**Biological or Microbial Carbon Pump? The role of phytoplankton stoichiometry in**

**ocean carbon sequestration**

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**Abstract**

Once fixed by photosynthesis carbon becomes part of the marine food web. The fate of this carbon has two possible outcomes: it may be respired and released back to the ocean and potentially the atmosphere as CO2 or retained in the ocean interior and/or marine sediments for extended time scales. The most important biologically mediated processes responsible for long term carbon storage in the ocean are the biological carbon pump (BCP) and the microbial carbon pump (MCP). While acting simultaneously in the ocean, the balance between these two mechanisms is thought to vary depending on the trophic state of the environment. Using previously published formulations, we propose a modelling framework to simulate variability in the MCP: BCP ratio as a function of external nutrients. Our results suggest that the role of the MCP might become more significant under future climate change conditions where increased stratification enhances the oligotrophic nature of the surface ocean. Based on these model results, we propose a conceptual framework in which the internal stoichiometry of phytoplankton, modulating both grazing pressure and DOM production (via phytoplankton exudation), plays a crucial role in regulating the MCP: BCP ratio.

**Introduction**

Assessing the capacity of the ocean to sequester atmospheric CO2 is one of the key challenges in oceanography. Consequently, a large number of studies have been dedicated to the physical and biological mechanisms capable of transporting carbon from the surface ocean to its interior (Volk and Hofert, 1985; Sarmiento and Gruber, 2006 and references therein). Amongst the biological processes, the so-called Biological Carbon Pump (BCP) is the most widely recognized and extensively studied (Honjo *et al*., 2008 and 2014; Henson *et al*., 2012; Lutz *et al*., 2007; Passow and Carlson 2012). However, the more recently defined Microbial Carbon Pump (MCP, Jiao *et al*., 2010) provides an additional biological mechanism which may significantly contribute to the capacity of the ocean to store atmospheric CO2.

Both the MCP and BCP facilitate the long term storage of carbon in the ocean, however, they are conceptually different. The BCP mainly describes the fixation of CO2 into particulate organic carbon (POC) and its subsequent transfer to the deep ocean through gravitational sinking. Conversely, the MCP does not imply any physical transportation but is instead based upon a bacterially mediated chemical transformation of dissolved organic matter (DOM) from labile (i.e., rapidly degradable) to recalcitrant forms (i.e., slowly degradable). Current estimates suggest that the carbon stored over long time scales (i.e., millennia) through the MCP is ~0.4% of the annual oceanic primary production (Legendre *et al*., 2015). The amount of carbon exported by the BCP over 2000 m depth ranges from 0.6 to 1.3% of the annual primary production (Legendre *et al*., 2015).

While the MCP and BCP occur simultaneously in the ocean, the balance between the two is thought to vary both spatially and temporally depending on the trophic conditions of the environment (Jiao *et al*, 2010). BCP is thought to dominate in conditions where nutrient sufficiency favours large plankton groups (e.g. diatoms) resulting in large, fast-sinking detritus (marine snow and/or fecal pellets). Conversely, oligotrophic regions dominated by small phytoplankton species (Jardillier *et al*., 2013) will support the microbial loop and fuel the MCP (Legendre *et al*., 2015). Nutrient deficiency is expected to enhance the MCP by substantially altering (“unbalancing”) phytoplankton stoichiometry and producing DOM with high C:N and C:P ratios. Additionally, stoichiometrically unbalanced phytoplankton is poor quality food for zooplankton (Urabe and Sterner 1996; Siuda and Dam, 2010; Loladze et al., 2000; Jones and Flynn 2005)*.* This in turn decreases zooplankton assimilation efficiency resulting in the release of carbon-rich detritus resistant to bacterial degradation (Jiao *et al*., 2014; Mitra, 2006; Polimene *et al*., 2015). Both the geographical distribution and intensity of nutrient limitation in the global ocean is projected to increase as a consequence of climate change induced increases in thermal stratification (Polovina *et al*., 2008); under this scenario a strengthening role for the MCP has been hypothesized for the future oceans (Jiao *et al*., 2010).

Although the MCP has been conceptually linked to nutrient supply and food quality for zooplankton (Jiao *et al*., 2014), a coherent modelling framework to test the effect of nutrients on the balance between MCP and BCP is still lacking. In this paper, we present a modelling framework able to simulate the partition of carbon between the BCP and the MCP as a function of nutrient concentrations. To achieve this, we combined the model of Polimene *et al.* (Polimene *et al.,* 2015) describing the effect of phytoplankton stoichiometry on grazing (a process called Stoichiometric Modulation of Predation, SMP, Mitra 2006) with a model describing the interaction between heterotrophic bacteria and DOM, including the production of recalcitrant DOC (Polimene *et al*., 2006). The two models (Polimene *et al.,* 2015 and 2006) were incorporated within the European Regional Seas Ecosystem Model (ERSEM, Butenschon *et al*., 2016) and share the same currencies (C, N, P) and underlying assumptions. Based on modelling results, we propose a conceptual framework in which the stoichiometry of primary producers is the central factor regulating the magnitude of the MCP relative to the BCP.

**The Model**

The combined model is composed of five biotic components (Fig. 1): one phytoplankton functional type, three zooplankton functional types (microzooplankton, mesozooplankton and heterotrophic nanoflagellates) and one heterotrophic bacteria functional type. The abiotic state variables include nutrients (ammonium, nitrate, phosphate and silicate), DOM (labile, semi-labile, and semi-refractory) and POM. The POM is further differentiated into three categories according to size/sinking velocity: i) small detritus, derived from phytoplankton (through stress-enhanced mortality), ii) medium size detritus, derived from microzooplankton (through mortality and not assimilated food) and iii) large size detritus derived from mesozooplankton (through mortality and not assimilated food, the latter implicitly accounting for faecal pellets). The formulation describing the predatory interaction between phytoplankton (P in Fig 1) and microzooplankton (Z1 in Fig 1) through the SMP is described in Polimene *et al*. (Polimene *et al*., 2015) and is summarised in SI. The SMP simulates a decrease in ingestion and assimilation efficiency when microzooplankton graze on carbon-rich (i.e., stoichiometrically unbalanced) phytoplankton. The bacterial production of recalcitrant DOC described through the cellular release of excess of carbon and the release of capsular material is detailed in Polimene *et al*. (Polimene *et al*., 2006) and succinctly described in SI. All the equations describing the dynamics of the other plankton functional types and abiotic state variables are fully described in Butenschon *et al.* (Butenschon *et al*., 2016). We refer the reader to these papers for a complete review of model equations, assumptions and underlying philosophy. Model parameters were typically taken from published literature (e.g., Polimene *et al*., 2006; Blackford *et al*., 2004; Polimene *et al*., 2015). The only parameters we tuned were the half saturation constant for food and the food threshold parameter for the functional type Z1. These two parameters were increased (to 40 and 10 mg C m-3, respectively) with respect to the values used in Polimene *et al*. (Polimene *et al*., 2015).

The model was run in an idealized box (Fig. 1) assumed to be homogenously mixed (i.e. zero-dimensional) under constant light and temperature. This assumption allowed us to focus explicitly on the effect of variation of nutrients on the BCP:MCP ratio. In particular, to investigate the impact of increasing nutrients on phytoplankton stoichiometry, grazing mortality, phytoplankton DOM release (through exudation), and BCP:MCP ratio, we explored a range of nutrient concentrations varying from 0.1 to 10 mmol m-3 of nitrate with phosphate at a fixed ratio of 1:16. The model was run with and without the effect of phytoplankton stoichiometry on grazing (SMP) to investigate the specific effect of this process on the simulated MCP: BCP ratio. In the simulations without SMP the standard ERSEM formulation describing grazing in microzooplankton was used (see SI and Butenschon *et al*., 2016). All the fluxes presented in Fig. 2 were calculated after the system had reached a steady-state condition. BCP (mg C m-2 d-1) was calculated through the flux of particulate organic carbon exported (via gravitational sinking) outside of the box represented in Fig. 1. MCP (mg C m-2 d-1) was computed by summing the bacterial production of semi-labile and semi-refractory DOM (as described in Polimene *et al*., 2006) within the box represented in Fig. 1. A sensitivity analysis on the effects of POM sinking velocities on the simulated MCP:BCP ratio is offered in the SI. Phytoplankton aggregation and sinking is not considered in our implementation. However, a sensitivity experiment investigating the effect of this process on the simulation of the MCP:BCP ratio is described in SI

It should be noted that the terms “semi-labile” and “semi-refractory” are used in Polimene *et al.* (Polimene *et al*., 2006) to identify DOC degradable on a time scale of weeks and months, respectively. As such, our model only describes the production of what is commonly classified as semi-labile DOM (Hansell, 2013) and does not consider the refractory DOM with a turn over time spanning from years to millennia. Refractory DOM is the main pool responsible for long term carbon sequestration

 in the ocean and, to the best of our knowledge, it is not considered in currently published model formulations. However, despite the lack of truly refractory DOM, the model we present accounts for the bacterially-mediated transformation of labile to non-labile DOM which is the fundamental process underpinning the MCP (Jiao *et al*., 2010). Here we consider the bacterially mediated production of non-labile DOC described in Polimene *et al*. (Polimene *et al.,* 2006) as the first step leading to the production of refractory DOM (Amon and Banner, 1996) and therefore a proxy of the MCP. Similarly, we recognize that the simulation of carbon sequestration through the BCP (i.e. carbon export below the permanent pycnocline or in the sediments) is not possible in our idealized system. For these reasons, it is important to stress that the simulated fluxes here labelled as MCP and BCP are processes underpinning biologically-mediated carbon sequestration but cannot be used for a quantitative assessment of the carbon effectively sequestered through the two pumps. We argue that this approximation is acceptable for the purely conceptual purpose of this paper.

**Results**

The simulated relationship between phytoplankton internal stoichiometry and the external concentration of nitrate given as initial condition is displayed in Fig. 2A. When nitrate is low (<5 mmol m-3 for the simulation with SMP enabled and lower than 3 mmol m-3 for the simulation with SMP disabled), phytoplankton tend to be unbalanced, reaching C:N ratios higher than the Redfield value of 6.6. The divergence from a balanced condition (C:N≈Redfield ratio) is more pronounced with SMP (C:N >9) than in the run without SMP (C:N ≈7). When nitrate concentration increases, the phytoplankton C:N ratio decreases reaching a threshold value which in our model is assumed to be half of the Redfield ratio (C:N≈3.5). Under high nutrient regimes (nitrate > 5 mmol m-3) the difference between the two simulation scenarios (with and without SMP) is negligible.

With SMP enabled, the ratio between the carbon exuded by phytoplankton and that transferred to the higher trophic levels through grazing changes from a value of ca. 1 to a value higher than 20 (Fig. 2B) once phytoplankton stoichiometry is unbalanced (Fig. 2A). However, in the absence of SMP, this ratio attains a value lower than 1, even under low nutrient regimes. The relationship between external nutrient concentrations and the BCP:MCP ratio is displayed in Fig. 2C. When nutrient concentration is low, the ratio between MCP:BCP increases, concomitantly with the internal C:N ratio and the ratio between exudation and grazing mortality. Model simulations suggest that the MCP:BCP ratio may increase up to 6 times if the external nutrients are decreased and the SMP is considered in the models. The MCP:BCP ratio ranges from 0.6 to 0.9 if the model is run without the SMP.

**Discussion**

Following the seminal work of Jiao and colleagues (Jiao *et al.,* 2010), the potential role played by the MCP in ocean carbon sequestration and its relative magnitude with respect to the BCP received particular attention (Legendre *et al*., 2015; Jiao *et al*., 2014; Hansell, 2013). However, modelling studies synergistically describing both the MCP and the BCP are at present lacking. We have created a modelling framework capable of simulating the nutrient-mediated variability of the MCP to BCP ratio as hypothesized by Jiao *et al*. (Jiao *et al*., 2010). The relationships between external nutrients and the MCP:BCP ratio emerge from the interplay between phytoplankton stoichiometry, zooplankton grazing, phytoplankton exudation and bacterial activity (Fig. 3). In particular, our model suggests that the effect of phytoplankton stoichiometry on microzooplankton grazing is crucial for determining the variability of the MCP:BCP ratio. Without SMP, the MCP:BCP ratio is much more stable, always remaining below 1 regardless of the external nutrient concentration. The importance of the stoichiometric balance of primary producers on trophic dynamics has previously been stressed in laboratory, field and modelling studies (Elser *et al*., 2000; Loladze *et al*., 2000; Mitra, 2006; Polimene *et al.*, 2015). Here we propose that variable stoichiometry within phytoplankton regulates the relative magnitude of the main two biologically-mediated mechanisms of carbon sequestration acting in the ocean. When nutrients are abundant, phytoplankton cells are stoichiometrically balanced (i.e., C:N:P ≈Redfield ratio of 106:16:1) and are effectively controlled by microzooplankton which can rely on highly nutritious food. Microzooplankton is then grazed upon by mesozooplankton and this triggers the evolution of the herbivorous food chain. Under these conditions, carbon export is dominated by zooplankton derived detritus (Fig. 3 and Fig. S1). The MCP plays a relatively minor role in this context as well balanced (nutrient rich) phytoplankton produce a limited amount of DOC. When external nutrients are low, the increase of the cellular C:N:P ratios decrease the nutritional quality of phytoplankton which become less palatable for the grazers. Such a decrease in “food quality” of phytoplankton decreases the grazing pressure from microzooplankton and therefore weakens the link between primary producers and metazoans. Ungrazed, carbon-rich phytoplankton release excess carbon through exudation in order to re-equilibrate internal stoichiometry. This increased production of carbon-rich material stimulates bacterial activity, eventually enhancing the MCP.

It should be noted that the reduction of the grazing activity may lead to an increase of POM in the system due to the increase of phytoplankton derived detritus due to natural mortality. This behaviour is simulated by the model under intermediate level of nutrient (i.e. nitrate > 1 and < 5 mmol m-3, Fig S1). Under these nutrient conditions, the simulated MCP:BCP ratio slightly decreases if higher sinking velocities for POC are assumed (see supporting information and Fig S3). However, the increase of MCP relative to BCP under very low nutrient conditions (i.e. nitrate < 1 mmol m-3) is always simulated by the model, regardless of the sinking velocity of particles (see supplementary information, Fig S2). Phytoplankton sinking due to stress-induced aggregation can also increase the export of particles through the BCP (Turner, 2015). For this reason, we performed an additional sensitivity experiment meant to investigate the effect of this process on the simulated MCP:BCP ratio (see supplementary information, Fig. S4). Model simulation indicated that although affecting the values of the simulated MCP:BCP ratio under low nutrient conditions (Fig. S4), the addition of phytoplankton sink does not change the general dynamics of the system.

While the proposed framework is conceptually plausible and readily usable for theoretical exercises, further model developments are required to simulate the dynamics underpinning the variability of the MCP: BCP ratio in a quantitative way. Detailed information of concentration and distribution of recalcitrant DOM are present in the literature (Hansell, 2013), however, much less is known about mechanisms and processes governing its production. Although some works have indicated that bacteria are capable of producing recalcitrant material over relatively short time scales (Tranvik, 1993; Ogawa *et al*., 2001; Osterholz *et al*., 2015), additional laboratory experiments investigating DOM processing by natural bacterial assemblages are required to identify the processes through which bacteria produce recalcitrant DOM and the environmental factors affecting them. In particular, the role played by the stoichiometry of the substrate in modulating bacteria growth efficiency and recalcitrant DOM production needs to be investigated. This needs to be coupled with laboratory experiments investigating zooplankton feeding behaviour as a function of phytoplankton stoichiometry so that the proposed conceptual framework may be validated. Experimental investigations designed to study all these processes in isolation are the foundations of a robust model development. Such experiments would provide physiological details required for the development of model formulations able to simulate emergent behaviours resulting from the interactions between phytoplankton, zooplankton and bacteria. When the individual model formulations are in place they will need to be merged and validated against observations. Mesocosm experiments provide a suitable framework to test the capability of models to simulate ecosystem interactions starting from single processes. The model thus validated can then be used to interpret environmental data and aid investigation of how climate change conditions may affect the capability of oceans to store anthropogenic carbon.

Even if we are still far from a reliable model capable of quantitatively describing the variability of the MCP:BCP ratio in the global ocean, we have demonstrated that existing process models represent a suitable platform to aid the design of future studies. Interestingly, the two models used were developed independently and for different purposes. This implies that the relationships shown in Fig 2 are emergent properties. This confirms that the addition of more detailed physiology in ecosystem models (e.g., the stoichiometric modulation of grazing) improve their capacity to simulate emergent behaviours (Allen and Polimene, 2011). Our results underline the importance of three elements essential to simulating the variability of carbon fluxes in response to environmental forcing. The first is a fully resolved microbial loop, the second is the variable stoichiometry within plankton functional types and organic matter and the third is a better representation of zooplankton behaviour and physiology. None of the dynamics shown could have been simulated without variable stoichiometry and SMP present in the model.

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**References**

Allen, J. I. and Polimene, L. (2011) Linking physiology to ecology: towards a new generation of plankton models. *J. Plankton Res*., **33**, 989–997.

Ammon, R.M.W. and Benner, R. (1996) Bacterial utilization of different size classes of dissolved organic matter. *Limnol. Oceanogr*., **41**, 41-51.

Blackford, J. C., Allen, J. I. and Gilbert, F. J. (2004) Ecosystem dynamics at six contrasting sites: a generic modelling study. J. Mar. Syst., 52, 191–215.

Butenschön, M., Clark, J., Aldridge, J. N. *et al*. (2016) ERSEM 15.06: a generic model for marine biogeochemistry and the ecosystem dynamics of the lower trophic levels, *Geosci. Model Dev*, in press.

C10011. doi.org/10.5670/oceanog.2014.78.

Elser, J.J., Sterner, R. W., Gorokhova, E. (2000) Biological stoichiometry from genes to ecosystems. *Ecology Letters*, **3**, 540-550.

Hansell, D.A. (2013) Recalcitrant dissolved organic fractions. *Ann. Rev. Mar. Sci*., **5**, 421-445.

Henson, S.A., Sanders, R. and Madsen, E. (2012) Global patterns in efficiency of particulate organic carbon export and transfer to the deep ocean. *Global Biogeochemical Cycles*, **26**, GB1028. http://dx.doi.org/10.1029/2011GB004099

Honjo, S., Eglinton, T.I., Taylor, *et al*. (2014). Understanding the role of the biological pump in the global carbon cycle: an imperative for ocean science. *Oceanography* 27 (3), 10–16. http://dx.

Honjo, S., Manganini, S.J., Krishfield, R.A. *et al*. (2008). Particulate organic carbon fluxes to the ocean interior and factors controlling the biological pump: a synthesis of global sediment trap programs since 1983. *Prog. Oceanog.,* **76**, 217–285.

Jardillier L.,, Zubkov, M.V., Pearman J. *et al*. (2010) . Significant CO2 fixation by small prymnesiophytes in the subtropical and tropical northeast Atlantic Ocean. *ISME J*. 9, 1180-92.

Jiao, N., Herndl, G.J., Hansell, *et al*. (2010). Microbial production of recalcitrant dissolved organic matter: long-term carbon storage in the global ocean. *Nature Reviews Microbiology,* **8**, 593–599

Jiao, N., Robinson, C., Azam, F. *et al*. (2014) Mechanisms of microbial carbon sequestration in the ocean – future research directions. *Biogeosciences*, **11**, 5285–5306. http://dx.doi.org/10.5194/bg-11-5285-2014.

Jones, R.H., Flynn, K.J. (2005). Nutritional status and diet composition affect the value

of diatoms as copepod prey. *Science* **307**, 1457–1459.

Legendre, L., Rivkin R.B., Weinbauer M.G. *et al*. (2015). The microbial carbon pump concept: Potential biogeochemical significance in the globally changing ocean. *Prog. Oceanog*., **134**, 432-450

linking energy flow with elemental cycling. *Bulletin of Mathematical Biology* **62**, 1137–1162.

Loladze, I., Kuang, Y. and Elser, J.J. (2000). Stoichiometry in producer-grazer system: linking energy flow with elemental cycling. *Bulletin of Mathematical Biology* **62**, 1137–1162.

Lutz, M.J., Caldeira, K., Dunbar, R.B., *et al*. (2007). Seasonal rhythms of net primary production and particulate organic carbon flux to depth describe the efficiency of biological pump in the global ocean. *Journal of Geophysical Research-Oceans* 112 (C10). http://dx.doi.org/10.1029/2006jc003706. Artn

Mitra, A. (2006). A multi-nutrient model for the description of the stoichiometric

modulation of predation in micro- and mesozooplankton. *J. Plankton Res*., **28**, 597–611.

Ogawa, H., Amagai, Y. , Koike, I. *et al*. (2001) Production of refractory dissolved organic matter by bacteria. *Science*, **292**, 917-920.

Osterholz, H., Niggemann, J., Giebel, H.A. et al. (2015) Inefficient microbial production of refractory dissolved organic matter in the ocean. *Nature Commun*., 6:7422 doi: 10.1038/ncomms8422

Passow, U. and Carlson, C.A.( 2012). The biological pump in a high CO2 world. *Mar. Ecol. Prog. Ser*., **470**, 249–271. <http://dx.doi.org/10.3354/meps09985>

Polimene, L., J. I. Allen, and Zavatarelli M. (2006), Model of interaction between dissolved organic carbon and bacteria in marine system. *Aquat.Microbiol. Ecol*., **43**, 127– 138.

Polimene, L., Mitra, A., Sailley, S.F *et al*. (2015). Decrease in diatom palatability contributes to bloom formation in the Western English Channel. *Prog. Oceanog*., **137**, 484-497

Polovina, J.J., Howell, E.A. and Abecassis, M. (2008). Ocean’s least productive waters are expanding. *Geophysical Research Letters*, **35**, L03618. http://dx.doi.org/10.1029/2007GL031745

Sarmiento, J.L. and Gruber, N. (2006) Ocean Biogeochemical Dynamics. Princeton University Press.

Siuda, N.S.A., Dam, G.H. (2010). Effects of omnivory and predator–prey elemental stoichiometry on planktonic trophic interactions. *Limnology and Oceanography*

**55**, 2107–2116.

Tranvik, L.J. (1993) Microbial transformation of labile dissolved organic matter onto humic-like matter in seawater. *FEMS Microbial Ecology*, **12**, 177-183

Turner, J.T. (2015) Zooplankton fecal pellets, marine snow, phytodetritus and the ocean’s biological pump*. Prog. Oceanogr*., **130**, 205-248

Urabe, J., Sterner, W.R. (1996). Regulation of herbivore growth by the balance of light and nutrients. PNAS **93**, 8465–8469.

Volk, T. and Hoffert, M.I. (1985) Ocean carbon pumps: analysis of relative strength and efficiencies of in ocean-driven circulation atmospheric CO2 changes. In: Sundquist, E.T., Broecker, W.S. (Eds.), The Carbon Cycle and Atmospheric CO2: Natural Variation Archean to Present. AGU Monograph 32. American Geophysical Union, Washington, DC, pp. 99–110

**Figure Captions**

**Fig. 1**. Box model used in this study. P=generic phytoplankton. Z1= microzooplankton; Z2=heterotrophic nanoflagellates; Z3=mesozooplankton (producing large detritus); B=heterotrophic bacteria; DOM= labile dissolved organic matter; RDOM=recalcitrant dissolved organic matter; POM=particulate organic matter. Dotted arrows indicate density dependent mortality closure (i.e. cannibalism). A simple remineralization closure re-injecting inorganic nutrients from exported detritus at a rate of 0.05 d-1 is applied at the lower boundary of the water column. All living functional types were initialised with 20 mg C m-3 with N and P assumed to follow Redfield stoichiometry. DOM was initialised with 80 mg C m-2 and 1 and 0.1 mmol m-3 of nitrogen and phosphorus, respectively. The initial conditions for RDOM and detritus were assumed to be zero. Blue arrows indicate the main pathways leading to the BCP while red arrows indicate the main pathways leading to the MCP.

**Fig. 2**. Simulated relationships between external nutrient concentration (nitrate) and (A) phytoplankton C:N ratio (mol mol-1), (B) phytoplankton exudation (release of DOC) to grazing mortality ratio (adim), and (C) MCP:BCP ratio (adim). In panel C, the grey line line marks the point where MCP and BCP are equal. For nitrate concentrations higher than 5 mmol m-3 the simulated ratios remained constant. The model was run with and without the Stoichiometric Modulation of Predation (SMP)

**Fig. 3**. Conceptual model linking phytoplankton stoichiometry to the biological and microbial carbon pumps. P=phytoplankton; Z=microzooplankton; mZ=mesozooplankton; B=bacteria; POM=particulate organic matter; DOM=dissolved organic matter; RDOM=recalcitrant DOM. The thickness of the arrows indicates the relative importance of each process in panel A and B. When C:N:P in phytoplankton is highly unbalanced (C:N >9, see Fig. 2), phytoplankton is less efficiently grazed by microzooplankton. This reduces the amount of mesozooplankton and large detritus in the system and the POC pool is dominated by phytoplankton derived detritus. At the same time, phytoplankton release excess of carbon (via exudation) which stimulate the bacterial production of recalcitrant DOM, enhancing the microbial carbon pump (Panel A)When cellular stoichiometry is close to the Redfield ratio (balanced), phytoplankton is effectively grazed by microzooplankton which, in turn, is preyed by mesozooplankton. Detrital carbon is mainly channelled into the POC pool which is dominated by large (fast sinking) particles (e.g. faecal pellets). In this context, carbon sequestration is mainly due to the biological carbon pump (Panel B)).

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**Fig. 1**

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**Fig. 2**

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**Fig 3**