density de-457 pendence in algal growth (Fig. 5, b., red line), and negative nonlinear inter-specific effects of the two predators (purple and blue line). We find that the growth rate of the flagellate is driven by a 459 positive effect of algal density, a negative effect of predation by the rotifer and intra-specific den-460 sity dependence (Fig. 5, e. and f.). The rotifer population is almost solely driven by a positive 461 nonlinear effect of flagellate density (Fig. 5, h., purple line). Overall, comparing results across the three replicates reveals that the effect of the rotifer population on the flagellate and algae, and the 463 effect of the algae on the rotifer, are the strongest and most consistent interactions (Fig. 6, table 464 2). The interactions of the flagellate with the algae, and its effect on the rotifer population varies 465 substantially across replicates (Fig. 6, table 2). Interestingly, intra-specific density dependence in rotifer and algae is also found to be inconsistent across the three replicates. 467

68 4.4 Case study 3: real di-trophic prey-predator oscillations

We present the analysis of the drivers of the hare-lynx population dynamics in figure 7. Cross-validation provides support for nonlinear effects in the per-capita growth rate of the hare and lynx.

We find that the hare population growth rate is mostly determined by a nonlinear negative effect of the lynx population (Fig. 7, b. and c. blue line), and by weak nonlinear positive density dependence (red line). The lynx growth rate is determined by a positive nonlinear effect of the hare (Fig. 7,

e. and f., red line), and to a lesser extent by negative nonlinear intra-specific density dependence (blue line).

4.5 Case study 4: drivers of the Maizuru bay community dynamics

We show the results of the NODE analysis of the drivers of the dynamics Maizuru bay community in figure 8. Our main finding is that the chameleon goby (*Tridentiger trigonocephalus*) has a strong negative effect on 8 of the 11 dominant species of the community. We find that *E. ercodes* also has a strong negative impact on other species in the community, although relatively smaller than that of the chamelon goby. We find a positive effect of sea bottom temperature on the growth rate of the chameleon goby. Other effects are found to be mostly positive and have a relatively smaller impact on community dynamics.

5 Discussion

Characterising ecological interactions from time series data is challenging. This is due to the fact
that interactions can be highly context-dependent processes (Song and Saavedra 2021), making it
difficult to identify parametric models that encapsulate their complexity (Wood 2001). Interactions estimated with parametric models are contingent on the parameterisation arbitrarily chosen
by the observer, and hence risk being biased (Wood 2001; Adamson and Morozov 2013). We
provide a novel method for estimating ecological interactions nonparametrically, by using neural
ordinary differential equations (NODEs) fitted with Bayesian neural gradient matching (BNGM).

First, we remove the cost of fitting NODEs by introducing BNGM, which allows for NODE fitting in only a few seconds. The method involves interpolating time series and dynamics with neural networks, and then fitting NODEs to interpolated dynamics with Bayesian regularisation. We further demonstrate that this approach is accurate, as NODEs approximate with minimal error the ecological interactions in artificial time series, where real interactions are known, performing better than three existing methods. Finally, we estimate the strength, direction, importance, and nonlinearity of ecological interactions in 3 natural and experimental systems, showing variation in ecological interactions within and across the time series.

Performance of NODEs fitted by BNGM compared to existing methods

Our approach relies on approximating population dynamics with NODEs and then computing their 501 sensitivity to a change in the density of the different populations in the system (Bonnaffé, Shel-502 don, and Coulson 2021). We demonstrate that NODEs accurately recover the dynamics, strength, 503 direction, and nonlinearity of ecological interactions in artificial tri-trophic prey-predator time se-504 ries, where truth is known. In particular, we find that the interactions between the prey and the two predators are nonlinear, and thereby oscillate throughout the time series, which is consistent 506 with the model, that features a resistance to predation at high prey density. We also recover the 507 interactions between the two predators, in spite of a slight tendency to overestimate the degree of 508 nonlinearity. To our knowledge, this is the first assessment of the accuracy of NODEs in recovering interactions between variables from time series data, as most of the work focuses on assessing the 510 accuracy of the fitting and forecasting of time series (e.g. Mai, Shattuck, and O'Hern 2016; Chen

et al. 2019; Treven et al. 2021; Frank 2022).

We find that fitting NODEs by BNGM provides higher estimation accuracies of ecological interactions, and reduces substantially fitting times compared to standard NODEs (Bonnaffé, Sheldon,
and Coulson 2021), and parametric ODEs (Rosenbaum et al. 2019). This difference is attributable
to three factors. First, BNGM alleviates the need for solving numerically the NODE system, which
makes it faster to evaluate the posterior distribution. Second, it allows for the calculation of analytical gradients of the posterior distribution, which greatly improve the speed and efficiency of the
gradient descent optimisation algorithm. Finally, it makes it possible to fit each variables independently on each other, which results in a simpler optimisation problem.

521 CCM remains faster than our approach in recovering estimates of ecological interactions (Sugihara et al. 2012), however its accuracy is lower. A possible explanation for this comes from the fact that CCM computes the sensitivity of the total population growth rate, rather than the per-capita growth rate, which can change estimated effects. Additionally, CCM relies on piecewise linear reconstructions of the state space (Deyle et al. 2015), whereas NODEs computes a global nonlinear approximation of the per-capita growth rate on the entire range covered by the data. We view the former as potentially more sensitive to local noise in the state space (Cenci, Sugihara, and Saavedra 2019), compared to the latter, which uses all evidence available to inform local inference.

Our BNGM approach extends standard gradient matching, by using artificial neural networks
(ANNs) as interpolating functions, and Bayesian regularisation to control the nonlinearity of the
processes (Cawley and Talbot 2007). The use of ANNs as interpolating functions sets it apart

from the initial approach of Ellner et al., who use splines to interpolate the time series before approximating the ODEs (Ellner, Seifu, and Smith 2002). ANNs are more general and flexible than 533 splines, as well as being easier to manipulate given that they are defined continuously on the state space, which is especially useful when handling multiple interactions between variables. Our ap-535 proach is related to that of Wu et al., who use ANNs to approximate both the states and ODEs of 536 prey-predator systems (Wu, Fukuhara, and Takeda 2005), as well as that of Treven and colleagues, 537 who developed the Gaussian process equivalent (Treven et al. 2021). In both approaches, they 538 train the interpolation functions at the same time as the NODEs, in order to constrain the interpola-539 tion of trajectories such that they can be achieved by the NODE system, which thereby introduces 540 dynamical coupling between state variables. One of the downsides of this approach is that misestimating one of the state variables of the model biases the estimation of the states and dynamics of other variables. To avoid this, we fit each interpolation and NODE independently to each time 543 series. In addition, this makes it possible to parallelise the code, resulting in potentially even faster computation. 545

Our approach opens new possibilities for nonparametric inference of ecological interactions from time series data. The lower fitting times makes it possible to tackle larger systems, quick and extensive model comparison, cross-validation, and apply more thorough statistical treatments of the uncertainty of these models, for instance by implementing Markov-chain Monte-Carlo (MCMC) sampling.

Ecological interactions in real prey-predator systems

We further tested NODEs in a real setting, by inferring ecological interactions across three replicated time series of an experimental tri-trophic system of algae, flagellate, and rotifer populations 553 (Hiltunen et al. 2013). Our approach reveals that only stronger interactions, namely the negative effects of the rotifer top predator on the other species, and the positive effect of algae on the rotifer, 555 are conserved across the three replicated time series. We also find evidence for nonlinearity in 556 the dynamics of the rotifer, as the positive effect of the algae on rotifer growth oscillates through-557 out the time series. This is consistent with the biology of the system, as the algae tends to form 558 anti-predation clumps at higher density, which would dampen the positive effect of algal density on 559 rotifer growth at high algal density (Yoshida et al. 2003; Hiltunen et al. 2013). We find it interesting 560 that the weaker interactions with the flagellate predator are not consistent across time series, given 561 the controlled laboratory conditions. This system is known to evolve rapidly, it is hence possible that fast evolution of the different populations from the onset of the time series may have driven 563 the system onto different attractors (Yoshida et al. 2003; Yoshida et al. 2007; Hiltunen et al. 2013). 564 Additionally, stochasticity in population dynamics may have a similar effect (Dallas et al. 2021). 565 Disentangling these two sources of variation would require refining the modelling framework, for instance by explicitly including evolution in the model (e.g. with the Price equation, Ellner, Geber, 567 and Hairston 2011), and by using neural stochastic differential equations (i.e. NSDEs, Rackauckas 568 et al. 2019) fitted with a particle filter. While these would constitute interesting developments, our 569 method is still a useful first step, identifying differences between the time series, and demonstrating 570 a reasonable amount of deterministic consistency in the dynamics, judging by the cross-validation and fits. 572

We also analysed the hare-lynx time series (Odum and Barrett 1972), as it is a common benchmark in the field of time series analysis, and provides a comparison point with our previous implemen-574 tation of NODEs (Bonnaffé, Sheldon, and Coulson 2021). As in our previous study, we found a predatory inter-specific interaction between lynx and hare, and negative intra-specific density dependence in the lynx. Evidence for positive density dependence in the hare was more limited than 577 previously found. We also found stronger evidence for nonlinearity, as intra- and inter-specific ef-578 fects oscillated throughout the time series, as a result of density dependence. This difference with 579 our previous study is due to the fact that our previous implementation of NODEs was based on sim-580 ulating the full NODE system, and hence imposed dynamical coupling between the variables. This 581 dynamical coupling comes at a cost, if one variable is not explained well by the model, it will bias 582 the interactions and dynamics of other variables. Here, the time series of lynx and hare are analysed independently, each state variable is interpolated as closely as desired, its effects on the dynamics 584 of other variables are hence even more robust to model misspecification than before. 585

We applied our approach to analyse the drivers of the dynamics of 11 species in the Maizuru bay

(Ushio et al. 2018). We inferred 11 × 12 ecological interactions, and 11 dependencies on water

temperature. We found that the chameleon goby had a strong negative impact on the other species

of the system, showing a strong competitive potential. This species is viewed as an aggressive

competitor (Ushio et al. 2018), and is considered an invasive species in places where it has been

introduced (Goren, Gayer, and Lazarus 2009). We also find a positive effect of temperature on

the growth rate of the chameleon goby, which suggests that warming could have indirect negative

effects on many species on Maizuru bay by favouring the reproduction of the goby.

Surprisingly, our results differ substantially from those obtained by Ushio et al. in their original analysis of the system with convergent cross mapping (Ushio et al. 2018). This may be due to a multitude of factors. First, we considered a different set of species, as some of the time series that Ushio and colleagues used were too sparse to be suitable for our analysis. Additionally, this difference may be explained by the disparity in the estimation accuracies revealed by our benchmark analysis, relating to fundamental mathematical differences between the two approaches, as discussed previously. If nothing else, our analysis of the Maizuru community dynamics demonstrates the usefulness of our BNGM method for fitting NODEs to a larger, more realistic system.

Overall, our approach provides a novel and powerful way of estimating interactions nonparametrically from time series data. The benefit of using NODEs is that they make no assumptions about the nature of the ecological interactions that drive the dynamics of the species (Chen et al. 2019;
Bonnaffé, Sheldon, and Coulson 2021). Hence, we have a better chance at estimating the actual value of the interactions, knowing that it is not subjected to potential incorrect model specifications (Jost and Ellner 2000; Ellner, Seifu, and Smith 2002; Wu, Fukuhara, and Takeda 2005; Kendall et al. 2005; Adamson and Morozov 2013).

609 Limits and prospects

One of the main difficulty in quantifying ecological interactions is to identify potential context dependences on other state variables (Song and Saavedra 2021), for example, whether predation rates are affected by temperature. Our approach allows for the quantification of context dependence,

which shows as nonlinear fluctuations of interactions throughout the time series. In the present work, we only report nonlinearity as evidence for context dependence in the interactions, but we 614 do not attempt to understand what it is attributable to. For instance, we identify nonlinear density 615 dependence in the effect of the algae on the rotifer, but we do not know whether this is due to a 616 change in the effect with algae density or rotifer density, or both. In order to disentangle these 617 higher order effects we could compute the Hessian of the system, namely the second order deriva-618 tive of the dynamics with respect to the different state variables. Though this procedure is simple 619 mathematically, it would result in 27 second order effects to analyse for the simple 3 species system 620 considered here. This type of analysis would get rapidly out of hand for larger systems. Further 621 work should hence consider how to handle these higher order effects, as a way to unveil context 622 dependence in ecological interactions. One further issue is that some interactions may depend on variables that are not observed. For

One further issue is that some interactions may depend on variables that are not observed. For instance, some population dynamics are strongly determined by their demographic state (Lande et al. 2002; Coulson et al. 2004), which would call for time series of the relevant demographic stages.

In the system considered here, the dynamics of algae in the rotifer system are most likely coupled with that of nitrogen, for which no time series was available (Hiltunen et al. 2013). Our method only accounts for observed variables, so that time series for all important variables are required, though unaccounted variables are captured to some extent by nonlinear fluctuations in interactions.

One interesting prospect would hence be to incorporate unobserved/latent state variables into the NODE system (Dupont, Doucet, and Teh 2019; Zhang et al. 2019; Frank 2022). Careful thought

has to be given here as whether to use an ODE or NODE for the latent states given that they are not constrained by observations.

A further question is whether we could use similar approaches to analyse systems larger than the 635 ones considered on this study. In particular, microbial communities feature thousands of species, 636 and hence potentially millions of interactions, which poses a real problem for inference with dynamical models (Ridenhour et al. 2017). Even the simplest linear ODE model would contain 638 millions of parameters, hence entering deep learning territory. We believe that our success there is 639 more readily limited by the availability of time series sufficiently long to identify this many inter-640 actions, rather than by our models. A possible next step could be to address the capacity of more complex NODE and ODE models to estimate interactions in large artificial communities (e.g. Co-642 enen et al. 2020), which could inform us on the relationship between model complexity and data 643 requirements in terms of time series length and sampling frequency.

We consider NODEs, which are only defined along the time dimension. The framework could
easily be extended to any other dimension by considering partial differential equations instead
(Rackauckas et al. 2019). For instance, in a spatial ecology context we could model the dynamics
of populations along two additional spatial dimensions. In an evolutionary context, we could model
the dynamics of populations in phenotype space, by adding phenotypic traits as an additional dimension. The BNGM method could be instrumental in fitting these models, which are notoriously
expensive to stimulate.

Conclusion

We provide a method, BNGM, which allows for NODE fitting in a matter of seconds. This is a crucial step for efficient model selection and uncertainty quantification in NODEs. We also demonstrate that NODEs allow for faster more accurate estimation of the direction, strength, and nonlinearity of ecological interactions than existing approaches, in a system where truth is known. Finally, we estimate ecological interactions in real prey predator systems, showing that only stronger interactions seem to be consistent across replicated time series, and that a single species can account for a large part of the changes in community dynamics. Our study allows for efficient NODE fitting, and confirms the power of NODEs in identifying dynamical coupling between populations.

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666 Data accessibility

All data and code will be made fully available at https://github.com/WillemBonnaffe/NODEBNGM,
as well as on https://datadryad.org/stash/dataset/doi:xxx.

669 Statement of authorship

Willem Bonnaffé designed the method, performed the analysis, wrote the manuscript; Tim Coulson
 led investigations, provided input for the manuscript, commented on the manuscript.

References

- Aarts, L. P. and P. V. D. Veer (2001). "Neural network method for solving partial differential equa-
- tions". In: Neural Processing Letters 14 (3), pp. 261–271.
- Adams, M. P. et al. (Apr. 2020). "Informing management decisions for ecological networks, using
- dynamic models calibrated to noisy time-series data". In: *Ecology Letters* 23 (4), pp. 607–619.
- Adamson, M. W. and A. Y. Morozov (2013). "When can we trust our model predictions? Un-
- earthing structural sensitivity in biological systems". In: *Proceedings of the Royal Society A:*
- Mathematical, Physical and Engineering Sciences 469 (2149), pp. 1–19.
- Arndt, H. (1993). "Rotifers as predators on components of the microbial web (bacteria, heterotrophic
- flagellates, ciliates) a review". In: *Hydrobiologia* 255-256 (1), pp. 231–246.
- Berryman, A. and P. Turchin (1997). "Detection of density dependence: comment". In: Ecology 78
- 683 (1), pp. 318–320.
- Berryman, A. A. (2002). "Population: a central concept for ecology?" In: Oikos 97 (3), pp. 439-
- 685 442.
- Berryman, A. A. (2003). "On principles, laws and theory in population ecology". In: Oikos 103
- 687 (3), pp. 695–701.
- Bonnaffé, W., S. Legendre, A. Danet, and E. Edeline (2021). "Comparison of size-structured and
- species-level trophic networks reveals antagonistic effects of temperature on vertical trophic
- diversity at the population and species level". In: *Oikos*, pp. 1–14.

- 691 Bonnaffé, W., M. Martin, M. Mugabo, S. Meylan, and J. F. L. Galliard (Dec. 2018). "Ontogenetic
- trajectories of body coloration reveal its function as a multicomponent nonsenescent signal". In:
- 693 Ecology and Evolution 8 (24), pp. 12299–12307.
- Bonnaffé, W., B. C. Sheldon, and T. Coulson (2021). "Neural ordinary differential equations for
- ecological and evolutionary time series analysis". In: *Methods in Ecology and Evolution* 2, pp. 1–
- 696 46.
- Bonsall, M. B., E. V. D. Meijden, and M. J. Crawley (2003). "Contrasting dynamics in the same
- plant-herbivore interaction". In: Proceedings of the National Academy of Sciences of the United
- 699 States of America 100 (25), pp. 14932–14936.
- Brook, B. W. and C. J. A. Bradshaw (2006). "Strength of evidence for density dependence in
- abundance time series of 1198 species". In: *Ecology* 87 (6), pp. 1445–1451.
- Brown, J. H., J. F. Gillooly, A. P. Allen, V. M. Savage, and G. B. West (2004). "Toward a metabolic
- theory of ecology". In: *Ecology* 85 (7), pp. 1771–1789.
- ₇₀₄ Bruijning, M., E. Jongejans, and M. M. Turcotte (2019). "Demographic responses underlying eco-
- evolutionary dynamics as revealed with inverse modelling". In: Journal of Animal Ecology 88
- 706 (5), pp. 768–779.
- ⁷⁰⁷ Cawley, G. C. and N. L. C. Talbot (2007). "Preventing over-fitting during model selection via
- bayesian regularisation of the hyper-parameters". In: Journal of Machine Learning Research 8,
- pp. 841–861.

- Cenci, S., G. Sugihara, and S. Saavedra (May 2019). "Regularized S-map for inference and fore-
- casting with noisy ecological time series". In: Methods in Ecology and Evolution 10 (5), pp. 650–
- 712 660.
- Chen, R. T. Q., Y. Rubanova, J. Bettencourt, and D. Duvenaud (2019). "Neural Ordinary Differen-
- tial Equations". In: *arXiv*, pp. 1–19.
- Coenen, A. R., S. K. Hu, E. Luo, D. Muratore, and J. S. Weitz (Apr. 2020). "A Primer for Micro-
- biome Time-Series Analysis". In: Frontiers in Genetics 11.
- Coulson, T., F. Guinness, J. Pemberton, and T. Clutton-Brock (2004). "The demographic conse-
- quences of releasing a population of red deer from culling". In: *Ecology* 85 (2), pp. 411–422.
- Dallas, T., B. A. Melbourne, G. Legault, and A. Hastings (2021). "Initial abundance and stochas-
- ticity influence competitive outcome in communities". In: Journal of Animal Ecology, pp. 1–
- 721 26.
- Deyle, E. R., R. M. May, S. B. Munch, and G. Sugihara (Jan. 2015). "Tracking and forecasting
- ecosystem interactions in real time". In: Proceedings of the Royal Society B: Biological Sciences
- 724 283, pp. 1–9.
- Dupont, E., A. Doucet, and Y. W. Teh (2019). "Augmented Neural ODEs". In: arXiv, pp. 1–11.
- Ellner, S. P., M. A. Geber, and N. G. J. Hairston (2011). "Does rapid evolution matter? Measuring
- the rate of contemporary evolution and its impacts on ecological dynamics". In: *Ecology Letters*
- 728 14 (6), pp. 603–614.
- Ellner, S. P., Y. Seifu, and R. H. Smith (2002). "Fitting Population Dynamic Models to Time-Series
- Data by Gradient Matching". In: *Ecology* 83 (8), p. 2256.

- Frank, S. A. (2022). "Automatic diierentiation and the optimization of diierential equation models
- in biology". In: *arXiv*, pp. 1–10.
- Funahashi, K. ichi and Y. Nakamura (1993). "Approximation of dynamical systems by continuous
- time recurrent neural networks". In: *Neural Networks* 6 (6), pp. 801–806.
- Goren, M., K. Gayer, and N. Lazarus (2009). "First record of the far East Chameleon goby tri-
- dentiger trigonocephalus (Gill, 1859) in the Mediterranean Sea". In: Aquatic Invasions 4 (2),
- pp. 413–415.
- Gross, K., A. R. Ives, and E. V. Nordheim (2005). "Estimating fluctuating vital rates from time-
- series data: A case study of aphid biocontrol". In: *Ecology* 86 (3), pp. 740–752.
- Hairston, N. G. J., S. P. Ellner, M. A. Geber, T. Yoshida, and J. A. Fox (2005). "Rapid evolution
- and the convergence of ecological and evolutionary time". In: *Ecology Letters* 8 (10), pp. 1114–
- 742 1127.
- Hiltunen, T., L. E. Jones, S. P. Ellner, and N. G. J. Hairston (2013). "Temporal dynamics of a simple
- community with intraguild predation: an experimental test". In: *Ecology* 94 (4), pp. 773–779.
- Hu, P., W. Yang, Y. Zhu, and L. Hong (2020). "Revealing hidden dynamics from time-series data
- by ODENet". In: *arXiv*, pp. 1–17.
- ⁷⁴⁷ Ives, A. R., B Dennis, K. L. Cottingham, and S. R. Carpenter (2003). "Estimating community
- stability and ecological interactions from time-series data". In: *Ecological Monographs* 73 (2),
- 749 pp. 301–330.

- Jonzén, N., P. Lundberg, E. Ranta, and V. Kaitala (Feb. 2002). "The irreducible uncertainty of
- the demography Environment interaction in ecology". In: *Proceedings of the Royal Society B:*
- 752 Biological Sciences 269 (1488), pp. 221–225.
- Jost, C. and S. P. Ellner (2000). "Testing for predator dependence in predator-prey dynamics: A
- non-parametric approach". In: Proceedings of the Royal Society B: Biological Sciences 267
- 755 (1453), pp. 1611–1620.
- Kendall, B. E. et al. (1999). "Why do populations cycle? A synthesis of statistical and mechanistic
- modeling approaches". In: *Ecology* 80 (6), pp. 1789–1805.
- Kendall, B. E. et al. (2005). "Population cycles in the pine looper moth: Dynamical tests of mech-
- anistic hypotheses". In: *Ecological Monographs* 75 (2), pp. 259–276.
- Lande, R, S Engen, B.-E Saether, F Filli, E Matthysen, and H Weimerskirch (2002). "Estimating
- Density Dependence from Population Time Series Using Demographic Theory and Life-History
- Data". In: *American Naturalist* 159, pp. 321–337.
- ₇₆₃ Lawton, J. H. (1999). "Are There General Laws in Ecology?" In: Oikos 84 (2), pp. 177–192.
- Lingjaerde, O. C. et al. (2001). "Exploring the density-dependent structure of blowfly populations
- by nonparametric additive modeling". In: *Ecology* 82 (9), pp. 2645–2658.
- Mai, M., M. D. Shattuck, and C. S. O'Hern (2016). "Reconstruction of Ordinary Differential Equa-
- tions From Time Series Data". In: *arXiv*, pp. 1–15.
- Moe, S. J., A. B. Kristoffersen, R. H. Smith, and N. C. Stenseth (2005). "From patterns to pro-
- cesses and back: Analysing density-dependent responses to an abiotic stressor by statistical and

- mechanistic modelling". In: Proceedings of the Royal Society B: Biological Sciences 272 (1577),
- pp. 2133–2142.
- Mysterud, A., T. Coulson, and N. C. Stenseth (2002). "The role of males in the dynamics of ungu-
- late populations". In: *Journal of Animal Ecology* 71, pp. 907–915.
- Novak, M. and D. B. Stouffer (Nov. 2021). "Geometric Complexity and the Information-Theoretic
- 775 Comparison of Functional-Response Models". In: Frontiers in Ecology and Evolution 9.
- Odum, E. P. and G. W. Barrett (1972). "Fundamentals of Ecology". In: The Journal of Wildlife
- 777 *Management* 36 (4), p. 1372.
- Pasquali, S. and C. Soresina (2018). "Estimation of the mortality rate functions from time series
- field data in a stage-structured demographic model for Lobesia botrana". In: *arXiv*, pp. 1–15.
- Pearce, T., F. Leibfried, A. Brintrup, M. Zaki, and A. Neely (2018). "Uncertainty in Neural Net-
- works: Approximately Bayesian Ensembling". In: *arXiv*, pp. 1–10.
- Rackauckas, C., M. Innes, Y. Ma, J. Bettencourt, L. White, and V. Dixit (Feb. 2019). "DiffEqFlux.jl
- A Julia Library for Neural Differential Equations". In: *arXiv*, pp. 1–17.
- Ridenhour, B. J. et al. (Nov. 2017). "Modeling time-series data from microbial communities". In:
- 785 *ISME Journal* 11 (11), pp. 2526–2537.
- Rosenbaum, B., M. Raatz, G. Weithoff, G. F. Fussmann, and U. Gaedke (2019). "Estimating param-
- eters from multiple time series of population dynamics using bayesian inference". In: Frontiers
- in Ecology and Evolution 6 (234), pp. 1–14.
- Royama, T (1984). "Population Dynamics of the Spruce Budworm Choristoneura Fumiferana". In:
- 790 *Ecological Monographs* 54 (4), pp. 429–462.

- Song, C., S. V. Ahn, R. P. Rohr, and S. Saavedra (May 2020). "Towards a Probabilistic Under-
- 792 standing About the Context-Dependency of Species Interactions". In: Trends in Ecology and
- 793 Evolution 35 (5), pp. 384–396.
- Song, C. and S. Saavedra (July 2021). "Bridging parametric and nonparametric measures of species
- interactions unveils new insights of non-equilibrium dynamics". In: Oikos 130 (7), pp. 1027–
- 796 1034.
- ⁷⁹⁷ Sugihara, G. et al. (2012). "Detecting causality in complex ecosystems". In: *Science* 338 (6106),
- 798 pp. 496–500.
- 799 Treven, L., P. Wenk, F. Dörfler, and A. Krause (2021). "Distributional Gradient Matching for Learn-
- ing Uncertain Neural Dynamics Models". In: *arXiv*, pp. 1–14.
- Turchin, P. (1999). "Population Regulation: A Synthetic View". In: Oikos 84 (1), pp. 153–159.
- (2001). "Does population ecology have general laws?" In: Oikos 94, pp. 17–26.
- Turchin, P. et al. (2003). "Dynamical effects of plant quality and parasitism on population cycles
- of larch budmoth". In: *Ecology* 84 (5), pp. 1207–1214.
- Ushio, M. et al. (Feb. 2018). "Fluctuating interaction network and time-varying stability of a natural
- fish community". In: *Nature* 554 (7692), pp. 360–363.
- Wood, S. N. (2001). "Partially specified ecological models". In: Ecological Monographs 71 (1),
- pp. 1–25.
- 809 Wu, J., M. Fukuhara, and T. Takeda (2005). "Parameter estimation of an ecological system by
- a neural network with residual minimization training". In: Ecological Modelling 189 (3-4),
- pp. 289–304.

- Yoshida, T., S. P. Ellner, L. E. Jones, B. J. M. Bohannan, R. E. Lenski, and N. G. J. Hairston (2007).
- "Cryptic population dynamics: Rapid evolution masks trophic interactions". In: *PLoS Biology* 5
- (9), pp. 1868–1879.
- Yoshida, T., L. E. Jones, S. P. Ellner, G. F. Fussmann, and N. G. J. Hairston (2003). "Rapid evo-
- lution drives ecological dynamics in a predator prey system". In: *Nature* 424 (July), pp. 303–
- 817 306.
- Zhang, H., X. Gao, J. Unterman, and T. Arodz (July 2019). "Approximation Capabilities of Neural
- ODEs and Invertible Residual Networks". In: *arXiv*, pp. 1–11.

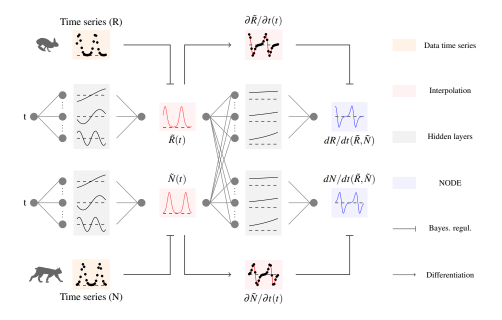


Figure 1: Overview of fitting neural ordinary differential equations (NODE) by Bayesian neural gradient matching (BNGM). In a first step we compute a continuous time approximation (interpolation) of each state variables, here the prey $\tilde{R}(t)$ and predator density $\tilde{N}(t)$ (red boxes). To do that we fit an ANN, that takes time as input, to each time series, via Bayesian regularisation. Interpolated dynamics of populations can then be computed by taking the derivative of the ANN with respect to time, $\partial \tilde{R}/\partial t$ and $\partial \tilde{N}/\partial t$. In a second step, we fit each NODE, dR/dt and dN/dt (blue boxes), to the interpolated dynamics. To do that we fit an ANN, which takes as input the interpolated variables $\tilde{R}(t)$ and $\tilde{N}(t)$, to the interpolated dynamics $\partial \tilde{R}/\partial t$ and $\partial \tilde{N}/\partial t$, via Bayesian regularisation.

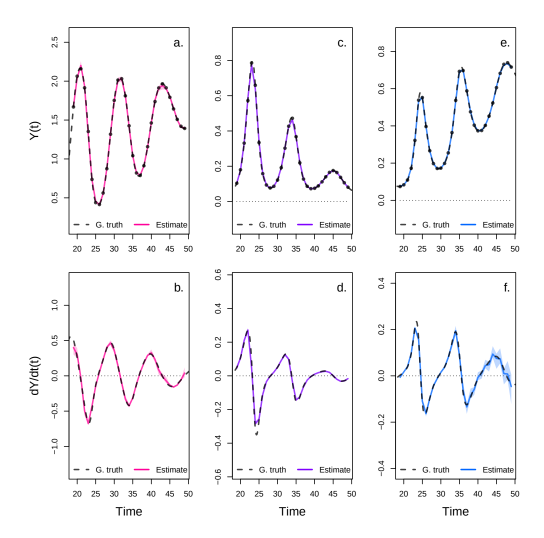


Figure 2: Interpolated density and dynamics of prey, intermediate, and top predators in the artificial system. This figure corresponds to the first step in the overview figure (Fig. 1). It shows the accuracy of the interpolated densities of prey (a.), intermediate (c.), and top predators (e.). We obtain interpolated densities by fitting observed densities (black dots) with ANNs that take time as input. The observed densities were obtained by sampling a tri-trophic prey-predator ODE model at regular time steps. We then derive interpolated dynamics (b., d., f.) by computing the temporal derivative of the interpolated densities with respect to time. In all graphs, the dashed line represents the ground truth, namely trajectories generated by the ODE model. The solid lines correspond to the interpolations. The shaded area shows the 90% confidence interval, obtained by approximately sampling the marginal posterior distributions.

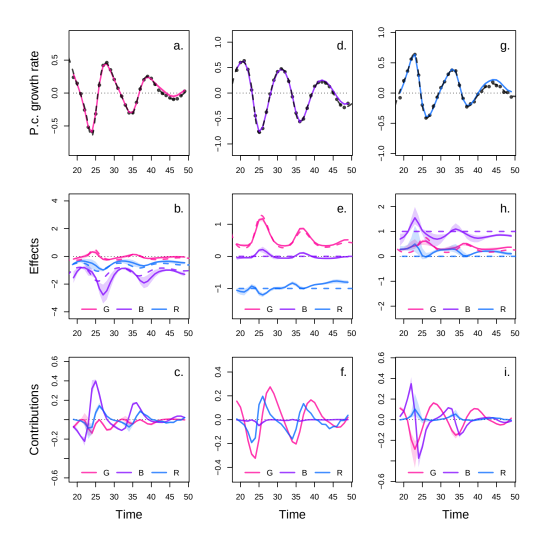


Figure 3: Drivers of dynamics of prey, intermediate, and top predator in the artificial system. This figure corresponds to the second step in the overview figure (Fig. 1). It displays the NODE nonparametric approximations of the per-capita growth rate of prey (a., b., c.), intermediate (d., e., f.), and top predators (g., h., i.). We obtain the NODE approximations (a., d., g., solid line) by fitting the interpolated per-capita growth rates (black dots) with ANNs that take population densities as input. We then estimate the direction of ecological interactions (effects, b., e., h.) by computing the derivative of the NODE approximations with respect to each density. Finally, we compute the strength of ecological interactions (contributions, c., f., i.) by multiplying the interpolated dynamics of each population (fig. 1, b., d., f.) with its effects. Dashed lines correspond to ground truth, obtained from the original trajectories of the tri-trophic ODE model. The shaded area shows the 90% confidence interval, obtained by approximately sampling the posterior distributions.

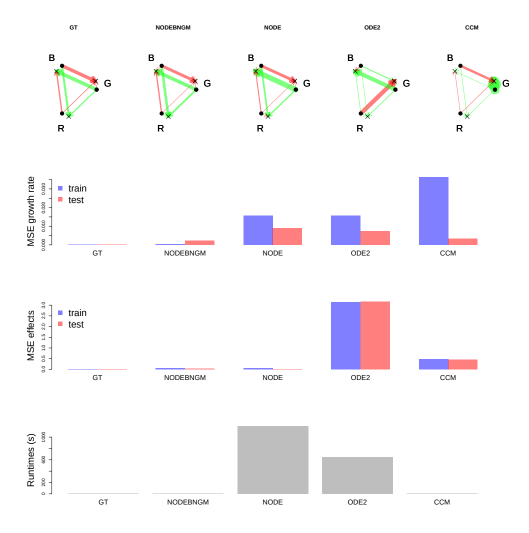


Figure 4: Runtimes and accuracy of NODEs fitted by BNGM compared to standard NODEs, **ODEs, and CCM.** The NODEBNGM method (nonparametric) involves fitting a NODE system by Bayesian neural gradient matching (BNGM). The NODE method (nonparametric) involves fitting a NODE system with an ODE solver. The ODE2 method (parametric) involves fitting an ODE system with quadratic functions of species densities with an ODE solver. The CCM method (nonparametric) involves computing locally linear approximations of the state space. For each method, we trained 30 models on the two first thirds of the artificial time series where ground truth is known (Fig. 2). We computed runtimes as the mean time (in seconds) required to train a single model. Using the best identified model, we predicted the growth rate and effects on the train and test set. We computed accuracies as the mean squared error of predictions vs ground truth (known from the equations that generated the data) (see Fig. S1-5 for more details). At the top, we show the dynamical interaction network of the system predicted by the best models, where G, B, R correspond to the prey, intermediate and top predator, respectively, Green and red colours correspond to positive and negative interactions, the width of arrows is proportional to relative total contribution to the growth rate of the population computed following the Geber method (Hairston et al. 2005; Bonnaffé, Sheldon, and Coulson 2021).

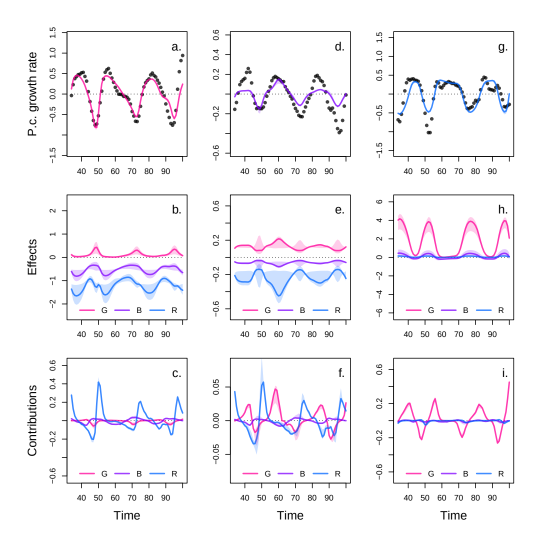


Figure 5: Drivers of dynamics of algae, flagellate, and rotifer in replicate B. This figure displays the NODE nonparametric approximations of the per-capita growth rate of algae (a., b., c.), flagellate (d., e., f.), and rotifer (g., h., i.). We obtain the NODE approximations (a., d., g., solid line) by fitting the interpolated per-capita growth rates (black dots) with ANNs that take population densities as input. We then estimate the direction of ecological interactions (effects, b., e., h.) by computing the derivative of the NODE approximations with respect to each density. Finally, we compute the strength of ecological interactions (contributions, c., f., i.) by multiplying the interpolated dynamics of each population with its effects. The shaded area shows the 90% confidence interval, obtained by approximately sampling the posterior distributions. The replicated time series were obtained by digitising the time series in Hiltunen et al. (2013).

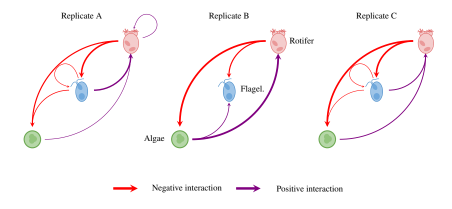


Figure 6: Interaction networks inferred from 3 replicated time series of algae, flagellate, and rotifers. This figure shows the direction and strength of ecological interactions inferred from 3 replicated sets of time series of algae, flagellate, and rotifer, using NODEs fitted by BNGM. The replicates A and C were analysed in the same way as replicate B (see fig. 5 for details). Red and purple arrows correspond to negative or positive mean effects. We estimated mean effects by averaging effects (i.e. derivative of NODE approximated per-capita growth rates with respect to each population density) across the time series. The width of the arrows is proportional to the relative strength of the ecological interaction. We compute the relative strength as the % of total contributions attributable to either algae, flagellate, or rotifer, obtained from summing the square of contributions of each species throughout the time series. For instance in replicate A, the relative strength of the effect of rotifer on algae is found by summing the square of the blue line in fig. 5 c., and comparing it to the sum of square of all contributions (Fig. 5 c., red, purple and blue lines). We provide the value of the mean effects and relative strengths in table 2. The replicated time series were obtained by digitising the time series in Hiltunen et al. (2013).

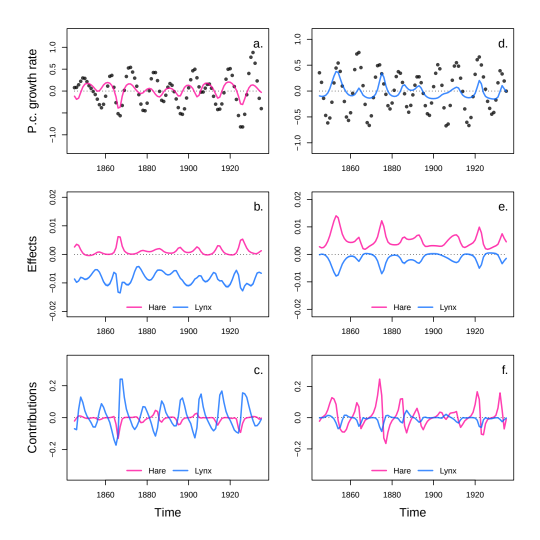


Figure 7: Drivers of dynamics of hare and lynx in the Odum and Barrett pelt count time series. This figure displays the NODE nonparametric approximations of the per-capita growth rate of hare (a., b., c.), and lynx (d., e., f.). We obtain the NODE approximations (a., d., solid line) by fitting the interpolated per-capita growth rates (black dots) with ANNs that take population densities as input. We then estimate the direction of ecological interactions (effects, b., e.) by computing the derivative of the NODE approximations with respect to each density. Finally, we compute the strength of ecological interactions (contributions, c., f.) by multiplying the interpolated dynamics of each population with its effects. The shaded area shows the 90% confidence interval, obtained by approximately sampling the posterior distributions.

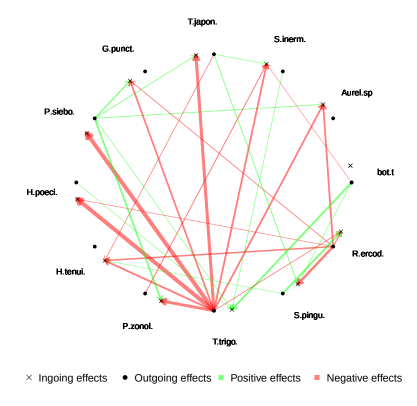


Figure 8: Dynamical interaction network of the Maizuru bay community. This figure summarises the results of the NODEBNGM analysis of the Maizuru bay community time series (Fig. S18). Bot.t corresponds to the temperature at sea bottom. Species are referred to by their shortened acronym. The main species are Aurelia sp., Sebastes inermis, Trachurus japonicus, Girella punctata, Pseudolabrus sieboldi, Halichoeres poecilopterus, Halichoeres tenuispinnis, Pterogobius zonoleucus, Tridentiger trigonocephalus, Sphyraena pinguis, and Rudarius ercodes. Red and green arrows correspond to negative or positive mean effects, obtained by averaging the sensitivity of the growth rate of a species to the density of other species across the time series. The width of the arrows is proportional to the relative total contribution (in %) of species density to the growth rate of other species, obtained by computing the relative sum the square of contributions across the time series. For the sake of clarity, we only display interactions which contribute to more than 10% of a change in the population growth rates.

Table 1: Summary of model runtimes. We measured the time required to perform 100 interpolations and 30 NODE fits to each variable in the systems. Replicate A, B, and C correspond to each replicated time series of the aglae, flagellate, and rotifer tri-trophic system (Hiltunen et al. 2013). The Hare-Lynx system correspond to the 90 years long time series of hare and lynx pelt counts (Odum and Barrett 1972). The Ushio system corresponds to the 100 time step long time series dataset of the 11 dominant species in the Maizuru bay community (Ushio et al. 2018). The number of time steps (N steps) is given for each time series. The total time per fit is obtain by dividing the total time in seconds by the number of fits (i.e. 130). These results were obtained on a macbook pro M1 MAX 2022, in base R (v4.0.2), with non-optimised code.

			Interpolation		NODE fit			
System	N var.	N steps	N fits	time (s)	N fits	time (s)	total	total p. fit
Replicate A	3	66	100	234.45	30	80.41	314.86	5.02
Replicate B	3	66	100	238.01	30	81.13	319.14	5.08
Replicate C	3	40	100	136.51	30	49.94	186.45	3.03
Hare-lynx	2	90	100	341.74	30	21.55	363.29	4.14
Ushio	12	100	100	806.12	30	604.42	1410.54	28.21

Table 2: Comparison of the direction and strentgh of ecological interactions estimated by BNGM across 3 replicated tri-trophic microcosms. Mean effects are obtained by averaging the effect of one species on the growth rate of another throughout the time series. The % of total contributions is obtained by summing the square of contributions of one species density to the growth of the other at each time step throughout the time series, then by computing the proportion of total change that it accounts for. The variables G, B, and R correspond to the population density of algae, flagellate, and rotifer respectively. r^2 corresponds to the r squared of the NODE nonparametric approximation of the pre-capita growth rate of the three species.

		G	В	R
Replicate A	r^2	0.11	0.37	0.47
Mean effects	on G	-0.08	-1.14	-1.13
	on B	0.28	-0.21	-0.66
	on R	0.60	1.09	0.32
% of total contributions	to G	0.01	0.34	0.65
	to B	0.02	0.04	0.93
	to R	0.26	0.66	0.08
Replicate B	r^2	0.52	0.4	0.65
Mean effects	on G	0.12	-0.53	-1.23
	on B	0.12	-0.06	-0.25
	on R	1.83	0.10	0.07
% of total contributions	to G	0.02	0.03	0.95
	to B	0.38	0.02	0.61
	to R	0.99	0.00	0.01
Replicate C	r^2	0.59	0.32	0.73
Mean effects	on G	0.09	-0.49	-1.96
	on B	0.04	-0.19	-0.65
	on R	1.07	0.79	-0.01
% of total contributions	to G	0.00	0.08	0.91
	to B	0.01	0.10	0.88
	to R	0.49	0.51	0.00

6 Supplementary

A Bayesian regularisation

The fitting of the models is performed in a Bayesian framework, considering normal error structure for the residuals, and normal prior density distributions on the parameters,

$$p(\boldsymbol{\omega}|\mathcal{D}) \propto p(\mathcal{D}|\boldsymbol{\omega})p(\boldsymbol{\omega}),$$
 (13)

where θ is the parameter vector of the model, and \mathcal{D} the evidence, namely the data that the model is fitted to. Assuming a normal likelihood for the residuals given the evidence we get

$$p(\mathcal{D}|\boldsymbol{\omega}) = \prod_{i=1}^{N} \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left\{-\frac{e_i(\mathcal{D}, \boldsymbol{\omega})^2}{2\sigma^2}\right\},\tag{14}$$

where $e_i(\mathcal{D}, \omega)$ are the residuals of the model given the parameters, and the evidence. In the case of the interpolation, the residuals correspond to the observation error $\varepsilon^{(o)}$ (equation 3). In the case of the NODE approximation, they correspond to the process error $\varepsilon^{(p)}$ (equation 7). N is the number of data points, either observations in the case of the interpolation, $N^{(o)}$, or interpolated points in the case of the NODE fitting, $N^{(p)}$.

The prior probability density functions for the parameters are given by

$$p(\boldsymbol{\omega}) = \prod_{j=1}^{M} \frac{1}{\sqrt{2\pi\delta_j^2}} \exp\left\{-\frac{\omega_j^2}{2\delta_j^2}\right\},\tag{15}$$

where M is the number of parameters in the models. The parameter δ_j controls the dispersion of the priors, and thereby the complexity/level of constraint of the model.

Bayesian regularisation simply amounts to constraining the values of the parameters in the model 834 to be close to a desired value. Usually, parameters are constrained by choosing normal priors 835 centered about 0. In this case, the standard deviation of the normal priors governs the range of values that the parameters can take, and hence constrains more or less strongly the behaviour of the 837 model (Cawley and Talbot 2007). There is no standard approach for choosing δ . Low values of 838 dispersion may increase constraint on parameters too drastically, which would lead to underfitting, 839 and result in a reduction of the variance of parameter estimates and bias mean estimates towards 0. In contrast, too high values of dispersion may lead to overfitting, by allowing for more complex 841 shapes. To account for this, we optimise models on the second-level of inference. This means that 842 we are finding the optimal value of δ , in addition to optimising the model parameters.

In practice, choosing the level of constraint is difficult, Cawley and Talbot hence developed a criterion to perform model selection on the second level of inference. They proposed to optimise the marginal posterior distribution by averaging out the dispersion of the priors. With an appropriate choice of prior, the dispersion can be integrated out, leaving us with a formula for the posterior that only depends on the parameters of the model,

$$\log P(\boldsymbol{\omega}|\mathcal{D}) \propto -\frac{N}{2}\log\left(\sum_{i=1}^{N}e_{i}(\mathcal{D},\boldsymbol{\omega})^{2}\right) - \frac{M}{2}\log\left(\sum_{j=1}^{M}\omega_{j}^{2}\right),\tag{16}$$

where $P(\boldsymbol{\omega}|\mathcal{D})$ denotes the marginal posterior density, \mathcal{D} denotes the evidence, N and M denote

the number of data points and parameters, respectively, e_i denote the residuals, and ω denote the parameters of the model. The construction is elegant because it is not sensitive to the choice of prior hyperparameters, and simple as it amounts to optimising the log of the sum of squares, rather than the sum of squares (in the case of normal ordinary least square).

The issue with this formula is that the marginal posterior density is not finite when the parameters are 0, which can lead to underfitting. In this paper we use a modified criterion, which corrects for that problem,

$$\log P(\boldsymbol{\omega}|\mathcal{D}) \propto -\frac{N}{2}\log\left(1 + \sum_{i=1}^{N} e_i(\mathcal{D}, \boldsymbol{\omega})^2\right) - \frac{M}{2}\log\left(1 + \sum_{j=1}^{M} \boldsymbol{\omega}_j^2\right),\tag{17}$$

where the marginal posterior density depends only on the residuals of the model when the parameters are equal to 0, and otherwise depends on both the parameters and the residuals. This construction can be obtained simply by assuming a gamma prior for the parameters $p(\xi) \propto \frac{1}{\xi} \exp{\{-\xi/2\}}$, where ξ is the regularisation parameter, instead of the improper Jeffreys' prior that Cawley and Talbot used in their original study, namely $p(\xi) \propto \frac{1}{\xi}$.

We provide below details of the calculation of this modified criterion, starting by averaging out the regularisation parameter. We let $\xi := 1/\delta^2$, assuming that all parameters are controlled by the

same regularisation parameter, which gives the following equation for the prior distribution,

$$p(\omega|\xi) = \prod_{j=1}^{M} \frac{1}{\sqrt{2\pi/\xi}} \exp\left\{-\frac{\xi}{2}\omega_j^2\right\}. \tag{18}$$

To follow the proof given by Cawley and Talbot we further use the following notation, d=M, $Z_{\Omega}:=(2\pi/\xi)^{d/2}$, and $\Omega:=\sum_{j}\omega_{j}^{2}$, which yields

$$p(\omega|\xi) = Z_{\Omega}(\xi)^{-1} \exp\left\{-\frac{\xi}{2}\Omega(\omega)\right\}. \tag{19}$$

We can then integrate out the regularisation parameter by computing the marginal prior distribu-

$$P(\omega) = \int p(\omega|\xi)p(\xi)d\xi. \tag{20}$$

This expression can be solved analytically with the right choice of prior. Cawley and Talbot use $p(\xi) \propto 1/\xi$, but instead we choose $p(\xi) \propto 1/\xi \exp\{-\xi/2\}$. By assuming positive support for ξ and expanding out the marginal prior distribution we get,

$$P(\omega) = \int_0^\infty \left(\frac{2\pi}{\xi}\right)^{-d/2} \exp\left\{-\frac{\xi}{2}\Omega(\omega)\right\} \frac{1}{\xi} \exp\left\{-\frac{\xi}{2}\right\} d\xi$$
$$= (2\pi)^{-d/2} \int_0^\infty \xi^{d/2-1} \exp\left\{-\frac{\xi}{2}\left(\Omega(\omega) + 1\right)\right\} d\xi$$
(21)

We use the gamma integral, $\int_0^\infty x^{\nu-1} \exp\{-\mu x\} = \Gamma(\nu)/\mu^{\nu}$ (Cawley and Talbot 2007), to obtain,

$$P(\omega) = (2\pi)^{-d/2} \frac{\Gamma(d/2)}{(1/2)^{d/2} (\Omega(\omega) + 1)^{d/2}}$$
(22)

Finally, by applying the logarithm and discarding constants, we obtain the final expression for the

marginal prior distribution,

$$\log P(\omega) \propto -\frac{d}{2} \log \left(\Omega(\omega) + 1\right)$$

$$\propto -\frac{M}{2} \log \left(1 + \sum_{j=0}^{M} \omega_{j}^{2}\right)$$
(23)

The same procedure can be applied to derive the marginal likelihood, by noting $d:=N,\,\xi:=1/\sigma^2,$

877
$$Z_\Omega:=(2\pi/\xi)^{d/2},\,\Omega(\omega):=\sum_i e_i^2.$$

B Complementary results benchmark analysis

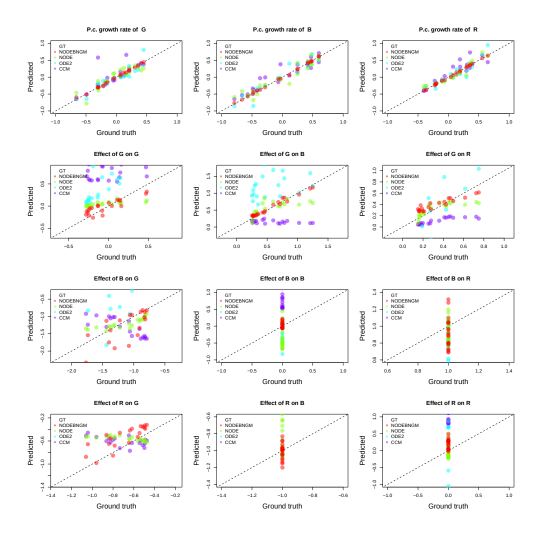


Figure S1: Train set accuracy of predicted per-capita growth rates and effects estimated by NODEBNGM, standard NODEs, ODE2, and CCM. The NODEBNGM method (nonparametric) involves fitting a NODE system by Bayesian neural gradient matching (BNGM). The NODE method (nonparametric) involves fitting a NODE system with an ODE solver. The ODE2 method (parametric) involves fitting an ODE system with quadratic functions of species densities with an ODE solver. The CCM method (nonparametric) involves computing locally linear approximations of the state space. For each method, we trained 30 models on the two first thirds of the artificial time series where ground truth is known (Fig. 2). For each plot, the x-axis corresponds to the ground truth, known from the equations that generated the artificial time series, and the y-axis corresponds to the prediction of the best model. Effects are computed as the sensitivity (i.e. derivative) of the per-capita growth rate with respect to each species density G, B, and R, namely the prey, intermediate and top predator.

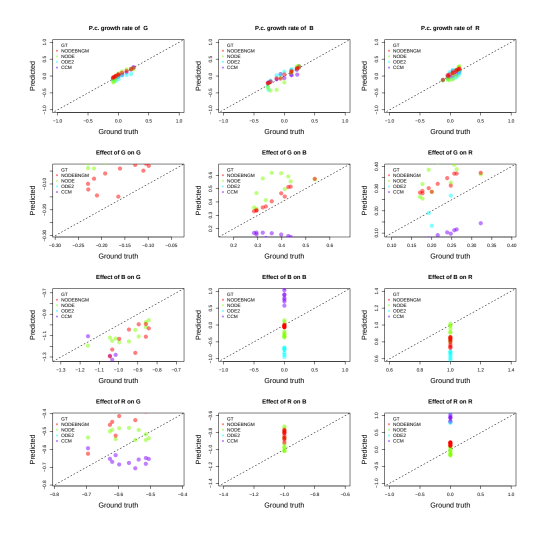


Figure S2: Test set accuracy of predicted per-capita growth rates and effects estimated by NODEBNGM, standard NODEs, ODE2, and CCM. The NODEBNGM method (nonparametric) involves fitting a NODE system by Bayesian neural gradient matching (BNGM). The NODE method (nonparametric) involves fitting a NODE system with an ODE solver. The ODE2 method (parametric) involves fitting an ODE system with quadratic functions of species densities with an ODE solver. The CCM method (nonparametric) involves computing locally linear approximations of the state space. For each method, we trained 30 models on the two first thirds of the artificial time series where ground truth is known (Fig. 2). For each plot, the x-axis corresponds to the ground truth, known from the equations that generated the artificial time series, and the y-axis corresponds to the prediction of the best model. Effects are computed as the sensitivity (i.e. derivative) of the per-capita growth rate with respect to each species density G, B, and R, namely the prey, intermediate and top predator.

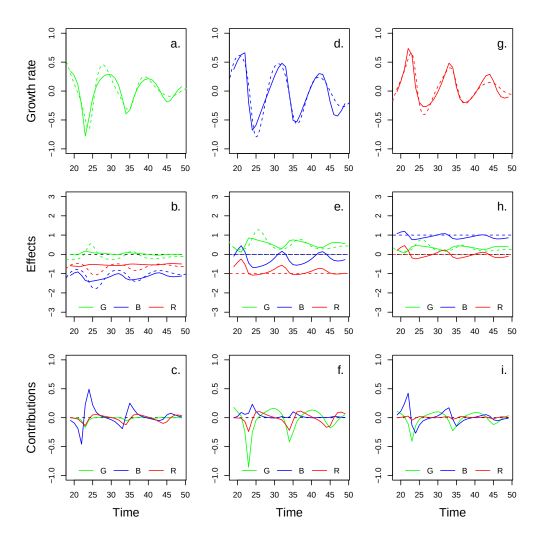


Figure S3: Effects and contributions inferred by standard NODEs. The standard NODE model nonparametrically approximates the per-capita growth rate of the 3 species with an ANN featuring a single layer, 3 input nodes, 10 hidden nodes, 3 outputs. 30 models are fitted to the two first third of the time series using BFGS and a Runge-Kutta ODE solver. The graphs present the predictions obtained for the model that maximises posterior density of the network parameters given the time series. We estimate the direction of ecological interactions (effects) by computing the derivative of the per-capita growth rate approximations with respect to each density. Finally, we compute the strength of ecological interactions (contributions) by multiplying the interpolated dynamics of each population with its effects.

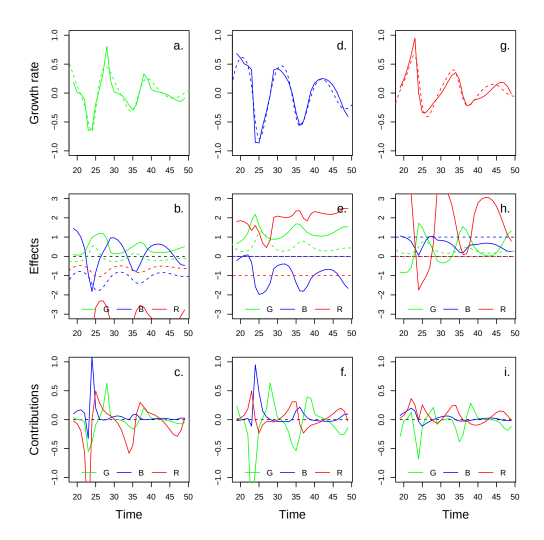


Figure S4: Effects and contributions inferred by parametric ODE. The ODE2 model parametrically approximates the per-capita growth rate of the 3 species with second order polynomial functions. 30 models are fitted to the two first third of the time series using BFGS and a Runge-Kutta ODE solver. The graphs present the predictions obtained for the model that maximises posterior density of the network parameters given the time series. We estimate the direction of ecological interactions (effects) by computing the derivative of the per-capita growth rate approximations with respect to each density. Finally, we compute the strength of ecological interactions (contributions) by multiplying the interpolated dynamics of each population with its effects.

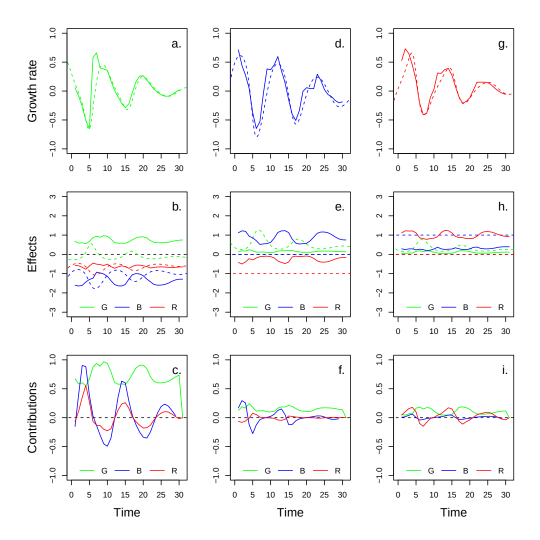


Figure S5: Effects and contributions inferred by CCM. The CCM method nonparametrically approximates the state space from which it derives the sensitivity of population dynamics to a change in the density of the species. We use the rEDM implementation and derived our code from the three species example provided in the package tutorial (v1.13.1, Sugihara et al. 2012). We calculated the dynamics and per capita growth rate using finite differences, as the standard library does not readily provide estimates. The effects correspond to the s-map coefficients. Finally, we compute the strength of ecological interactions (contributions) by multiplying the interpolated dynamics of each population with its effects.

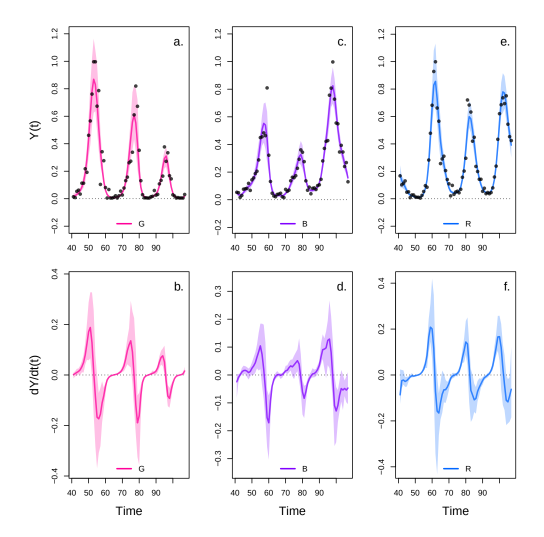


Figure S6: Interpolation of state and dynamics of algae, flagellate, and rotifer density in replicate A. Graph a., c., and e. display the neural interpolation of the population density of algae (G), flagellate (B), and rotifer (R), respectively (obtained with Eq. 7). Graph b., d., and f. show the corresponding interpolated dynamics, obtained by differentiating the interpolation of the states with respect to time (Eq. 5). The shaded areas represent the 90% confidence interval on estimates, obtained by anchored ensembling of the log marginal posterior distribution (Eq. 7) (Pearce et al. 2018). Time series are obtained from digitising the time series in Hiltunen et al. 2013.

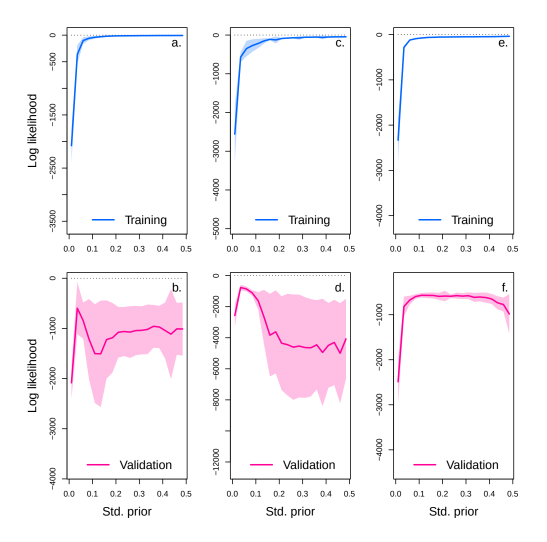


Figure S7: Cross-validation plot of the NODE analysis of replicate A. The x-axis of the graphs correspond to the standard deviation of the prior distribution of the NODE parameters, which constrains the nonlinearity of the nonparametric approximation of the NODEs. Small values of standard deviation correspond to a linear model, while higher values (towards 0.5) correspond to a highly nonlinear model. Time series of algae, flagellate, and rotifer are split in half to create a train set and a test set. The model is fitted to the train set for increasing value of standard deviation (from 0.05 to 0.5 by 0.05 increments), and evaluated on the test set. Graph a., c., and e. show the log likelihood of the NODE system fitted by BNGM to the train set of algae, flagellate, and rotifer, respectively. Graph b., d., and f. show the log likelihood of the fitted NODE, evaluated on the corresponding test set. The shaded areas represent the 90% confidence interval on estimates, obtained by anchored ensembling of the log posterior distribution (Eq. 8) (Pearce et al. 2018).

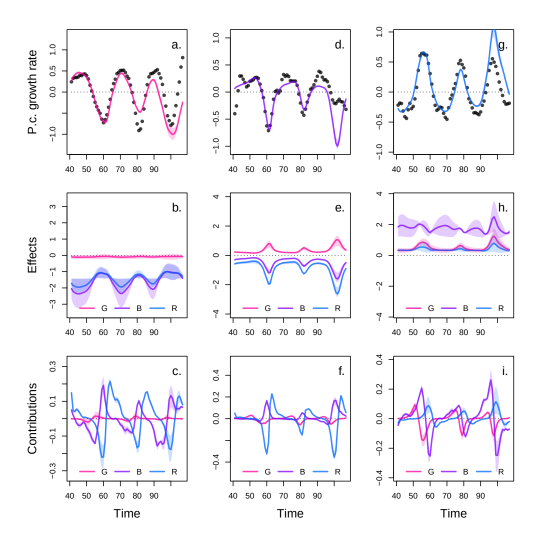


Figure S8: Drivers of dynamics of algae, flagellate, and rotifer in replicate A. This figure displays the NODE nonparametric approximations of the per-capita growth rate of algae (a., b., c.), flagellate (d., e., f.), and rotifer (g., h., i.). We obtain the NODE approximations (a., d., g., solid line) by fitting the interpolated per-capita growth rates (black dots) with ANNs that take population densities as input. We then estimate the direction of ecological interactions (effects, b., e., h.) by computing the derivative of the NODE approximations with respect to each density. Finally, we compute the strength of ecological interactions (contributions, c., f., i.) by multiplying the interpolated dynamics of each population with its effects. The shaded area shows the 90% confidence interval, obtained by approximately sampling the posterior distributions. The replicated time series were obtained by digitising the time series in Hiltunen et al. (2013).

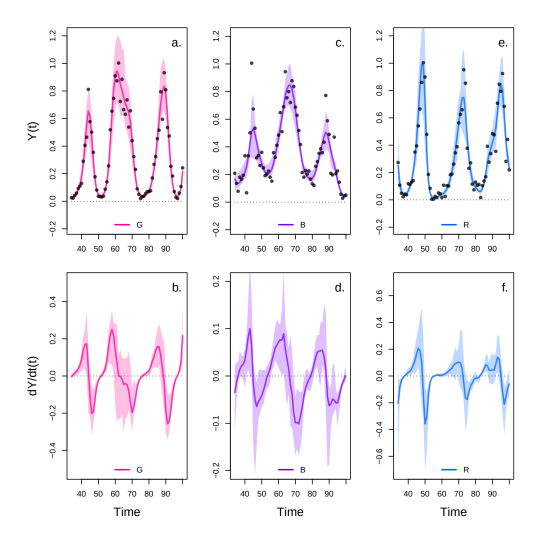


Figure S9: Interpolation of state and dynamics of algae, flagellate, and rotifer density in replicate B. Graph a., c., and e. display the neural interpolation of the population density of algae (G), flagellate (B), and rotifer (R), respectively (obtained with Eq. 7). Graph b., d., and f. show the corresponding interpolated dynamics, obtained by differentiating the interpolation of the states with respect to time (Eq. 5). The shaded areas represent the 90% confidence interval on estimates, obtained by anchored ensembling of the log marginal posterior distribution (Eq. 7) (Pearce et al. 2018). Time series are obtained from digitising the time series in Hiltunen et al. 2013.

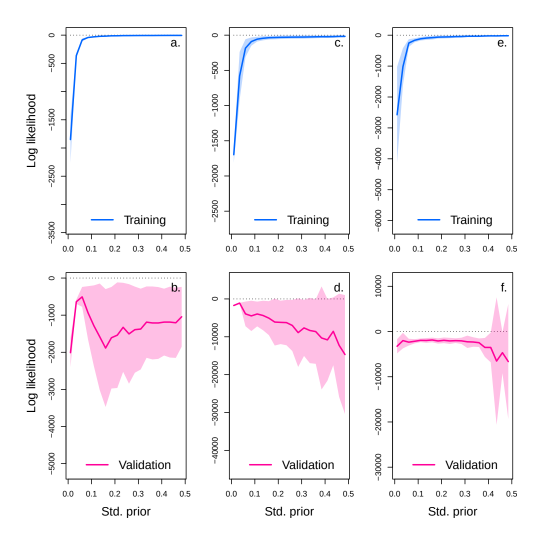


Figure S10: Cross-validation plot of the NODE analysis of replicate B. The x-axis of the graphs correspond to the standard deviation of the prior distribution of the NODE parameters, which constrains the nonlinearity of the nonparametric approximation of the NODEs. Small values of standard deviation correspond to a linear model, while higher values (towards 0.5) correspond to a highly nonlinear model. Time series of algae, flagellate, and rotifer are split in half to create a train set and a test set. The model is fitted to the train set for increasing value of standard deviation (from 0.05 to 0.5 by 0.05 increments), and evaluated on the test set. Graph a., c., and e. show the log likelihood of the NODE system fitted by BNGM to the train set of algae, flagellate, and rotifer, respectively. Graph b., d., and f. show the log likelihood of the fitted NODE, evaluated on the corresponding test set. The shaded areas represent the 90% confidence interval on estimates, obtained by anchored ensembling of the log posterior distribution (Eq. 8) (Pearce et al. 2018).

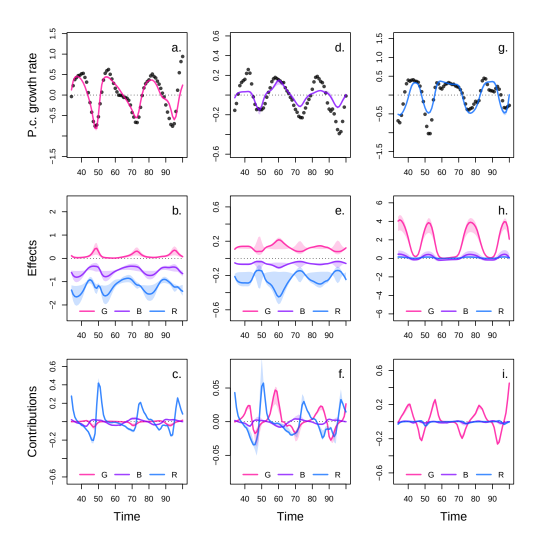


Figure S11: Drivers of dynamics of algae, flagellate, and rotifer in replicate B. This figure displays the NODE nonparametric approximations of the per-capita growth rate of algae (a., b., c.), flagellate (d., e., f.), and rotifer (g., h., i.). We obtain the NODE approximations (a., d., g., solid line) by fitting the interpolated per-capita growth rates (black dots) with ANNs that take population densities as input. We then estimate the direction of ecological interactions (effects, b., e., h.) by computing the derivative of the NODE approximations with respect to each density. Finally, we compute the strength of ecological interactions (contributions, c., f., i.) by multiplying the interpolated dynamics of each population with its effects. The shaded area shows the 90% confidence interval, obtained by approximately sampling the posterior distributions. The replicated time series were obtained by digitising the time series in Hiltunen et al. (2013).

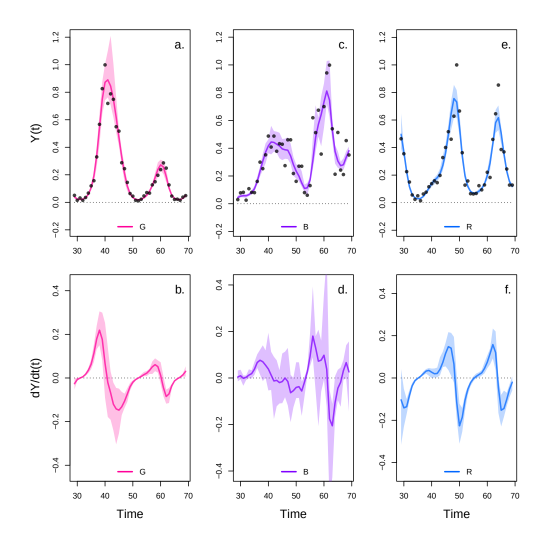


Figure S12: Interpolation of state and dynamics of algae, flagellate, and rotifer density in replicate B. Graph a., c., and e. display the neural interpolation of the population density of algae (G), flagellate (B), and rotifer (R), respectively (obtained with Eq. 7). Graph b., d., and f. show the corresponding interpolated dynamics, obtained by differentiating the interpolation of the states with respect to time (Eq. 5). The shaded areas represent the 90% confidence interval on estimates, obtained by anchored ensembling of the log marginal posterior distribution (Eq. 7) (Pearce et al. 2018). Time series are obtained from digitising the time series in Hiltunen et al. 2013.

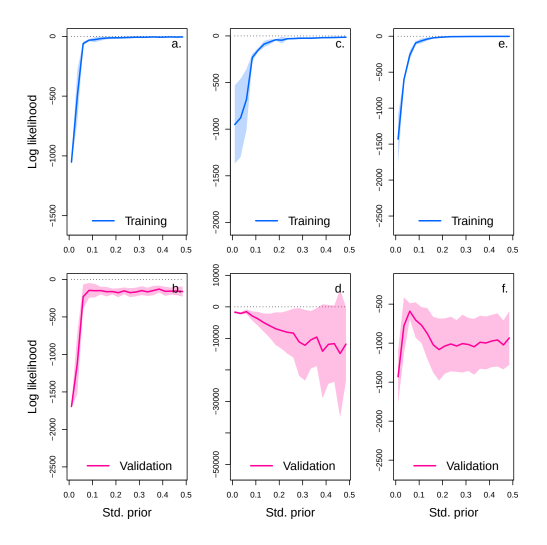


Figure S13: Cross-validation plot of the NODE analysis of replicate C. The x-axis of the graphs correspond to the standard deviation of the prior distribution of the NODE parameters, which constrains the nonlinearity of the nonparametric approximation of the NODEs. Small values of standard deviation correspond to a linear model, while higher values (towards 0.5) correspond to a highly nonlinear model. Time series of algae, flagellate, and rotifer are split in half to create a train set and a test set. The model is fitted to the train set for increasing value of standard deviation (from 0.05 to 0.5 by 0.05 increments), and evaluated on the test set. Graph a., c., and e. show the log likelihood of the NODE system fitted by BNGM to the train set of algae, flagellate, and rotifer, respectively. Graph b., d., and f. show the log likelihood of the fitted NODE, evaluated on the corresponding test set. The shaded areas represent the 90% confidence interval on estimates, obtained by anchored ensembling of the log posterior distribution (Eq. 8) (Pearce et al. 2018).

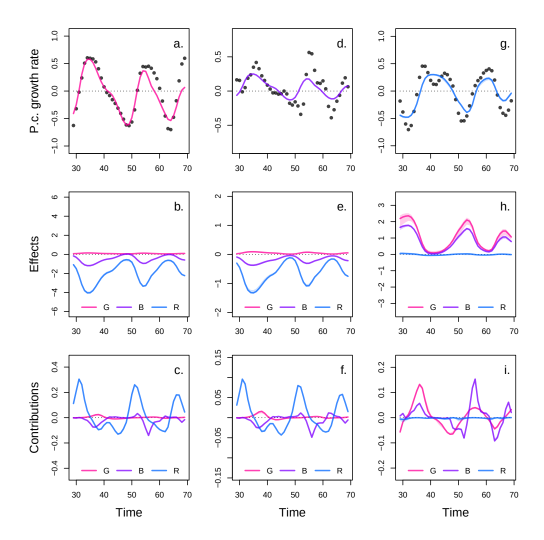


Figure S14: Drivers of dynamics of algae, flagellate, and rotifer in replicate C. This figure displays the NODE nonparametric approximations of the per-capita growth rate of algae (a., b., c.), flagellate (d., e., f.), and rotifer (g., h., i.). We obtain the NODE approximations (a., d., g., solid line) by fitting the interpolated per-capita growth rates (black dots) with ANNs that take population densities as input. We then estimate the direction of ecological interactions (effects, b., e., h.) by computing the derivative of the NODE approximations with respect to each density. Finally, we compute the strength of ecological interactions (contributions, c., f., i.) by multiplying the interpolated dynamics of each population with its effects. The shaded area shows the 90% confidence interval, obtained by approximately sampling the posterior distributions. The replicated time series were obtained by digitising the time series in Hiltunen et al. (2013).

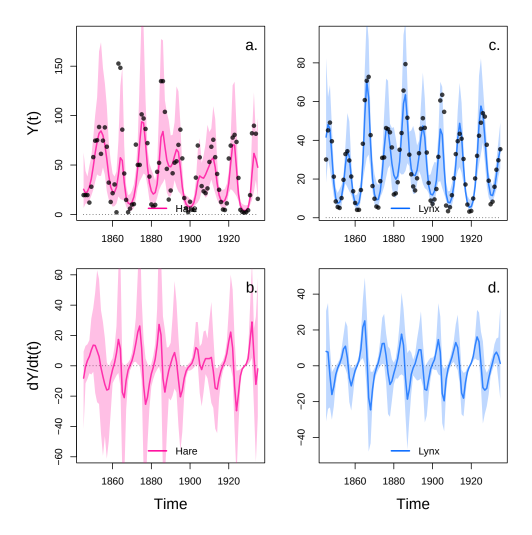


Figure S15: Interpolation of state and dynamics of hare and lynx. Graph a. and c. display the neural interpolation of the population density of hare and lynx respectively (obtained with Eq. 7). Graph b. and d. show the corresponding interpolated dynamics, obtained by differentiating the interpolation of the states with respect to time (Eq. 5). The shaded areas represent the 90% confidence interval on estimates, obtained by anchored ensembling of the log marginal posterior distribution (Eq. 7) (Pearce et al. 2018). Time series are obtained from Bonnaffé, Sheldon, and Coulson 2021, originally from Odum and Barrett 1972.

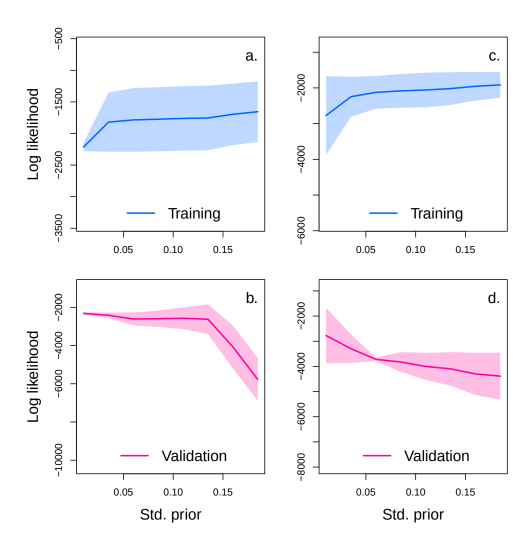


Figure S16: Cross-validation plot of the NODE analysis of the hare-lynx system. The x-axis of the graphs correspond to the standard deviation of the prior distribution of the NODE parameters, which constrains the nonlinearity of the nonparametric approximation of the NODEs. Small values of standard deviation correspond to a linear model, while higher values (towards 0.5) correspond to a highly nonlinear model. Time series of algae, flagellate, and rotifer are split in half to create a train set and a test set. The model is fitted to the train set for increasing value of standard deviation (from 0.0 to 0.2 by 0.025 increments), and evaluated on the test set. Graph a., c., and e. show the log likelihood of the NODE system fitted by BNGM to the train set of algae, flagellate, and rotifer, respectively. Graph b., d., and f. show the log likelihood of the fitted NODE, evaluated on the corresponding test set. The shaded areas represent the 90% confidence interval on estimates, obtained by anchored ensembling of the log posterior distribution (Eq. 8) (Pearce et al. 2018).

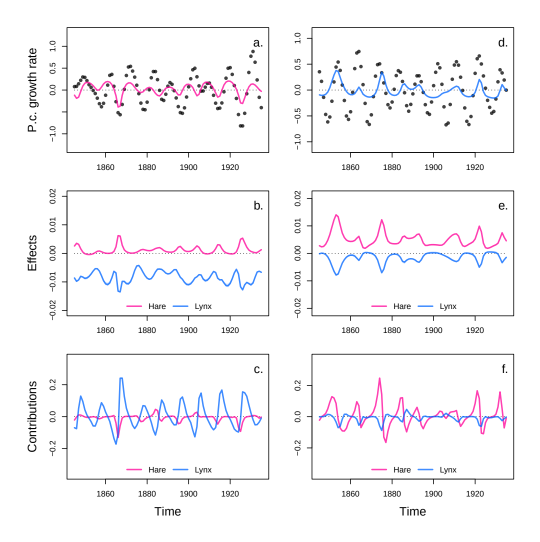


Figure S17: Drivers of dynamics of hare and lynx in the Odum and Barrett pelt count time series. This figure displays the NODE nonparametric approximations of the per-capita growth rate of hare (a., b., c.), and lynx (d., e., f.). We obtain the NODE approximations (a., d., solid line) by fitting the interpolated per-capita growth rates (black dots) with ANNs that take population densities as input. We then estimate the direction of ecological interactions (effects, b., e.) by computing the derivative of the NODE approximations with respect to each density. Finally, we compute the strength of ecological interactions (contributions, c., f.) by multiplying the interpolated dynamics of each population with its effects. The shaded area shows the 90% confidence interval, obtained by approximately sampling the posterior distributions.

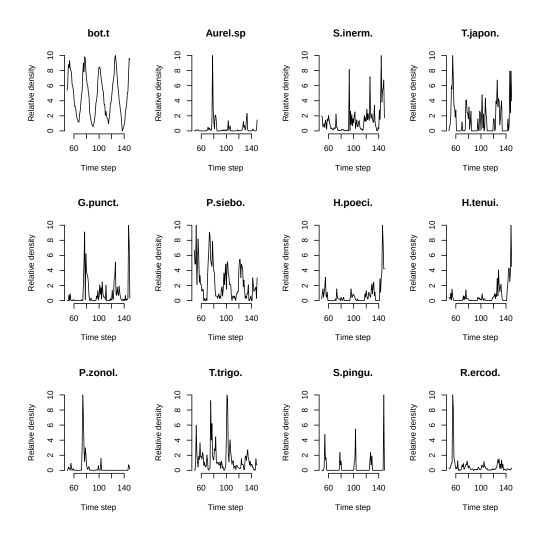


Figure S18: Time series of the fish community in Ushio et al. 2018. The time series was collected for 12-years on a fortnightly basis, for 15 dominant species in the Maizuru bay in Japan. We focus on the 11 species and the 100 time steps with the least sparse abundance records. Bot.t corresponds to water temperature near the bottom. The main species are *Aurelia sp.*, *Sebastes inermis*, *Trachurus japonicus*, *Girella punctata*, *Pseudolabrus sieboldi*, *Halichoeres poecilopterus*, *Halichoeres tenuispinnis*, *Pterogobius zonoleucus*, *Tridentiger trigonocephalus*, *Sphyraena pinguis*, and *Rudarius ercodes*.

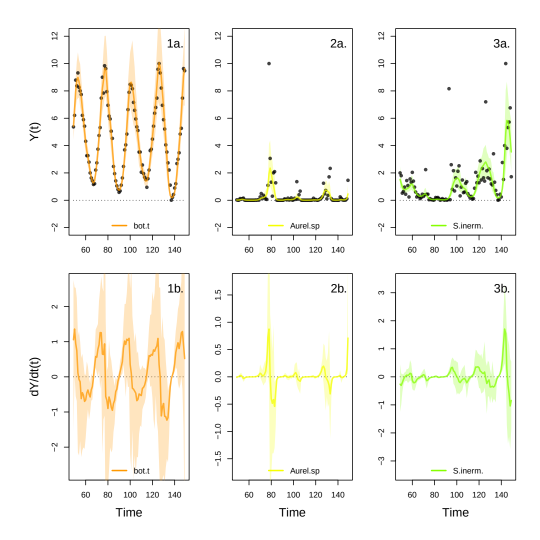


Figure S19: Interpolation of state and dynamics of species abundance in the Maizuru bay community. Graphs a. display the neural interpolations of the population density (obtained with Eq. 7). Graphs b. show the corresponding interpolated dynamics, obtained by differentiating the interpolation of the states with respect to time (Eq. 5). The shaded areas represent the 90% confidence interval on estimates, obtained by anchored ensembling of the log marginal posterior distribution (Eq. 7)

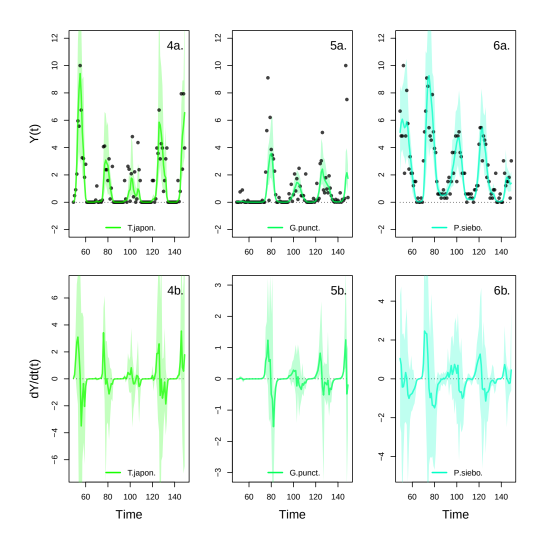


Figure S20: Interpolation of state and dynamics of species abundance in the Maizuru bay community. Graphs a. display the neural interpolations of the population density (obtained with Eq. 7). Graphs b. show the corresponding interpolated dynamics, obtained by differentiating the interpolation of the states with respect to time (Eq. 5). The shaded areas represent the 90% confidence interval on estimates, obtained by anchored ensembling of the log marginal posterior distribution (Eq. 7)

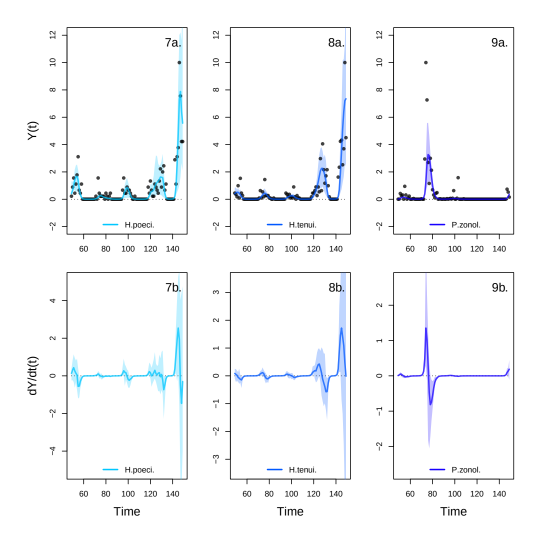


Figure S21: Interpolation of state and dynamics of species abundance in the Maizuru bay community. Graphs a. display the neural interpolations of the population density (obtained with Eq. 7). Graphs b. show the corresponding interpolated dynamics, obtained by differentiating the interpolation of the states with respect to time (Eq. 5). The shaded areas represent the 90% confidence interval on estimates, obtained by anchored ensembling of the log marginal posterior distribution (Eq. 7)

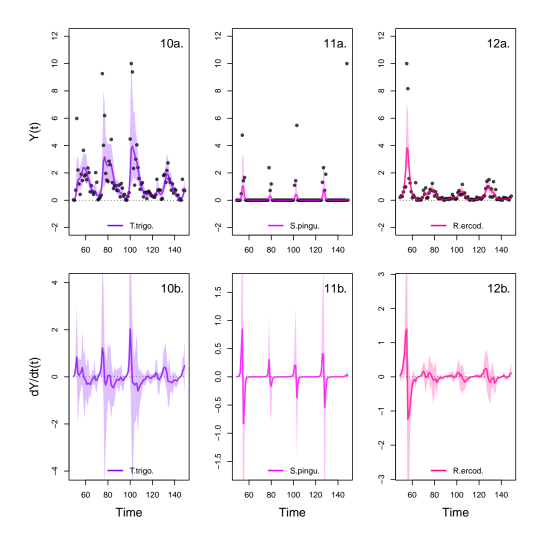


Figure S22: Interpolation of state and dynamics of species abundance in the Maizuru bay community. Graphs a. display the neural interpolations of the population density (obtained with Eq. 7). Graphs b. show the corresponding interpolated dynamics, obtained by differentiating the interpolation of the states with respect to time (Eq. 5). The shaded areas represent the 90% confidence interval on estimates, obtained by anchored ensembling of the log marginal posterior distribution (Eq. 7)

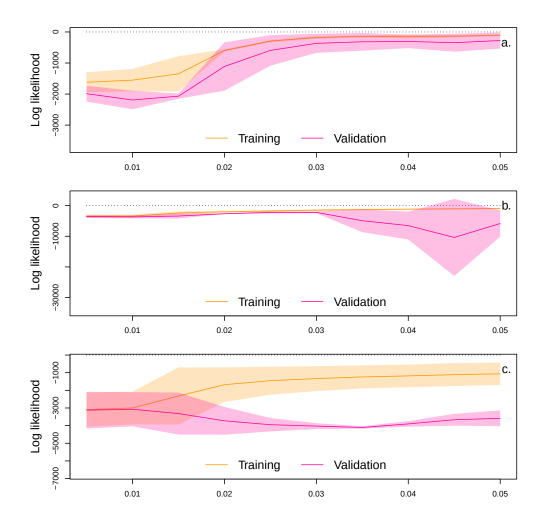


Figure S23: Cross-validation plot of the NODE analysis of the Maizuru bay community. The x-axis of the graphs correspond to the standard deviation of the prior distribution of the NODE parameters, which constrains the nonlinearity of the nonparametric approximation of the NODEs. Small values of standard deviation correspond to a linear model, while higher values correspond to a highly nonlinear model. Time series are split in three thirds to create a train, validation, and test set. The model is fitted to the train set (i.e. first third) for increasing value of standard deviation (from 0.0 to 0.1 by 0.01 increments), and evaluated on the validation set. The operation is repeated by swapping the training and validation set. The graphs show the log likelihood of the NODE system fitted by BNGM to the train set (in orange), and evaluated on the corresponding validation set (in red). The shaded areas represent the 90% confidence interval on estimates, obtained by anchored ensembling of the log posterior distribution (Eq. 8) (Pearce et al. 2018).

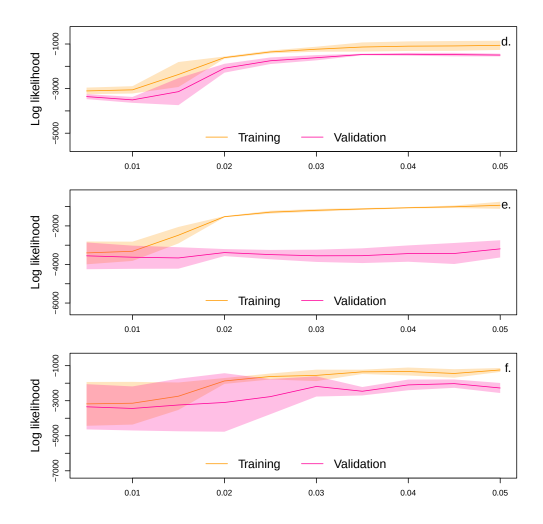


Figure S24: Cross-validation plot of the NODE analysis of the Maizuru bay community. The x-axis of the graphs correspond to the standard deviation of the prior distribution of the NODE parameters, which constrains the nonlinearity of the nonparametric approximation of the NODEs. Small values of standard deviation correspond to a linear model, while higher values correspond to a highly nonlinear model. Time series are split in three thirds to create a train, validation, and test set. The model is fitted to the train set (i.e. first third) for increasing value of standard deviation (from 0.0 to 0.1 by 0.01 increments), and evaluated on the validation set. The operation is repeated by swapping the training and validation set. The graphs show the log likelihood of the NODE system fitted by BNGM to the train set (in orange), and evaluated on the corresponding validation set (in red). The shaded areas represent the 90% confidence interval on estimates, obtained by anchored ensembling of the log posterior distribution (Eq. 8) (Pearce et al. 2018).

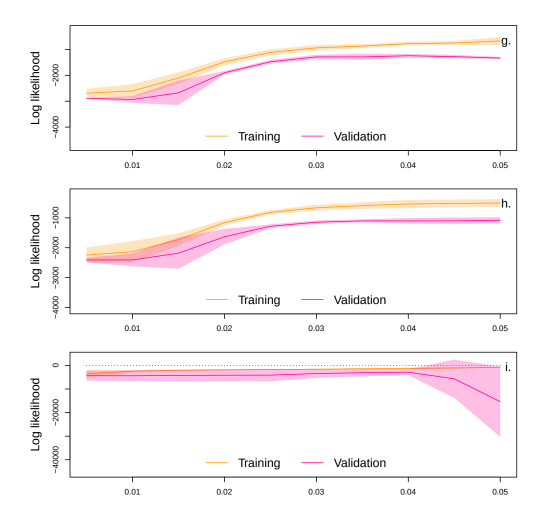


Figure S25: Cross-validation plot of the NODE analysis of the Maizuru bay community. The x-axis of the graphs correspond to the standard deviation of the prior distribution of the NODE parameters, which constrains the nonlinearity of the nonparametric approximation of the NODEs. Small values of standard deviation correspond to a linear model, while higher values correspond to a highly nonlinear model. Time series are split in three thirds to create a train, validation, and test set. The model is fitted to the train set (i.e. first third) for increasing value of standard deviation (from 0.0 to 0.1 by 0.01 increments), and evaluated on the validation set. The operation is repeated by swapping the training and validation set. The graphs show the log likelihood of the NODE system fitted by BNGM to the train set (in orange), and evaluated on the corresponding validation set (in red). The shaded areas represent the 90% confidence interval on estimates, obtained by anchored ensembling of the log posterior distribution (Eq. 8) (Pearce et al. 2018).

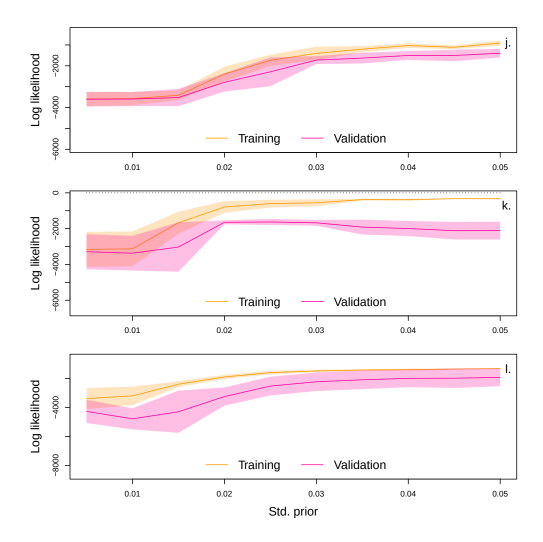


Figure S26: Cross-validation plot of the NODE analysis of the Maizuru bay community. The x-axis of the graphs correspond to the standard deviation of the prior distribution of the NODE parameters, which constrains the nonlinearity of the nonparametric approximation of the NODEs. Small values of standard deviation correspond to a linear model, while higher values correspond to a highly nonlinear model. Time series are split in three thirds to create a train, validation, and test set. The model is fitted to the train set (i.e. first third) for increasing value of standard deviation (from 0.0 to 0.1 by 0.01 increments), and evaluated on the validation set. The operation is repeated by swapping the training and validation set. The graphs show the log likelihood of the NODE system fitted by BNGM to the train set (in orange), and evaluated on the corresponding validation set (in red). The shaded areas represent the 90% confidence interval on estimates, obtained by anchored ensembling of the log posterior distribution (Eq. 8) (Pearce et al. 2018).

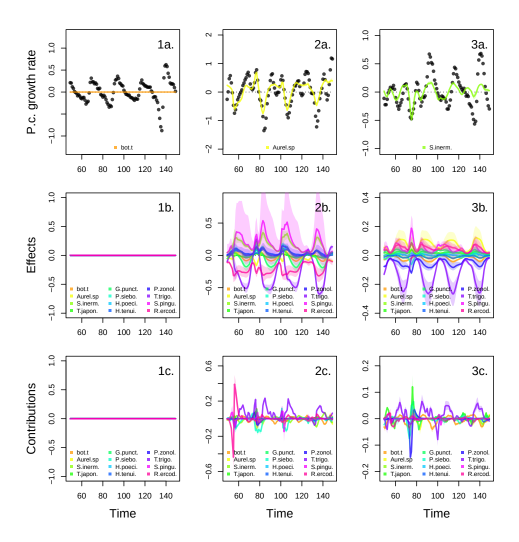


Figure S27: Drivers of dynamics of species abundance in the Maizuru bay community. This figure displays the NODE nonparametric approximations of the per-capita growth rates (2-3a.). We obtain the NODE approximations (2-3a., solid line) by fitting the interpolated per-capita growth rates (black dots) with ANNs that take population densities as input. We then estimate the direction of ecological interactions (effects, 2-3b.) by computing the derivative of the NODE approximations with respect to each density. Finally, we compute the strength of ecological interactions (contributions, 2-3c.) by multiplying the interpolated dynamics of each population with its effects. The shaded area shows the 90% confidence interval, obtained by approximately sampling the posterior distributions.

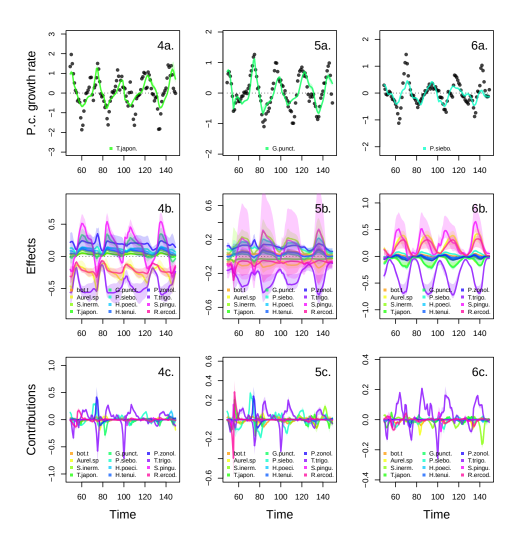


Figure S28: Drivers of dynamics of species abundance in the Maizuru bay community. This figure displays the NODE nonparametric approximations of the per-capita growth rates (4-6a.). We obtain the NODE approximations (4-6a., solid line) by fitting the interpolated per-capita growth rates (black dots) with ANNs that take population densities as input. We then estimate the direction of ecological interactions (effects, 2-6b.) by computing the derivative of the NODE approximations with respect to each density. Finally, we compute the strength of ecological interactions (contributions, 2-6c.) by multiplying the interpolated dynamics of each population with its effects. The shaded area shows the 90% confidence interval, obtained by approximately sampling the posterior distributions.

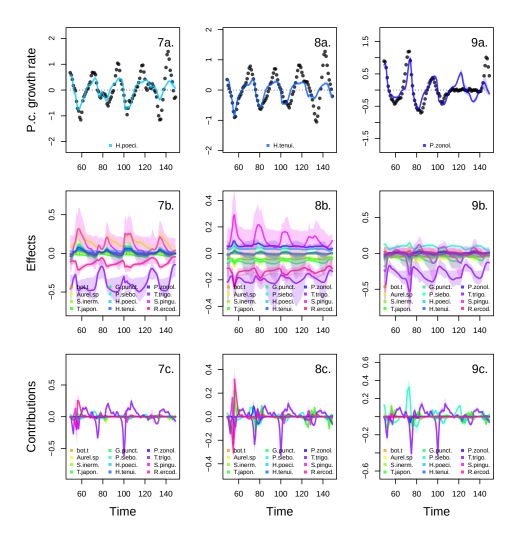


Figure S29: Drivers of dynamics of species abundance in the Maizuru bay community. This figure displays the NODE nonparametric approximations of the per-capita growth rates (7-9a.). We obtain the NODE approximations (7-9a., solid line) by fitting the interpolated per-capita growth rates (black dots) with ANNs that take population densities as input. We then estimate the direction of ecological interactions (effects, 7-9b.) by computing the derivative of the NODE approximations with respect to each density. Finally, we compute the strength of ecological interactions (contributions, 7-9c.) by multiplying the interpolated dynamics of each population with its effects. The shaded area shows the 90% confidence interval, obtained by approximately sampling the posterior distributions.

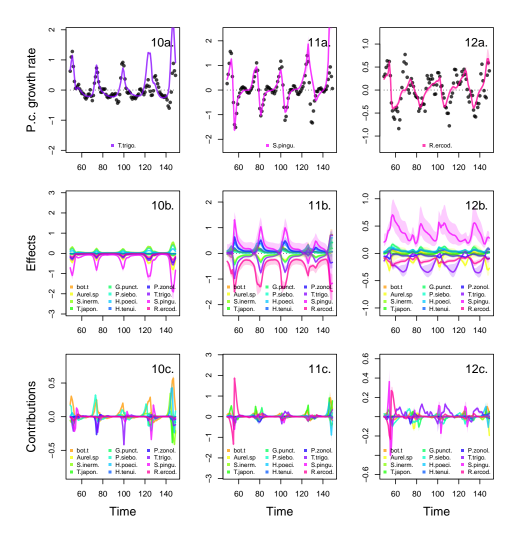


Figure S30: Drivers of dynamics of species abundance in the Maizuru bay community. This figure displays the NODE nonparametric approximations of the per-capita growth rates (10-12a.). We obtain the NODE approximations (10-12a., solid line) by fitting the interpolated per-capita growth rates (black dots) with ANNs that take population densities as input. We then estimate the direction of ecological interactions (effects, 10-12b.) by computing the derivative of the NODE approximations with respect to each density. Finally, we compute the strength of ecological interactions (contributions, 10-12c.) by multiplying the interpolated dynamics of each population with its effects. The shaded area shows the 90% confidence interval, obtained by approximately sampling the posterior distributions.