Interactive comment on “Response of bacterioplankton activity in an Arctic fjord system to elevated $p$CO$_2$: results from a mesocosm perturbation study” by J. Piontek et al.

Anonymous Referee #1

Received and published: 24 September 2012

General comment

This study reports temporal variations of bacterial protein production and activity of three extracellular enzymes in the Kongsfjorden, Svalbard. This study was conducted in the frame of a mesocosm pCO2 perturbation experiment using a natural plankton community. The authors demonstrate that (1) bacterial community was initially limited by the availability of organic carbon, (2) activity of beta-glucosidase and leucine-aminopeptidase increased with decreasing seawater pH, (3) Q10 for beta-glucosidase, leucine-aminopeptidase and bacterial protein production decreased with increasing carbon exudation rate. They give two important suggestions: (1) future changes in seawater temperature and pH have a similar potential to increase extracellular enzyme activity, (2) future pH decrease have a potential to enhance consumption of labile organic matter by bacteria.

I think that the presentation of the 'fjord' data (outside the mesocosms) does not fit to the scope of this ms. Hence, it appears to me that the current ms is unfocused. Once the mesocosm experiment starts, the plankton community inside and outside the mesocosms are under different conditions. Provided the scope of this study and of the EPOCA experiment, I had the difficulty to understand the importance of (for example) the comparison of bacterial production between the mesocosms and in situ on day 20 in this ms. I recommend to reduce the presentation of in situ data.

Q10 of bacterial protein production (BPP): I think that the water temperature at the in situ incubation site was not constant during each incubation. If the incubation temperature varied during 24 h incubation for BPP, the authors should mention the range of temperature variation and explain how the daily change of temperature was taken into account for the calculation of Q10 of BPP. The authors conclude the significant effect of water temperature on BPP, however they present the data of BPP at 2°C (not at in situ temperature) for analyzing the effect of pCO2 on BPP (see Fig. 2). They need to give rationale for this.

Other comments

Abstract P 10468, L9-10: It appears that this sentence means that the extracellular enzyme was highest at moderate acidification level in this experiment. But the results suggest that the extracellular enzyme was highest at the higher pCO levels (Fig. 6).

Introduction P10469, L4: Iversen and Seuthe -> Rokkan Iversen and Seuthe

P10469, L4-6: Neither Rokkan Iversen and Seuthe (2011) nor Seuthe et al. (2011) report that bacteria were subject to intense grazing by heterotrophic dinoflagellates and ciliates during the vernal bloom in April. Please specify the reference.
P10470, L10: It is not clear which period “past marine research” means. However, it should be noted that, as the authors cited in the ms, for example Liu et al. (2010) did meta-analysis of the published papers about the effect of ocean acidification on the structure and functioning of microbial communities.

P10471, L15-16: This sentence says that pCO2 in the enclosed seawater was initially in a range of 250-1085 µatm. But in fact the authors set up a pCO2 gradient in a range of 250-1085 µatm by the stepwise addition of CO2-enriched seawater during several days.

P10471, L20-21: Specify if whole plankton community or certain groups developed, and what the nutrient deplete condition mean here (e.g. concentration, type of nutrients). Add relevant reference.

P10372, L17-18: It would be useful for readers to mention how different pH levels affected the calibration factor of MUF and AMC fluorescence. It is suggested to mention how the authors measured blank fluorescence.

P10472, L25-26: Enzymatic rates were given as nmol/l in the ms, so that the unit of mean and standard deviation of rates should be nmol/l. “a standard deviation of 9%” should be rewritten.

P10473, L4-5: Specify the number of live samples and killed-control samples.

P10473, L11-12: Give the information of the light condition in a temperature-controlled walk-in room and in situ during the incubation.

P10474, L23: Specify if “20 µmol/l” is glucose concentration or carbon concentration.

P10475, L3: It is unclear why the authors applied the diluted acid instead of CO2 gas for pH adjustment in the acidification assay. This is apparently different from the pH modification in the mesocosm experiment.

P10476, L4-7: It is unclear why delta-hydrolysis potential was calculated by the difference in enzyme activity between the acidified mesocosms and the two control mesocosms (M3 and M7) on each day. The two mesocosms (M3 and M7) were control in terms of CO2 manipulation but received nutrient enrichment on day 13. The response of plankton community may be different even between the control mesocosms during one month incubation. In this context, it would be better to show the integrated values of enzyme activity in the two control mesocosms as well.

Results P10476, Enrichment assays: The limiting resource should be identified based on statistical analysis. The current ms shows the statistical analysis only on Lines4-5, Page 10477. The result of the statistical test should be shown in the text and Figure 1.

P10478, L7-8: Figs. 4-5 do not support the description ‘activities remained rather constant between days 12 and 20’.

P10478, L16-17: The method used in this study measured not only ‘bacterial enzymes released into seawater’ and also bacterial enzymes in particulate fraction.

P10478, L19-21: Fig. 6 shows that an elevated enzymatic potential was observed in the two highest pCO2 mesocosms rather than “the three mesocosms of highest pCO2” during the first 20 days.

P10479, L14-17: It seems that the data on DOC and DON are (or will be) published in an accompanying paper. If yes, add the reference. The same for Fig. 7 legend.

P10479, L18-21: It is very difficult to understand that a substrate concentration of 200 µmol/l did not saturate alkaline phosphatase activity, so that the data are shown based on the measurement of alkaline phosphatase activity at a non-saturation substrate concentration of 10 µmol/l. Isn’t it 100 µmol/l?
The production of alkaline phosphatase is generally enhanced under low concentration of phosphate. It would be interesting to compare the relationship between alkaline phosphatase activity and phosphate concentration.

Add reference.

Specify if no significant differences of water temperature, Q10 for extracellular enzyme, and BPP between the mesocosms were tested by statistical analysis.

’revailing’ -> ‘revealing’?

‘bPP’ -> ‘BPP’?

Add ‘Fig. 11’ at the end of the sentence.

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may assume the bulk organic carbon channelled mesozooplankton and then exported from the surface.

Fig. 1: The upper panel shows BPP and bacterial abundance. The legend should mention bacterial abundance as well.

Fig. 2: The lower panel indicates specific growth rate (/d) on the right axis. The legend should mention specific growth rate as well.

Fig. 11: The unit of bacterial protein production (BPP) is fgC/cell which does not correspond to the first definition. In the legend, primary production should be time-integrated primary production. The integration period should be added.

Fig.13 is not referred in the text. I do not think that this figure is useful for the ms.

Interactive comment on Biogeosciences Discuss., 9, 10467, 2012.