#### SPECIAL FEATURE

ECOLOGICAL WILEY

Check for updates

Promoting idea sharing via Idea Paper

# Idea paper: Predicting culturability of microbes from population dynamics under field conditions

### Masayuki Ushio<sup>1,2,3</sup>

<sup>1</sup>Hakubi Center, Kyoto University, Kyoto, Japan

<sup>2</sup>Center for Ecological Research, Kyoto University, Otsu, Shiga, Japan

<sup>3</sup>PRESTO, Japan Science and Technology Agency, Kawaguchi, Saitama, Japan

#### Correspondence

Masayuki Ushio, Hakubi Center, Kyoto University, Kyoto, 606-8501, Japan. Email: ong8181@gmail.com

#### **Funding information**

Kyoto University, Grant/Award Number: Hakubi Project; Precursory Research for Embryonic Science and Technology, Grant/Award Number: JPMJPR16O2

#### **Abstract**

Isolation and cultivation of microbes from environmental samples have been fundamental and important for species identification and investigating functions and ecology of target microbes. Although cultivation and isolation of microbes are not easy, the natural environment can "culture" any endemic microbes, and thus key information for culturing and isolating microbes may be encoded in the natural population dynamics of microbes. In this article, I present the idea that culturability of microbes may be inferred by quantifying the dynamic properties of microbes using nonlinear time series analysis and empirical dynamic modeling (EDM). To briefly demonstrate the idea, I analyzed high-frequency, quantitative microbial time series obtained for experimental rice plots. I selected bacterial phyla that included sufficient numbers of microbial taxa and analyzed 398 microbial taxa using EDM. The nine phyla analyzed generally followed a similar pattern: many microbial taxa fell into the "Simple" dynamics category, and a small proportion of taxa were categorized in "Simple but nonlinear" or "Nearly random" dynamics categories. The present analysis suggested that many microbes in the study system might be cultivated by modifying a relatively small number of conditions. However, the present idea as well as the result is preliminary, and more precise taxonomic information (i.e., species-level identification) and a culturability data set will help to validate the idea. If the present idea was found to be valid, a priori evaluation of the culturability of microbes would become possible, which would avoid unnecessary costs (labor, time and money) of attempts to cultivate microbes.

#### KEYWORDS

culturability, DNA, empirical dynamic modeling, microbes, time series

### 1 | RESEARCH QUESTION

Isolation and cultivation of microbial species from environmental samples have been fundamental for identifying microbial species and investigating functions and ecology of a target microbial species. However, unfortunately, the majority of microbes in the environment

cannot be cultured easily (e.g., Amann, Ludwig, & Schleifer, 1995), and improving the recovery of microbes from environmental samples is an important but difficult and labor-, time- and money-consuming work. Therefore, if a priori evaluations of the culturability of microbes would become possible, it would contribute to avoiding unnecessary costs and improving the recovery of

microbes from environments. For example, a cultivation design that includes a large number of combinations of nutrients in media might not be necessary if the target microbes were classified as "easily cultured."

#### 2 | VALUE

A priori evaluations of the culturability of microbes would contribute to reducing potential costs of cultivation. Also, a framework for quantifying culturability would enable analysis of the relationship between the culturability and genetic factors, which could potentially contribute to understanding why particular microbial species are difficult to culture while others are not.

## 3 | RELEVANT HYPOTHESIS (AND APPROACHES)

Although isolation and cultivation of microbes are not easy, natural environments can "culture" any endemic microbes, and thus key information for isolating and culturing microbes may be encoded in microbes' interactions with natural habitats and resultant population dynamics in nature (or under field conditions). Indeed, the diffusion chamber approach, which allows exchanges of chemicals between media and their natural habitat, can simulate the natural environment and can improve the recovery of some microbes from environment (Bollmann, Lewis, & Epstein, 2007; Kaeberlein, Lewis, & Epstein, 2002). These findings implied that microbes in nature respond to biotic/abiotic variables in their environment and the interactions among microbes and the environment drive their population dynamics. Therefore, conversely, the natural population dynamics of microbes contain integrated information on biotic and abiotic variables and their interactions in the environment. The question is, however, how we can extract the information encoded in the population dynamics of microbes.

Information encoded in population dynamics can be extracted using empirical dynamic modeling (EDM) that is based on attractor reconstruction and designed for analyzing nonlinear dynamics (e.g., Sugihara, May, Ye, et al., 2012). Core tools of EDM are the simplex projection (Sugihara & May, 1990) and S-map (Sugihara, 1994), which enable quantifying (a) the best embedding dimension and (b) state dependence of time series data. Briefly, the best embedding dimension (denoted by E) includes information on how many potential variables might be involved in the process (i.e., complexity or dimensionality), and state dependency (quantified by the nonlinear weighting parameter,  $\theta$ ) includes information on how

state dependent the process is. A previous study demonstrated that E and  $\theta$  are effective indices to distinguish random environmental fluctuations from low dimensional, nonlinear dynamics of organisms (Hsieh, Glaser, Lucas, & Sugihara, 2005). Detailed information on how simplex projection and S-map are performed is described in previous studies (Sugihara, 1994; Sugihara & May, 1990). Also, an overview of EDM is described in Chang, Ushio, and Hsieh (2017).

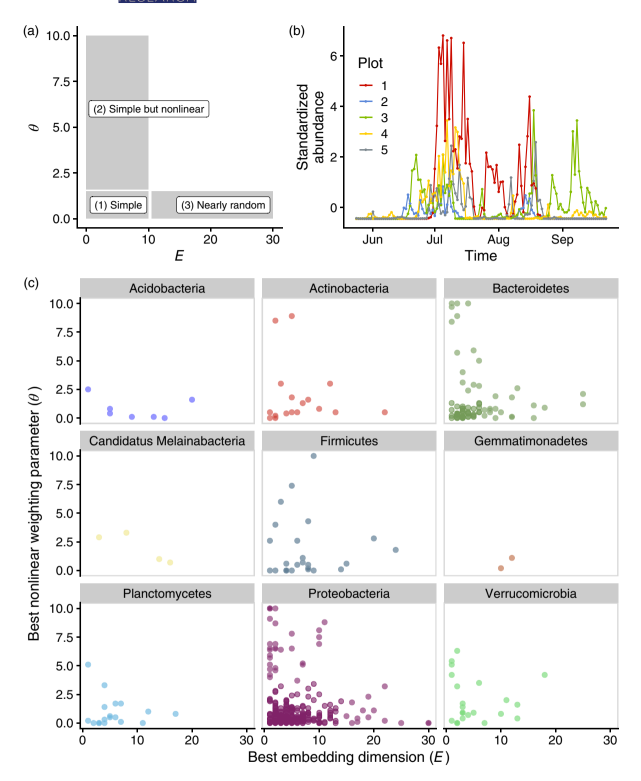
#### 4 | NEW RESEARCH IDEA

In this article, I present the idea that culturability of microbes may be inferred using population dynamics of microbes and EDM. Specifically, I expect that the complexity (E) and state dependency  $(\theta)$  of population dynamics would provide information on the culturability of microbes.

# 5 | HOW TO ANSWER THE QUESTION THROUGH THE RESEARCH APPROACH

I expect three possible dynamics regarding the combinations of E and  $\theta$  by following Hsieh et al. (2005): (a) "Simple" (i.e., low-dimensional dynamics), (b) "Simple but nonlinear" and (c) "Nearly random" dynamics (Figure 1a). First, if the population dynamics of a target species show small E and  $\theta$ , then the number of potential variables might be small and the species responds to cultivation conditions relatively linearly ("Simple" dynamics). In this case, the target microbe might be cultivated by examining a relatively small number of cultivation conditions. Second, if the population dynamics show small E but large  $\theta$ , then the number of potential variables might be small but the growth of the species is state dependent ("Simple but nonlinear" dynamics). In this case, the target microbe could respond to cultivation conditions nonlinearly, and thus careful considerations of cultivation conditions would be required. Third, if the population dynamics show large E but small  $\theta$ , then the dynamics are nearly random and the target microbe might have randomly immigrated from outside the system ("Nearly random" dynamics).

To briefly demonstrate the idea, I analyzed highly frequent, quantitative microbial time series taken from experimental rice plots established at the Center for Ecological Research, Kyoto University, Japan (Ushio, unpublished). Briefly, water samples were collected from five rice plots every day from 23 May to 22 September in 2017 (610 samples in total) and filtered using filter



**FIGURE 1** (a) A conceptual explanation of the relationships between E and  $\theta$  and dynamics properties. Note that the thresholds between the categories are arbitrarily determined. (b) An example of time series analyzed in the present study. Changes in abundance (i.e., estimated DNA copy numbers) of a microbial amplicon sequence variant (ASV) belonging to Bacteroides are shown. Different colors indicate different rice plots where water samples were taken. (c) The relationships between E and  $\theta$  and microbial taxa. Each panel indicates a particular phylum, and each point indicates the best E and  $\theta$  of each ASV [Color figure can be viewed at wileyonlinelibrary.com]

cartridges (pore size =  $\varphi$ 0.22 µm). Then, DNAs were extracted, amplified a portion of the 16S rRNA genes and sequenced using Illumina MiSeq. In the library preparation process, artificial standard DNAs with known concentrations were included to estimate the copy numbers of microbial DNAs (Ushio, 2019). The sequences generated were analyzed using the amplicon sequence variant (ASV) approach (Callahan, McMurdie, Rosen, et al., 2016). I selected bacterial phyla that are abundant and contain sufficient numbers of ASVs, which resulted in 398 bacterial ASVs belonging to nine phyla. An example of the microbial time series is shown in Figure 1b. For the time series selected, simplex projection and S-map were applied to determine the best E and  $\theta$  using rEDM packages (Ye, Clark, Deyle, et al., 2018) of R (R Core Team, 2019). The whole data set is being analyzed for different purposes and thus is currently not publicly available.

The nine phyla analyzed followed a similar pattern (Figure 1c): many bacterial ASVs fall into the "Simple" category, and a relatively small proportion of ASVs were categorized as "Simple but nonlinear" or "Nearly random." This result might indicate that, among the selected bacterial ASVs, many microbes could be cultivated by modifying a relatively small number of conditions, which is contrary to the current general consensus that most environmental microbes are difficult to cultivate and isolate (Amann et al., 1995; but see Martiny, 2019). However, technical issues (e.g., biases generated by an amplicon sequencing approach) may partly contribute to this pattern, and thus careful interpretations of the results are necessary (see debates in the following references: Martiny, 2019; Steen, Crits-Christoph, Carini, et al., 2019).

Here I list data that are required to verify my idea more thoroughly. First, microbes should be identified at least at the species level. This is because the culturability would be a property of a species, and thus genus- or family-level identification is insufficient. The present data analyzed were generated by a short-read marker-gene (amplicon) analysis, and thus the phylogenetic resolution was often not sufficient. Recently introduced long-read sequencing (e.g., full-length 16S rRNA sequencing) would be more appropriate to achieve this goal (e.g., MinION by Oxford Nanopore or Sequel by PacBio). Species-level dynamic properties would be useful to classify and predict previously ignored aspects of microbial properties. Second, accurate information on which microbial species and how the microbial species are cultivated is required. Information on which microbial species is cultivated would be available in public databases, but developing the cultivation method often requires detailed tuning and thus the method as well as its complexity would be difficult to fully document/digitalize. Compiling fundamental information about cultivation methods (e.g., temperature

resources) would help to develop a culturability database, which could also contribute to understanding of the potential habitat range of microbes in nature. Overall, if the present idea were validated by overcoming the above limitations, dynamic properties of microbes could be linked to culturability and a priori evaluation of the culturability of microbes would become possible.

#### **ACKNOWLEDGMENTS**

I would like to thank Asako Kawai for assistance in the sampling and experiments and Hiroki Yamanaka for providing an opportunity to use Illumina MiSeq. This study was financially supported by PRESTO (JPMJPR16O2) from the Japan Science and Technology Agency (JST) and the Hakubi project of Kyoto University.

#### CONFLICT OF INTEREST

The author declare no conflict of interest.

#### ORCID

Masayuki Ushio https://orcid.org/0000-0003-4831-7181

#### REFERENCES

- Amann, R. I., Ludwig, W., & Schleifer, K. H. (1995). Phylogenetic identification and in situ detection of individual microbial cells without cultivation. *Microbiological Reviews*, *59*, 143–169.
- Bollmann, A., Lewis, K., & Epstein, S. S. (2007). Incubation of environmental samples in a diffusion chamber increases the diversity of recovered isolates. *Applied and Environmental Microbiology*, 73, 6386–6390.
- Callahan, B. J., McMurdie, P. J., Rosen, M. J., Han, A. W., Johnson, A. J. A., & Holmes, S. P. (2016). DADA2: High-resolution sample inference from Illumina amplicon data. *Nature Methods*, 13, 581–583
- Chang, C.-W., Ushio, M., & Hsieh, C. (2017). Empirical dynamic modeling for beginners. *Ecological Research*, 32, 785–796.
- Hsieh, C., Glaser, S. M., Lucas, A. J., & Sugihara, G. (2005). Distinguishing random environmental fluctuations from ecological catastrophes for the North Pacific Ocean. *Nature*, 435, 336–340.
- Kaeberlein, T., Lewis, K., & Epstein, S. S. (2002). Isolating uncultivable; microorganisms in pure culture in a simulated natural environment. *Science*, 296, 1127–1129.
- Martiny, A. C. (2019). High proportions of bacteria are culturable across major biomes. *The ISME Journal*, *13*, 2125–2128.
- R Core Team. (2019). R: A language and environment for statistical computing. Vienna: Austria.
- Steen, A. D., Crits-Christoph, A., Carini, P., DeAngelis, K. M., Fierer, N., Lloyd, K. G., & Cameron Thrash, J. (2019). High proportions of bacteria and archaea across most biomes remain uncultured. *The ISME Journal*, *13*, 3126–3130.
- Sugihara, G. (1994). Nonlinear forecasting for the classification of natural time series. *Philosophical Transactions of the Royal Society A: Mathematical, Physical and Engineering Sciences, 348,* 477–495.
- Sugihara, G., May, R., Ye, H., Hsieh, C., Deyle, E., Fogarty, M., & Munch, S. (2012). Detecting causality in complex ecosystems. *Science*, *338*, 496–500.

- Sugihara, G., & May, R. M. (1990). Nonlinear forecasting as a way of distinguishing chaos from measurement error in time series. *Nature*, *344*, 734–741.
- Ushio, M. (2019). Use of a filter cartridge combined with intracartridge bead-beating improves detection of microbial DNA from water samples. *Methods in Ecology and Evolution*, 10, 1142–1156.
- Ye H., Clark A., Deyle E., Munch, S., Keyes, O., Cai, J., ... Sugihara, G. (2018). rEDM: Applications of empirical dynamic modeling from time series. doi:https://doi.org/10.5281/zenodo.1935847

**How to cite this article:** Ushio M. Idea paper: Predicting culturability of microbes from population dynamics under field conditions. *Ecological Research.* 2020;35:586–590. <a href="https://doi.org/10.1111/1440-1703.12104">https://doi.org/10.1111/1440-1703.12104</a>