

Mesocosm Examples

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Abstract

The **QPress** package provides facilities for modelling the qualitative impact of a press perturbation upon a network model. This document illustrates the comparison of alternative models in terms of their responses to perturbation. Two examples are provided for experimental mesocosms in (I) a lake environment (Hulot et al., 2000), and (II) Antarctica (Davidson et al. unpublished).

1 Introduction

An experimental mesocosm is an ecological tool that enables the experimenter to assess assemblage-level responses to environmental change. Mesocosm experiments are performed in the ‘natural environment’ rather than in a laboratory, but in an enclosure that is small enough that key variables can be directly controlled. Because the responses of component species to perturbation can be observed directly in a mesocosm setting, the results from such studies lend themselves to analysis using qualitative network modelling. In particular, these analyses can be used to make inferences regarding the trophic and competitive interactions that drive assemblage-level responses to perturbation in a mesocosm setting and potentially more broadly.

In this document we use the **QPress** package to analyse two sets of models corresponding to two separate mesocosm experiments. Example I is for experimental mesocosms in a lake in France that were subjected to high and low nutrient treatments (Hulot et al., 2000). Example II examines the effects of elevated concentrations of CO₂ on the Southern Ocean microbial loop, studied in environmentally controlled tanks (referred to as ‘minicosms’ by the authors) in Antarctica (Davidson et al. unpublished).

2 Example I: Lake mesocosm

Hulot et al. (2000) describe an eight-variable lake mesocosm model comprising phosphorous, three

algal groups, two size-classes of herbivores, invertebrate carnivores and carnivorous fish (Figure~1). Unambiguous, directional responses of model components to increased phosphorous observed in the mesocosm experiment were increases in each of: phosphorous (Phos), large herbivores (Herb2), fish (Carn2) and periphyton (AlgP). Melbourne-Thomas et al. (2012) compare the ability of Hulot et al.'s original model (Figure~1a) and a variant of the model with one edge added and one edge deleted (Figure~1b) to reproduce these changes in response to a positive perturbation to phosphorous. Specifically, the variant of the original model (model (b) in Figure~1b) includes a self-limitation effect for invertebrate carnivores, and the predator-prey edge from fish to invertebrate carnivores has been deleted.

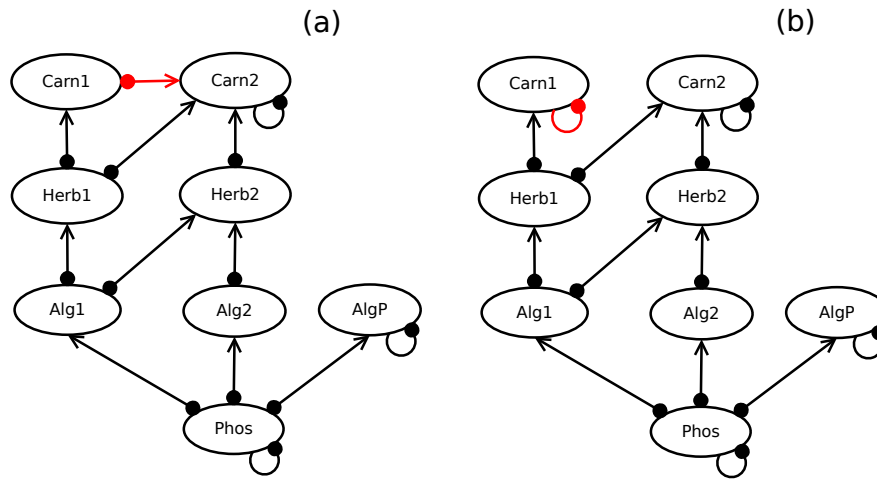


Figure 1: Two versions of the eight-variable lake mesocosm model analyzed by Melbourne-Thomas et al. (2012) and Hosack et al. (2008), and originally presented by Hulot et al. (2000). Abbreviations are: Alg1, edible algae; Alg2, protected algae; AlgP, periphyton; Carn1, invertebrate carnivores; Carn2, fish; Herb1, small herbivores; Herb2, large herbivores; Phos, phosphorous.

The models shown in Figure~1 were created using Dia and so are read using `model.dia`. Here we assign the two model objects to `lake.a` and `lake.b`.

```
> library(QPress)
> lake.a <- model.dia("Lake-a.dia")
> lake.b <- model.dia("Lake-b.dia")
```

We then simulate for model (a) and use `impact.barplot` to view simulation outcomes corresponding with observations from the mesocosm experiment (Figure~2).

```
> simlake.a <- system.simulate(1000, lake.a)
> impact.barplot(simlake.a)
```

Doing the same for model (b) gives Figure~3.

Approximately 50% of the simulations match with observations for model (a), whereas for model (b) this increases to over 80%. This suggests that model (b) provides a more parsimonious representation of the lake mesocosm system. Melbourne-Thomas et al. (2012) provide a direct comparison of the two models using Bayes factors.

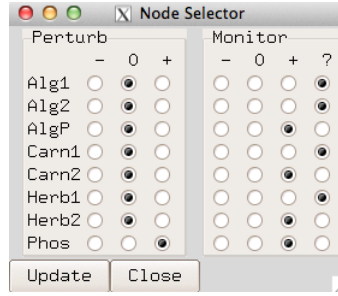


Figure 2: Node selector for the lake mesocosm example.

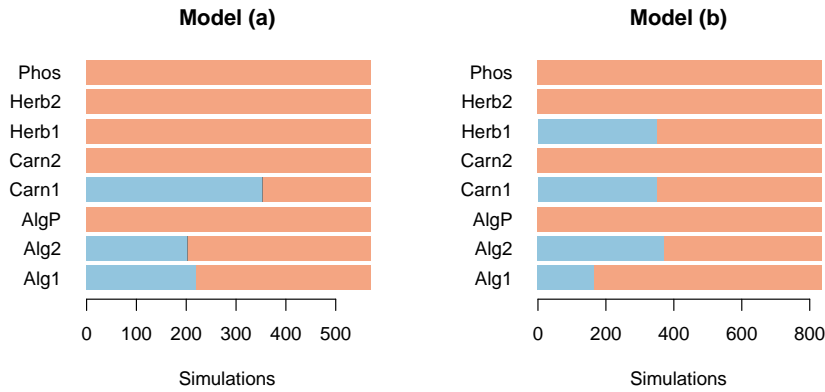


Figure 3: Simulation outcomes from the two lake mesocosm models in response to a positive press perturbation to phosphorous. Each plot shows the fraction of positive (orange), and negative (blue) outcomes at each node. The x-axis indicates the number of simulations (out of 1000 total) that match with the selection for ‘Monitor’ shown in Figure 2.

3 Example II: Antarctic mesocosm

This example is based on experiments conducted at Australia’s Davis Station in Antarctica, which assessed the responses of a natural community of Antarctic marine microbes from near-shore waters to elevated concentrations of CO₂ using environmentally controlled tanks (mesocosms). Davidson et al. (in prep) describe the main components of the Southern Ocean microbial loop represented in these mesocosms and the ecological interactions between them. Specifically:

- Small and large phytoplankton cells consume macronutrients and iron (Fe), although iron is potentially less important for growth of small phytoplankton;
- Small phytoplankton are consumed by heterotrophic nanoflagellates and microzooplankton;
- Large phytoplankton (diatoms) are consumed by microzooplankton;
- Bacteria break down DOC (dissolved organic carbon) and are consumed by heterotrophic nanoflagellates;
- Phytoplankton, microzooplankton and heterotrophic nanoflagellates supply the DOC pool.

These interactions are represented in a model that was built using Dia (black-coloured edges in Figure 4).

When this system was subjected to elevated CO₂ concentrations in experimental mesocosms, the following responses were observed:

- Small phytoplankton increased;
- Large phytoplankton decreased;
- Heterotrophic nanoflagellates decreased; and
- Bacteria increased.

In the model shown in Figure~4 we assume that the direct effects of CO_2 correspond with these observed changes.

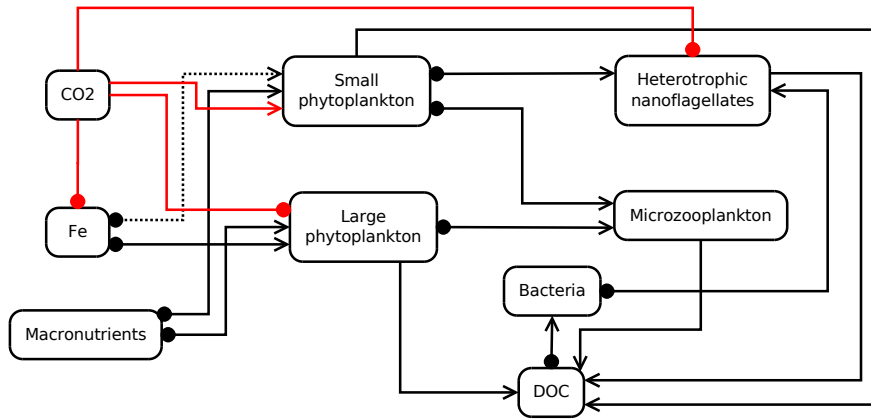


Figure 4: Model (a) representing the effects of CO_2 on the Southern Ocean microbial loop, determined directly from observed responses in experimental mesocosms. These direct effects are shown in red.

We then analyse the response this model to a positive press perturbation to CO_2 . In this case we enforce a specific ordering of the nodes, and also enforce self-limitation for each model variable.

```
> ## Enforce the ordering of nodes.
> labels <- c("CO2","Fe","Macronutrients","Small phytoplankton",
+           "Large phytoplankton","Heterotrophic nanoflagellates",
+           "Microzooplankton","Bacteria","DOC")
> antarctic.a <- model.dia("Antarctic-a.dia",labels=labels)
> antarctic.a <- enforce.limitation(antarctic.a)
```

As for Example I, we then simulate and use `impact.barplot` (with the selections shown in Figure~5) to produce Figure~6.

```
> simantarctic.a <- system.simulate(1000,antarctic.a)
> impact.barplot(simantarctic.a)
```

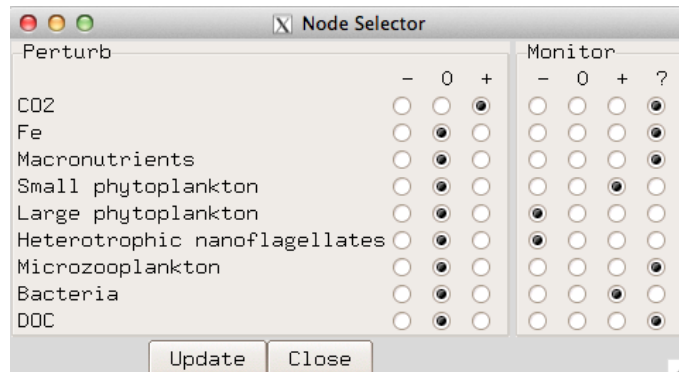


Figure 5: Node selector for the Antarctic mesocosm example.

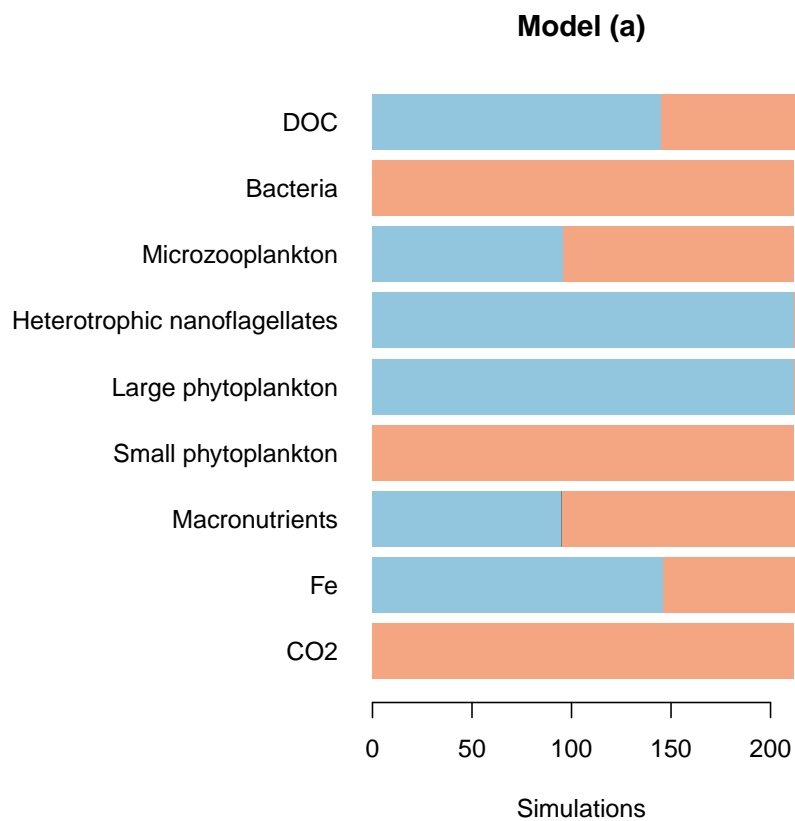


Figure 6: Simulation outcomes in response to a positive press perturbation to CO₂ for Antarctic mesocosm model (a). Each plot shows the fraction of positive (orange), and negative (blue) outcomes at each node.

For this version of the model, less than 20% of the simulations match with our ‘Monitor’ criteria, suggesting that it does not provide a particularly good representation of the Antarctic microbial loop system or its response to increased CO₂ concentrations. Given that increases in dissolved CO₂ are generally expected to have negative impacts on the growth of marine organisms, we could assume a direct negative effect of CO₂ on each of large phytoplankton, small phytoplankton, heterotrophic nanoflagellates, and also the pool of available iron (Figure~7).

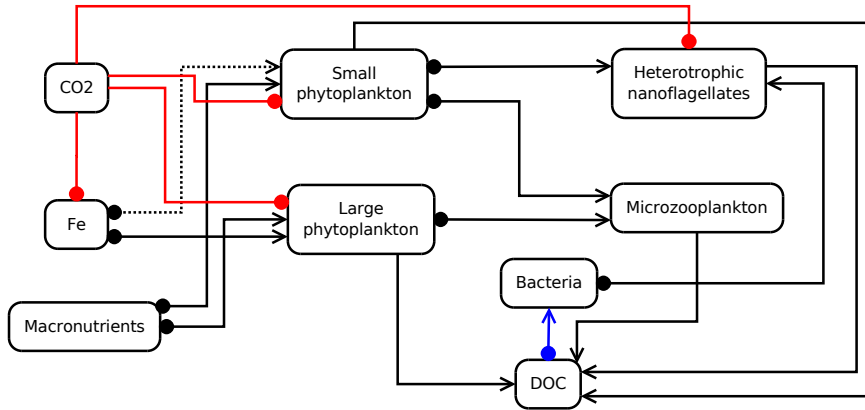


Figure 7: Alternative model (b) representing the effects of CO₂ on the Southern Ocean microbial loop.

Reading this alternative representation from a Dia file

```
> antarctic.b <- model.dia("Antarctic-b.dia", labels=labels)
> antarctic.b <- enforce.limitation(antarctic.b)
```

and exploring simulation outcomes with `impact.barplot` (with the same ‘Monitor’ criteria as before) produces Figure~8.

```
> simantarctic.b <- system.simulate(1000, antarctic.b)
> impact.barplot(simantarctic.b)
```

In this case, we still only see approximately 20% of simulations that meet our criteria, suggesting that we need to re-think the structure of our model. Davidson et al. (in prep) indicate that bacteria “transform dissolved organic carbon (DOC)... thereby repackaging carbon and making it available to higher trophic levels”. Looking at our original model, we haven’t quite captured this in our representation of interactions between bacteria, DOC and macronutrients. In a third version of our microbial loop model (model (c) – Figure~9) we therefore assume a negative effect of bacteria on DOC and a positive effect of bacteria on macronutrients (which includes carbon).

Reading a third Dia version of the model

```
> antarctic.c <- model.dia("Antarctic-c.dia", labels=labels)
> antarctic.c <- enforce.limitation(antarctic.c)
```

and repeating the simulation and `impact.barplot` steps from above gives Figure~10.

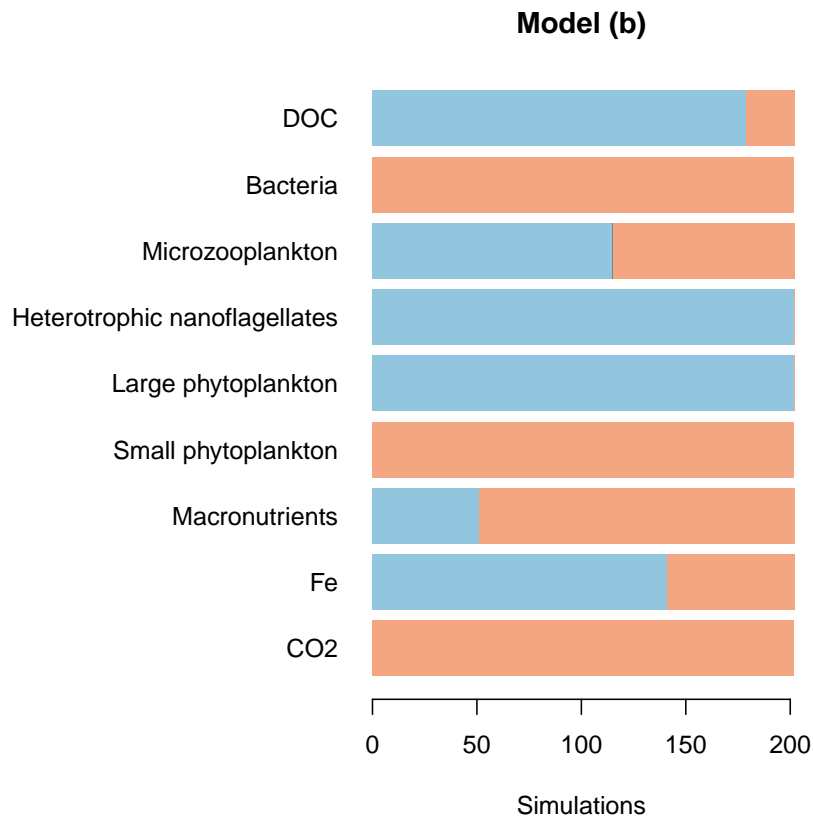


Figure 8: Simulation outcomes in response to a positive press perturbation to the CO_2 for the alternative microbial loop model (b) shown in Figure~7.

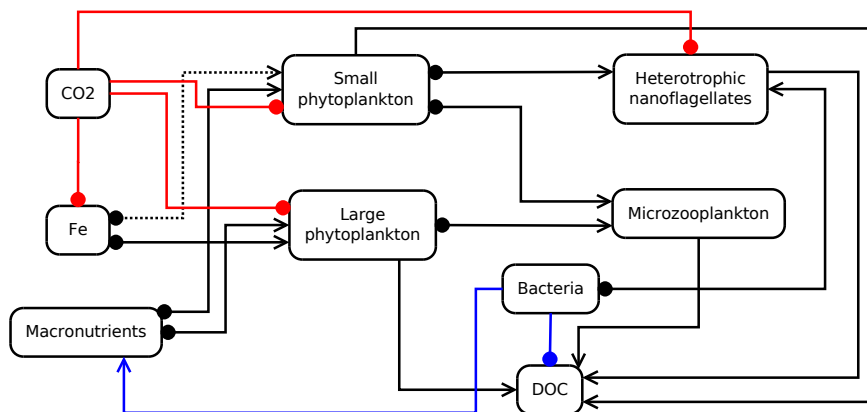


Figure 9: Alternative model (c) representing the effects of CO_2 on the Southern Ocean microbial loop.

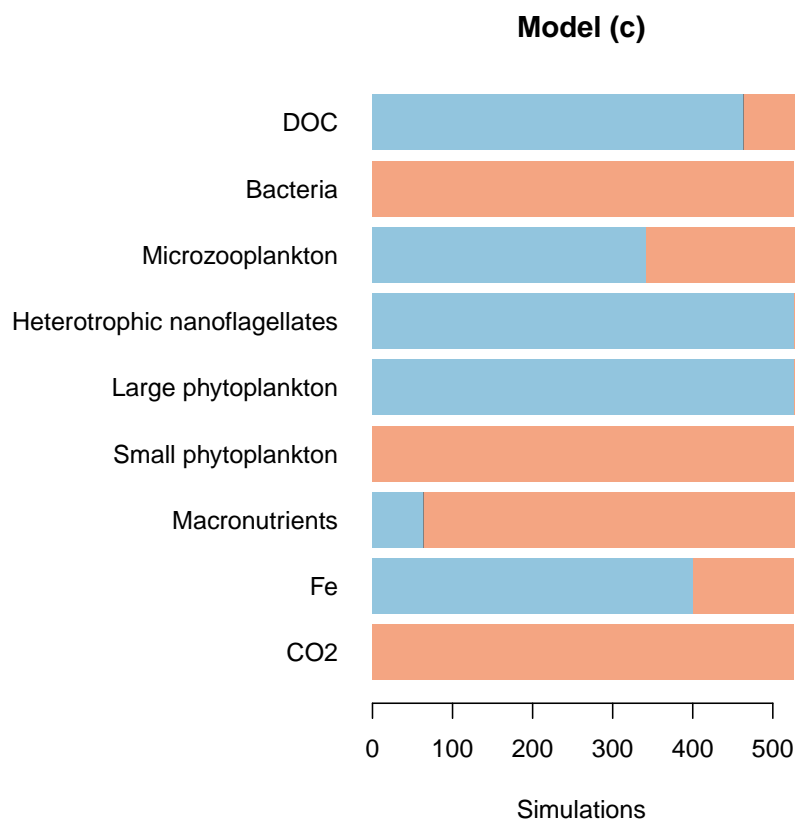


Figure 10: Simulation outcomes in response to a positive press perturbation to the CO_2 for the alternative microbial loop model (c) shown in Figure~9.

References

- Hosack, G.~R., Hayes, K.~R., and Dambacher, J.~M. (2008). Assessing model structure uncertainty through an analysis of system feedback and Bayesian networks. *Ecological Applications*, 18(4):1070–1082.
- Hulot, F.~D., Lacroix, G., Lescher-Moutoué, F., and Loreau, M. (2000). Functional diversity governs ecosystem response to nutrient enrichment. *Nature*, 405:340–344.
- Melbourne-Thomas, J., Wotherspoon, S., Raymond, B., and Constable, A. (2012). Comprehensive evaluation of model uncertainty in qualitative network analyses. *Ecological Monographs*, 82(4):505–519.