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Special Topic: Marine Carbon Sequestration and Climate Change

Evolving paradigms in biological carbon cycling in the ocean

Chuanlun Zhang1,∗, Hongyue Dang2, Farooq Azam3, Ronald Benner4, Louis Legendre5, Uta Passow6, Luca Polimene7, Carol Robinson8, Curtis A. Suttle9 and Nianzhi Jiao2,∗

1Department of Ocean Science and Engineering, Southern University of Science and Technology, Shenzhen 518055, China; 2State Key Laboratory of Marine Environmental Science, Institute of Marine Microbes and Ecospheres, College of Ocean and Earth Sciences, Xiamen University, Xiamen 36102, China; 3Scripps Institution of Oceanography, University of California San Diego, La Jolla, CA 92039, USA; 4Department of Biological Sciences and School of the Earth, Ocean and Environment, University of South Carolina, Columbia, SC 29208, USA; 5Sorbonne Universités, Laboratoire d’Océanographie de Villefranche, LOV, 06230 Villefranche-sur-Mer, France; 6Marine Science Institute, University of California Santa Barbara, CA 93106, USA; 7Plymouth Marine Laboratory, Prospect Place, The Hoe, Plymouth, PL1 3DH, UK; 8School of Environmental Sciences, University of East Anglia, Norwich, NR4 7TJ, UK and 9Departments of Earth, Ocean and Atmospheric Sciences, Botany, and Microbiology and Immunology, and the Institute for the Oceans and Fisheries, University of British Columbia, Vancouver, British Columbia, V6T 1Z4, Canada

∗Corresponding authors. E-mails: zhangcl@sustc.edu.cn; jiao@xmu.edu.cn

ABSTRACT

Carbon is a keystone element in global biogeochemical cycles. It plays a fundamental role in biotic and abiotic processes in the ocean, which intertwine to mediate the chemistry and redox status of carbon in the ocean and the atmosphere. The interactions between abiotic and biogenic carbon (e.g. CO2, CaCO3, organic matter) in the ocean are complex, and there is a half-century-old enigma about the existence of a huge reservoir of recalcitrant dissolved organic carbon (RDOC) that equates to the magnitude of the pool of atmospheric CO2. The concepts of the biological carbon pump (BCP) and the microbial loop (ML) shaped our understanding of the marine carbon cycle. The more recent concept of the microbial carbon pump (MCP), which is closely connected to those of the BCP and the ML, explicitly considers the significance of the ocean’s RDOC reservoir and provides a mechanistic framework for the exploration of its formation and persistence. Understanding of the MCP has benefited from advanced ‘omics’ and novel research in biological oceanography and microbial biogeochemistry. The need to predict the ocean’s response to climate change makes an integrative understanding of the BCP, ML and MCP a high priority. In this review, we summarize and discuss progress since the proposal of the MCP in 2010 and formulate research questions for the future.

Keywords: biological carbon pump, microbial loop, microbial carbon pump, ocean carbon cycle, global climate change

INTRODUCTION

The modern ocean accounts for ∼50% of global photosynthesis, with its primary production of organic matter forming the core of the ocean carbon cycle. Thus the ocean has a major influence on the chemistry and redox status of the atmosphere through the net uptake of atmospheric CO2 and net release of molecular oxygen. An early estimate showed that about 25% of the ocean’s primary production was transported to the interior of the ocean (below the euphotic zone) via the biological carbon pump (BCP) [1]; later on, this number was changed to 10–15% for gravitational sinking with another 5% each for passive transport by water motion and active transport by vertical migrators [2]. Carbon transported to the deep ocean (>1000 m) is sequestered on timescales of > 100 years up to 1000 years (i.e. the residence time of deep waters). About 0.3% of the ocean’s primary production is buried in marine sediments [3,4], some of which eventually forms a major reservoir of organic matter that persists for hundreds of millions of years in rock formations (Fig. 1).

Since the industrial revolution, the ocean is estimated to have taken up approximately 25% of the anthropogenic CO2 [5], resulting in ocean acidification with consequences for biogeochemical and climatological processes and the ocean carbon cycle [6,7,1,8]. Global warming and ocean acidification and their respective consequences influence the functioning of the BCP, a major pathway for
Figure 1. Cycling of biologically produced organic carbon (POC and DOC) in the ocean and links between the seafloor and the atmosphere: the BCP, which transports organic matter from the surface to the interior of the ocean; the MCP, which converts parts of labile organic carbon into RDOC via microbial activities, mainly by heterotrophic archaea and bacteria, and associated viruses.

EVOLUTION OF OUR UNDERSTANDING OF THE MICROBIAL ROLE IN DOC GENERATION AND DEGRADATION

Understanding of the ocean’s carbon cycle in the late twentieth century was largely promoted by the BCP (called ‘soft tissue pump’ in [28]) and the ML [29]. The term ‘pump’ was initially used to refer to the movement of carbon against a concentration gradient between the surface ocean and the deep ocean [28]. Both concepts find their roots in Dugdale and Goering [30], who recognized new (BCP) and regenerated (ML) production in the ocean.

The BCP begins in the euphotic zone where photosynthetic organisms fix dissolved CO₂ to produce particulate organic carbon (POC) (Fig. 2). Particulate organic matter (POM) consists of both living and non-living components, and most of it is respired to CO₂ by metabolic processes in the epipelagic ecosystem. The subsequent export of a small fraction of the POM is carried out by gravitational flux, vertical migrations of zooplankton and physical subduction of water masses, which remove the organic matter to deeper regions where it accumulates or is respired. The respiratory CO₂ at depth is removed from contact with the atmosphere for a period corresponding to the residence time of deep waters, namely tens to hundreds of years below 100 m and thousands of years below 1000 m (Fig. 2). In addition, organic matter in particulate or dissolved form reaching the latter depth via the BCP...
Increasing atmospheric CO₂ concentration raises several questions: (1) Will the ocean continue to take up carbon? (2) At what rate? (3) For how long will the exported carbon remain removed from the atmosphere? These questions address the functioning and efficiency of the future BCP. Global warming and past carbon sequestration (ocean acidification) will also change the BCP leading to the next question: (4) How will the biological pump respond to the consequences of increased carbon input combined with warming? [31]. One scenario suggests that, in the coming decades, decreasing phytoplankton cell size will decrease the downward POC flux from the surface ocean, while changes in zooplankton community structure will decrease the downward POC flux in subsurface waters [32]. However, other predictions suggest alternative outcomes and the answers to these questions are still discussed controversially in the scientific community. A recent report on a transformative understanding of the ocean’s BCP to the US National Science Foundation [33] recommended three major research directions addressing (i) food web regulation of export, (ii) the dissolved-particulate continuum, and (iii) variability of organic transport in space and time. Several large programs, such as the ongoing US-EXPORTS [2] and the UK COMICS [34] programs as well as many other efforts are currently focusing on the BCP.

Though many forms of vertical export can be related to BCP, it mainly focuses on particles that move downward through physical and biological forces (i.e. by gravity and transport by vertically migrating zooplankton). The ML, on the other hand, intimately links intricate interactions between microorganisms and their physical and chemical surroundings [29,35]. The ML focuses on carbon cycling in the water column where bacteria (actually referring to both bacteria and archaea), protozoa and viruses determine the fate of DOM [35]. It was estimated that bacteria could channel up to 50% of marine primary production into the ML, highlighting their importance in the ocean’s carbon cycle [35,36]. Similarly, Legendre and Rivkin [37] found

Figure 2. Schematic depiction of the BCP, the ML and the MCP. The remineralization length scale in the left part of the figure shows the return of resired CO₂ back to the surface, from three depth zones (modified from [31]).
Table 1. Definitions and major impacts of the BCP, ML and MCP.

| Concept              | Definition                                                                 | Major impacts and focus                                                                 | Reference   |
|----------------------|-----------------------------------------------------------------------------|----------------------------------------------------------------------------------------|-------------|
| Biological pump      | A complex ecosystem process that transports particulate organic carbon from the epipelagic zone to the deep interior of the ocean and further to the ocean floor | Sequestration of atmospheric CO₂ through vertical transportation of living biomass to marine sediments; focusing on sediment storage | [28,42]     |
| Microbial loop       | A ‘feedback’ pathway of loss of the primary production to the environment in the form of dissolved organic matter and the utilization of the latter by bacteria that feed the protozoa, which enter the food chain | The role of bacteria in sequestering nutrients from the environment, which are consumed by protozoa; focusing on organismal populations above thermocline | [29,35]     |
| Microbial carbon pump| A conceptual framework for understanding the role of microbial processes in the production of recalcitrant dissolved organic matter in the ocean water column | Sequestration of atmospheric CO₂ through transformation of labile organic matter to recalcitrant organic matter; focusing on capacity of the ocean to store atmospheric CO₂ | [9,50]      |

that heterotrophic microbes always dominate respiration in the euphotic zone, even when most particulate primary production is grazed by metazoans. The ML intertwines with the grazing food web and provides a mechanism to retain nutrients such as N and P in the highly stratified upper oligotrophic oceans by recycling them through pico-phytoplankton, bacteria and microzooplankton [29] (Fig. 2).

The MCP complements and connects the concepts of BCP and ML, additionally including the idea of the VS, into a more integrated concept of the cycling of biogenic carbon in the ocean. The VS, which refers to the release of carbon and nutrients back into the environment due to cell lysis, is tightly connected to the BCP, the ML and the MCP because cell lysis transforms living POM into DOM and non-living POM [38,18]. As much as a quarter of the C fixed by phytoplankton is estimated to flow through the VS [18], thereby promoting ecosystem respiration [39]. The released DOM and POM are largely of bacterial origin and, hence, relative to bacterial requirements (because of the carbon required for respiration), have too little carbon relative to other nutrients. This shortage of carbon is exacerbated because of the recalcitrant nature (e.g. cell-wall material) of some of the carbon released by cell lysis. Therefore, as the lysis products are processed by the ML, the more accessible DOM is metabolized, releasing inorganic nutrients, altering pathways of nutrient cycling [40,41] and enriching the pool of less labile DOC. This process directly couples the VS to the ML and MCP, and has been termed the ‘shunt and pump’ [17].

The BCP, ML and MCP have distinct ecological or biogeochemical meanings (Table 1) and each has influenced multiple research disciplines (Table 2).

These three concepts are fundamental in developing global biogeochemical and ecological models that rely on understanding organismal biology and the interactions between the POC and DOC pools (Fig. 3).

Several reviews provide thorough descriptions of the BCP and the ML (e.g. [36,42,43,31,11]). Here, we focus on recent progress concerning the MCP in the context of the BCP and ML.

PROGRESS ON THE MCP DURING THE LAST EIGHT YEARS

During the last eight years, our understanding of the MCP has advanced appreciably (e.g. [44–49,26,50,51]), specifically addressing some of the questions raised in Jiao et al. [9]. In particular, substantial progress has been made on the composition of recalcitrant DOM, the mechanisms of its formation, the nature of its interactions with ML biogeochemistry and the associated community shifts and trophic dynamics. There were also gains in our understanding of the microbial processing of DOM at various taxonomic and functional group levels (e.g. [52–54]) (Table 3). The state of the art of these topics will be discussed in the remainder of this review.

IDENTIFICATION AND QUANTIFICATION OF THE COMPOSITION OF RDOM

According to Hansell et al. [14], less than 1% of the DOC in the ocean is labile and 94% is refractory, while the remaining 5% is classified as semi-labile (note: [13] divided the DOC into labile,
Table 2. Impacts of the three original publications that defined the ML [29], BCP [28] and MCP [9] in different research disciplines, based on the definition of the disciplines in the Web of Science [v.5.29] Core Collection Result Analysis (http://apps.webofknowledge.com). Data were up to 6 July 2018. The percentage value for each discipline is the standardized percentage of the citations in a discipline (minimal 10 citations) vs. the total citations of BCP, ML or MCP since their publication.

| Research discipline defined by Web of Science | ML % of 3051 citations since 1983 | BCP % of 308 citations since 1985 | MCP % of 357 citations since 2010 |
|---------------------------------------------|---------------------------------|---------------------------------|---------------------------------|
| Microbiology, Biodiversity, Biotechnology    | 18                              | 1                               | 17                              |
| Environmental Sciences, Ecology              | 23                              | 21                              | 22                              |
| Geology Geochimistry Geophysics, Chemistry   | 3                               | 27                              | 23                              |
| Marine Freshwater Biology                    | 32                              | 8                               | 9                               |
| Meteorology Atmospheric Sciences             | 0                               | 12                              | 1                               |
| Oceanography, Science Technology other topics| 24                              | 30                              | 28                              |

Characterization of specific biochemicals in RDOM

Carbohydrates, amino acids and amino sugars
Early studies examined the composition of RDOC based on measurements of common biochemicals, such as carbohydrates, amino acids and lipids. Ogawa et al. [55] reported the transformation of labile substrates (D-glucose and D-glutamate) into refractory forms of hydrolysable neutral sugars, amino sugars and amino acids that persisted after 1 year in bioassay experiments. The concentrations of these compounds were later confirmed to be similar to those reported for natural deep ocean waters [63] and represented less than 2% of the total RDOC in low-molecular-weight DOC [16]. In particular, D-enantiomers of amino acids have been observed to contribute to the RDOC pool and are predominantly derived from bacterial sources [56,63]. The ratio of the D-amino acids vs. L-amino acids has been used as a proxy for the degree of recalcitrance, which increases dramatically from bulk POM to the refractory low-molecular-weight DOM [16] (Table 4).

Microbial lipids
Microbial lipids may be important compounds contributing to the RDOC pool in the ocean [64]. Some lipids are much more resistant to degradation than carbohydrates or proteins (hydrolysed to amino acids) [16] and can be preserved in sediments or rocks for millions or billions of years [65–67]. Most studies of microbial lipids have been conducted in sediments or POM (e.g. [68–71]) because of the requirement for a large amount of organic material for lipid analysis. Selective accumulation of the refractory lipid-like material in the water column has been demonstrated by the increasing alkanes in the pyrolyzates of sinking POC as depth increased in the
Mediterranean Sea [72]. Alkanes from Proterozoic rocks were also identified as biomarkers of heterotrophic bacteria [66]. These biomarkers might have been derived from MCP activity that contributed to the large DOC pool that may have been 100–1000 times greater than in the modern ocean [73–75]. Lipid-like macromolecules in the deep ocean have similar radiocarbon ages and δ13C values as the majority (∼70%) of the uncharacterized acid-insoluble fraction, indicating that the bulk POC may be compositionally similar to the lipid-like macromolecules [64] (Table 4).
Table 3. Progress made over the past 8 years related to microbial carbon pump framework.

| Question asked in Jiao et al. [9]                                                                 | Progress made* | Note                                                                                                                                                              | Representative papers |
|------------------------------------------------------------------------------------------------|----------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------|
| What are the concentrations, compositions and spatiotemporal variations of recalcitrant dissolved organic matter (RDOM) in the ocean? | ✓ ✓ ✓          | Considerable progress has been made in addressing this question with the past 8 years (see Table 4 for more information)                                         | [16,49,87]            |
| Is the RDOM inventory currently changing and, if so, is the rate of change fast enough for RDOM to serve as an evolving reservoir for stored carbon?   | ✓              | The projected global warming is likely to enhance MCP, which may produce more RDOC in future ocean. However, current information is insufficient to make any concrete conclusion     | [137,149]            |
| Why do heterotrophic bacteria and archaea not degrade RDOM in 'microbial timeframes'?          | ✓ ✓           | Degradation of RDOM by heterotrophic bacteria or archaea is constrained by a specific environment (RDOCt) or low concentration (RDOCc)                             | [46,47,110,111]       |
| What are the structural and biochemical constraints on degradability?                           | ✓ ✓ ✓          | Progress has been made in addressing this question with the past 8 years (see Table 4 for more information)                                                         | [16,26,87,49]         |
| What environmental conditions make RDOM more or less degradable?                               | ✓              | The rate of RDOM degradation is affected by multiple variables, among which temperature, sunlight, pH, redox, nutrient availability and water movement all play a role | [46,111,150]         |
| Can we predict the chemical composition of the degradation products?                           | ✓              | Several proxies have been developed for characterization of the composition of RDOC. However, the exact composition of RDOC is still largely unknown                     | [49,87,102,105]       |
| What is the taxon-specific variation in the degradability of RDOM?                             | ✓ ✓ ✓          | The best examples are copiotrophic organisms that prefer to degrade carbohydrates and oligotrophic organisms preferring to utilize nitrogen-containing compounds         | [52,118]              |
| Is ecosystem energy supply also a constraint on RDOM degradability?                            | ✓              | Energy supply at the ecosystem level would be an ultimate constraint on RDOM production and degradation. This question is poorly addressed so far                       | [53,134]              |
| How does organic-matter flux through the microbial loop affect RDOM lability?                  | ✓              | The flux of labile DOM through the microbial loop can serve as a priming agent enhancing RDOM removal                                                             | [111]                 |

*The level of progress made most (✓ ✓ ✓ ✓) or least (✓) since Jiao et al. [9].

The greater ages of lipid-like material than carbohydrate- and protein-like substances were also observed in the DOM pool of the open Atlantic and Pacific Oceans [76]. In particular, the deep-water lipid extract was 13–14 kyr older than the corresponding protein- and carbohydrate-like components in the DOM. This lipid extract was also up to 1 kyr older than the high-molecular-weight DOM. However, the δ13C values of the high-molecular-weight DOC were more similar to the carbohydrate- and protein-like substances than to the lipid extracts, in contrast to the observations of POC [64]. This suggests that deep-ocean POM and DOM have different origins, with the latter having undergone more extensive recycling [76] (Table 4).

Hwang et al. [64] and Loh et al. [76] did not identify specific lipid compositions in either the POM or DOM fractions. However, numerous studies focusing on POM have shown diverse lipid biomarkers from planktonic archaea, bacteria and phytoplankton [77–81,69,82,83]. In particular, crenarchaeol was identified as a major glycerol dialkyl glycerol tetraether (GDGT) biomarker for planktonic Thaumarchaeota that are present in the global ocean at
Table 4. Summary of compounds or proxies for description of recalcitrant-dissolved organic matter.

| Compounds or proxy                      | Description or method                                                                 | References |
|-----------------------------------------|---------------------------------------------------------------------------------------|------------|
| Aminosugars, amino acids                | Total hydrolysable, accounts less than 5% of RDOM                                     | [16,55]    |
| D-L-amino acids                         | Ratio of D-enantiomers of amino acids vs. L-enantiomers of amino acids                 | [16,63]    |
| High D:L ratio in RDOM                  |                                                                                       |            |
| Lipid-like macromolecules in POM        | Acid-insoluble fraction                                                               | [64]       |
| Lipid-like extracts in DOM              | Organic solvent extracts, contributing to 0.1–0.3% of DOM_HMW                        | [76]       |
| Glycerol dialkyl glycerol tetraethers, Crenarchaeol | Total lipid extracts in 0.2–0.7 μm fraction                                               | [86]       |
| Carotenoid degradation product          | Solid-phase extracted (SPE) DOM using comprehensive gas chromatography coupled to mass spectrometry; contribute to ~4% of total DOM | [87]       |
| DOC:DON                                 | The percentage of DOC mineralization is negatively correlated with the initial percentage of total fluorescence | [89]       |
| TDAA (%DOC)                             | TDAA (%DOC) is defined as the ratio of carbon in total dissolved amino acids to the whole DOC. DOM is thought to be refractory when TDAA (%DOC) is less than 0.7% | [91,92]    |
| % Humic-like fluorescent (F) DOM        | The percentage of DOC mineralization is negatively correlated with the initial contribution of humic-like fluorescence to the total fluorescence | [89,96,97] |
| Humic-like FDOM                         | Humic-like FDOM (Ex = 320 nm, Em = 420 nm) are thought to be bio-refractory            | [94]       |
| % Protein-like FDOM                     | The percentage of DOC mineralization is positively correlated with the initial contribution of protein-like fluorescence to the total fluorescence | [89,96,97,151] |
| Specific UV absorbance (SUVA)           | SUVA positively correlates with aromaticity and thus the recalcitrance of DOM         | [89,95–97] |
| Size-age-reactivity continuum           | Increasing decomposition along the flow of organic carbon from larger to smaller size classes results in greater chemical complexity, less biological reactivity and older radiocarbon ages of the organic matter | [16,58,152] |
| CRAM (carboxyl-rich alicyclic molecules)| CRAM are thought to be refractory and derived from terpenoids. They are defined with these criteria: DBE/C = 0.30–0.68; DBE/H = 0.20–0.95; DBE/O = 0.77–1.75. CRAM account for 8% of the RDOC in the ocean | [88]       |
| IOS (island of stability)               | IOS (falling into the area with the same criteria as CRAM) is thought to be the most stable combination of elements in a distinct window of H/C (1.17 ± 0.13), O/C (0.52 ± 0.10) and molecular mass (360 ± 28 and 497 ± 51 Da). IOS compounds contribute about 50% of SPE-DOM | [49]       |
| Degradation Index (IDEg)                | IDEg was developed to compare the degradation state of marine SPE-DOM samples analysed with FT–ICR MS based on correlation between peak intensity and 14C. IDEg = Σ(magnitudes NEGIdeg)/Σ(magnitudes(NEGIdeg + POSIdeg)) Since higher IDEg indicates older age, IDEg could be positively related to DOM recalcitrance | [102]      |
| Terrestrial indicator compounds         | The 184 terrestrial formulae, identified in most river samples and ocean samples based on correlation between peak intensity and 13C, are thought to be resistant to degradation. It contributes 2–3% to SPE-DOM in ocean samples | [105]      |
a total inventory of $10^{28}$ cells [84]. GDGTs can be preserved in sediments for millions of years [85] and can be a significant component of the lipids in the RDOC pool (Table 4). Because *Thaumarchaeota* cell size is small, they are more abundant in the DOM fraction (operationally defined as the fraction passing through a ∼0.7-μm filter) than the particulate organic fraction [86]. Measurements of the dissolved phases of lipids give total GDGT abundance in the tens of nanograms per liter range [86]; however, once the organisms die, their core lipids may be incorporated into larger particles (0.7- to 60-μm size fraction) that can be more quickly transported into the deeper ocean and buried in marine sediments (Table 4). The same mechanism may apply to bacterial lipid accumulation in the POM fraction that is preserved in marine sediments. It is unknown, however, how much bacterial or archaeal lipids are actually present in the uncharacterized fraction of the RDOM because the uncharacterized RDOM is largely acid-insoluble and cannot be identified by regular gas chromatography or liquid chromatography mass spectrometry.

**Coupling between molecular size and radiocarbon age of DOC**

It has been observed that the distribution of total organic carbon in the global ocean is heavily skewed toward the nanometer size range [16]. A hypothesis is that bioavailability of the organic matter decreases with decreasing size and alteration of the organic molecules (Fig. 3 insert), meaning that smaller-sized classes of organic molecules are more slowly remineralized by microorganisms [98,16]. This has been confirmed by approaches coupling the chemical composition and radiocarbon content of marine organic matter in different size fractions ([76,58]. In Loh et al. [76], seawater from different depths of the central North Pacific and the Sargasso Sea region of the North Atlantic showed that the δ14C values ranged from −5 to −434‰ for high-molecular-weight DOM and from −210 to −539‰ for low-molecular-weight DOM, with the latter being older than the former by 1650–1850 kyr. The low-molecular-weight DOM was also the most abundant (77–95%) fraction of total DOM, consistent with the overall dominance of RDOM in the ocean [14]. Walker et al. [58] examined the C:N ratio and 14C age of organic matter in different size classes from the coastal, surface and deep waters of the Pacific Ocean. In all three environments, larger particles were characterized by young ages and nitrogen enrichment and smaller molecules by older ages and nitrogen depletion. The size–age–composition relationship was also observed in marine sediments with pore water DOC being dominated by low-molecular-weight DOM [99].
In addition to the relationships between size, age and composition, a recent study observed declining concentrations of high-molecular-weight DOM correlated with increasing apparent oxygen utilization along the shallow overturning circulation cell of the Mediterranean Sea [93]. Decreases in high-molecular-weight DOM accounted for about 30% of DOM mineralization. The apparent low-molecular-weight DOM experienced little mineralization, indicating microbes primarily utilized high-molecular-weight molecules, whereas the smaller size classes resisted degradation and were the primary source of recalcitrant DOM in the deep ocean [93].

Characterization of RDOM composition using FT–ICR MS

It is well established that RDOM is composed of less than 10% of common biomolecules such as carbohydrates, amino acids or lipids (see discussion above). Proxies such as the DOC:DOM ratio, TDAA (%DOC), fluorescent DOM or the size-age relationship provide insights about the composition and reactivity of DOM, but additional analytical approaches are needed to understand RDOM composition. One approach, Fourier transform ion cyclotron resonance mass spectrometry (FT–ICR MS), has gained popularity in recent years because it identifies thousands of molecular formulae, which can be further analysed in detail. FT–ICR MS was proposed over 20 years ago [100] and has been increasingly applied in the characterization of changes in DOM composition in both terrestrial and marine environments and along environmental gradients [49,88,101–106].

A number of proxies have been developed based on characterization of DOM using FT–ICR MS. CRAM are commonly believed to be refractory and occur as the most abundant components of DOM in the deep ocean. Using the FT–ICR MS technique, Hertkorn et al. [88] identified over 613 CRAM (Table 4), which can be constrained by the double bond equivalent (DBE) normalized to C (DBE/C = 0.30–0.68), H (DBE/H = 0.20–0.95) or O (DBE/O = 0.77–1.75) within the van Krevelen diagram. These compounds are characterized by abundant carboxyl groups and alicyclic rings commonly found in terpenoids that occur as membrane constituents or secondary metabolites in diverse prokaryotic and eukaryotic organisms [107]. Such findings can be linked to the GC/GC MS analysis of the CDP that can account for 4% of the RDOM component [87], which agrees with the estimate that CRAM account for 8% of the DOC [88]. Lechtenfeld et al. [49] further identified 361 most stable molecular formulae, called the ‘island of stability’ (IOS) (Table 4) within the CRAM domain [49] in the Atlantic and Southern Ocean waters. These molecules are deemed potential indicators of refractory DOM in the Southern Ocean; however, it is unknown whether the same IOS compounds exist in other oceanic environments.

Another proxy called the degradation index (I_{DEG}) was developed by Flerus et al. [102] to describe the degradation status of marine DOM analysed with FT–ICR MS from solid-phase extraction (SPE) samples (Table 4). I_{DEG} was calculated using 10 mass peak magnitudes that have either significant linear positive or negative correlation with the Δ^{14}C values of the SPE-DOM. The value of I_{DEG} ranges between 0 and 1 with higher I_{DEG} indicating older age and greater recalcitrance of the DOM. Analysis of seawater at 37° N and 14° W from the eastern Atlantic Ocean showed that I_{DEG} values increased from 0.756 at 400–500 m to 0.808 at 4000–5000 m, consistent with the notion that DOM from deeper water is more refractory than shallower water. Likewise, the I_{DEX} was developed based on the SPE-DOM samples from the Atlantic Ocean, which needs to be verified in other oceanic regions [102].

Lastly, Medeiros et al. [105] identified 184 molecular formulae (Table 4) using FT–ICR MS and used them to indicate riverine inputs in the deep North Atlantic and North Pacific Oceans. These compounds are most enriched in river water and correlated well with known terrigenous tracers in the deep ocean waters, based on which the authors concluded that terrigenous organic matter can be preserved in the deep ocean [105]. This observation is consistent with the deep-ocean distributions of dissolved lignin phenols—biomarkers derived from terrestrial plants [108].

FT–ICR MS and nuclear magnetic resonance spectroscopy have been used together to trace the source of deep-ocean RDOC from surface primary production. Zhao et al. [109] observed that cultured picocyanobacteria, *Synechococcus* and *Prochlorococcus*, released fluorescent DOM that underwent similar photo-degradation behavior when compared with deep-ocean fluorescent DOM (Table 4). Ultrahigh-resolution mass spectrometry and nuclear magnetic resonance spectroscopy revealed abundant nitrogen-containing compounds in *Synechococcus* DOM, which may originate from degradation products of the fluorescent phycobilin pigments. Their results suggested that picocyanobacteria are likely to be important sources of marine autochthonous fluorescent DOM, which may accumulate in the deep ocean as RDOC [109].

Proxies of RDOM in carbon cycle studies must be used with caution given the current constraints in
defining the composition and reactivity of RDOC. Jiao et al. [46] used the term RDOC\textsubscript{t} to describe RDOC compounds maintaining recalcitrance in a specific environmental context and used RDOC\textsubscript{c} to describe RDOC compounds being inaccessible to microbes due to their extremely low concentrations. It was debated whether low concentration of any DOC compound is the predominant reason for RDOC to remain recalcitrant in the ocean [47,110]. Recent evidence indicates that only a small fraction of RDOC molecules are too dilute for microbial utilization and that environmental conditions, including exposure to photochemical alterations in surface waters and varying microbial communities, are critical for the removal of RDOC from the ocean [111]. The size–age–composition relationship that organic-matter size is negatively correlated with radiocarbon age and carbon:nitrogen ratios also supports the dominant role of chemical composition (RDOC\textsubscript{t}) in determining the long persistence of the RDOC pool [58,112].

In addition, if the majority of deep oceanic DOC is RDOC\textsubscript{c}, namely the dilution hypothesis dominates deep oceanic DOC persistence, the $\Delta^{14}C$ in the deep ocean calculated from a mass balance model of deep oceanic diluted DOC would be difficult to reconcile with the observed $\Delta^{14}C$ (4000–6000 years) for deep oceanic DOC [113]. This is because, with this observed age constraint, the box model of diluted DOC in the deep ocean would result in either (i) labile DOC comprising a relatively large fraction of bulk DOC but with radiocarbon ages similar to or older than bulk radiocarbon ages or (ii) a smaller labile DOC pool with much younger radiocarbon ages; the latter would be most consistent with a variety of other observations [114].

**MECHANISMS AND PROCESSES OF RDOC PRODUCTION**

Studies on the MCP have attempted to address the grand challenges of dissecting the composition of the bulk RDOM and identifying the diverse microbial populations responsible for the fate and complexity of RDOM; both are still largely ‘black boxes’. The research community has reached a consensus that in-depth and integrative characterization of both complex DOM compounds and microbial communities is a prerequisite for exploring the relationship between microbial community composition and the processing of DOM [27,115]. Hopes are high to unveil the intimate linkages between the two black boxes by using the advanced technologies provided by both genomics and bioinformatics, and by mass spectrometry capabilities [25,27,51,116].

Here we present some of the latest advances on focused groups of marine organisms as well as community shifts and trophic dynamics associated with RDOM production.

**Carbon metabolism of known organisms**

Bacterial metabolism of organic matter is constrained by their physiological capability and biochemical pathways for processing organic molecules. The most studied marine bacteria have been the ‘eutrophic’ *Roseobacter* clade and the ‘oligotrophic’ SAR11 clade of marine alphaproteobacteria [117]; both are numerically dominant and functionally important groups of marine bacteria [52]. These clades have distinct patterns of DOC utilization, with *Roseobacter* clade strains mostly taking up carbohydrates and SAR11 preferring nitrogen-containing DOC such as amino acids, which are attributed to different capabilities of ATP binding cassette transporters among these organisms [45,52,118]. Two other studied groups of marine bacteria are the *Gammaproteobacteria* and the Cytophaga-Flavobacterium-Bacteroides, which are known to be capable of metabolizing macromolecules through the TonB-dependent transporter proteins [52,118]. Cottrell and Kirchman [43] observed in estuarine and coastal environments that the Cytophaga-Flavobacterium-Bacteroides cluster showed overrepresentation in the assemblage consuming chitin, N-acetylglucosamine and protein but underrepresentation in the assemblage consuming amino acids. Tang et al. [119] demonstrated through multi-omics analysis and cultivation experiments that the *Bacteroidetes* strain *Gramella flava* JLT2011 (*Flavobacteria*) has the ability to grow on a wide range of polysaccharides such as xylan and homogalacturonan from pectin, which are operated by different polysaccharide utilization loci (PUL) or PUL-like systems. *Flavobacteria* have also been demonstrated to be a major contributor for the utilization of exopolysaccharides that represent an important source of organic carbon in marine ecosystems [120]. However, *Flavobacteria* could not completely utilize exopolysaccharides and fluorescent DOM (e.g. humic acid-like substances) produced during metabolism of exopolysaccharides, which may be refractory and may contribute to the carbon storage in the oceans [120]. While these model organisms provide specific knowledge of carbon compounds they metabolize, it is uncertain how these compounds can be identified in natural environments where complex community interactions occur (see below).
Carbon metabolism of natural populations

Studies using individual organisms under laboratory conditions often focus on limited substrates of known compositions. However, the situation is much more complex for natural populations regarding which bacteria may utilize which carbon compounds and whether such compounds in turn may affect specific bacterial community composition [121]. Multiple reports demonstrate that specific carbon compounds can select for particular species or groups of organisms under different environmental conditions [122]. For example, low-molecular-weight molecules (e.g. monomers amino acids, sugars, short chain fatty acids) can be easily transported across cell membranes and may be utilized by most heterotrophic bacteria or archaea. However, it has been demonstrated that different low-molecular-weight organic compounds stimulated growth of different types of bacteria, leading to the suggestion that changing composition of the DOC pool can selectively alter the community structure of bacterioplankton [121]. This is consistent with observations of the distribution of Roseobacter or SAR11 types of organisms selecting for different types of organic substrates (see above). However, it also has been demonstrated that it is the quantity and not the quality of phytoplankton-derived DOC that selects for different types of bacteria in a given range (10–100 μM) of substrate concentrations [54].

The importance of community composition for the fate of DOM has also been shown [53,115]. For example, in incubation experiments using only <1.0-μm microbial populations, DOM composition was dominated by compounds with lipid and peptide characteristics; whereas, in incubations with the presence of organisms larger than 1.0 μm, the DOM composition from the culture experiment was nearly identical to that in the natural water, indicating the role of larger microorganisms in constraining DOM composition in the marine environment [53]. These studies highlight the importance of both microbial community structure and composition or abundance of DOM in the marine system, which should allow distinction between RDOCt and RDOCe to better understand the MCP framework (see above).

The interplay between bacterial community and DOM composition is also examined by comparing particle-attached vs. free-living organisms using genomic tools [123–128]. Despite our awareness of the different ecological strategies of particle-associated and free-living microbes (e.g. [129]), we know little about the principles behind the phylogenetic differences and life strategies between free-living and particle-attached microbes in the marine environment [126,130]. Particle-associated microbes are capable of utilizing a variety of substrates under nutrient-rich conditions. Free-living heterotrophs, on the other hand, often face a massive pool of refractory dissolved organic molecules under oligotrophic conditions [130,131]. However, Zhang et al. [132] observed that the composition of POM was more strongly related to the free-living than to the particle-attached bacterial community, which indicates that POM composition may significantly influence the free-living bacterial community through the release of labile or semi-labile organic matter from particles contributing to the bioavailability of DOC [132]. The nutritional status of the environment may also affect the difference between particle-attached and free-living populations. For example, in the deep ocean when substrates (ammonia, for example) are scarce, particles provide concentrated life-supporting microenvironments. Microorganisms adapted to a particle-attached lifestyle show the dominance of extracellular hydrolytic enzymes; free-living bacteria, on the other hand, are characterized by hydrolytic enzymes typically bound to the cell surface [130]. In the eutrophic surface ocean and estuaries, substrates or nutrients are abundant and organisms were found to be similar between particle-attached and free-living populations [129,133].

Microbes–DOM interaction at the ecosystem level

The finding of Kujawinski et al. [53] that incubation experiments using the whole water community resulted in DOM composition similar to the natural water composition highlights the need to examine the microbes–DOM interaction at the ecosystem scale (Fig. 3). This is convincingly demonstrated by a long-term large volume (> 100 tons) water column (12 m in depth) incubation, which showed solid evidence of the effective microbial transformation of organic matter from labile to refractory states [48]. Another study provides metagenomic evidence of system-level dynamics of microbes–DOM interactions, utilizing the Tara Ocean data that included comprehensive sequences of eukaryotic, prokaryotic and viral lineages from samples collected within the euphotic zone of ocean waters [134]. The increased carbon export in this water column was found to correlate not only with bacteria, particularly Synechococcus, but also several unicellular eukaryotic microorganisms including three Rhizaria lineages and three dinoflagellate lineages that have previously not been
believed to play important roles for carbon flux. Also important is the finding of a correlation between the abundance of *Synechococcus* phages and increased carbon export at depth, indicating that phage induced cell lysis promotes particle sinking through enhanced aggregate formation [17], thus increasing carbon export to the deep ocean [134]. The importance of viruses in deeper water is also highlighted by Zhang *et al.* [135], who considered viral particles as ‘bottom-up’ agents fueling the ML in the deep ocean.

Another comprehensive study [136] examined the genomic and transcriptional responses of microbial communities to high-molecular-weight DOM addition in samples from the surface ocean. These authors observed specific resource partitioning of DOM by the bacterial species *Idiomarina* and *Alteromonas* spp. that were most highly represented at the early time points and *Methylophaga* at the final point of the experiment. Their results demonstrated a temporal succession of taxa, metabolic pathways and chemical transformations associated with high-molecular-weight DOM turnover, suggesting that the cycling of marine DOM may require a coordinated and cooperative effort between different bacterial ‘specialists’.

**CASE STUDIES OF INTERACTIONS BETWEEN BCP, ML AND MCP**

**Case Study 1: MCP dynamics associated with upwelling activities**

Jiao *et al.* [46] hypothesized that microbial activity plays a significant role in mediating the source and/or sink of CO₂ in a productive upwelling region. This hypothesis was tested by measuring multiple biogeochemical parameters at two cyclonic eddy-induced upwelling sites in the western South China Sea, which allowed the formulation of a scenario model of MCP processes under different upwelling conditions.

In the western South China Sea, satellite altimetric data identified intensification of two cold-core cyclonic eddies, CE1 (decaying) and CE2 (growing), during sample collection [46]. In the case of the decaying eddy CE1 (modeling scenario 1, Fig. 4), no phytoplankton bloom occurred and *Prochlorococcus* dominated. The small-sized non-sinking organic particles favored the transfer of energy and organic matter through the ML pathway than through the BCP. The enhanced production of labile organic carbon due to upwelled nutrients and phytoplankton growth stimulated microbial respiration (e.g. net community respiration) and decreased POC flux, which suggested that the MCP is the prevailing mechanism for carbon sequestration. In the case of a growing eddy, CE2 (modeling scenario 2, Fig. 4), the rapid growth of phytoplankton caused enhancement of POC downward export flux, where the BCP was the prevailing mechanism for carbon sequestration. Further research is needed to validate these models for general applications.

**Case Study 2: Modeling the MCP functions**

Lu *et al.* [137] made an attempt to analyse the MCP-related variables and processes using a coupled physical-ecosystem model that used data collected in the South China Sea and assumed a constant annual production of RDOC of ~0.2 Pg C for global oceans [50]. They also ran the model with different scenarios simulating rising sea-surface temperature and compared the BCP and MCP rates and their relative contributions to carbon sequestration.

The model coupled a physical model from the operational Taiwan Strait Nowcast\Forecast system [138,139] and a biogeochemistry model based on the Carbon, Silicon, Nitrogen Ecosystem module [140], which was modified to incorporate an explicit RDOC pool and the MCP processes (Fig. 5). With the constraint of a bulk RDOC concentration of 40 µM [13], and the satellite-based value of primary production, this model estimated the ratio of MCP to BCP (at the depth of 1000 m) to be 1:6.08 in the South China Sea. The annual production rate of RDOC by the MCP averaged over the whole South China Sea domain was estimated to be 1.55 mg C m⁻² d⁻¹. The BCP, on the other hand, sequestered 9.43 mg C m⁻² d⁻¹.

**FUTURE RESEARCH FOCI AND PROSPECTS**

Jiao *et al.* [9] highlighted nine major questions regarding MCP processes, which have been addressed at different levels over the past 8 years (Table 3). There is an urgent need to better understand the impacts of global-scale environmental change, including ocean warming and acidification and related deoxygenation and changes in nutrients availability on carbon cycling in the ocean [141]. A central question is how microbial processes contribute to the transformation of organic carbon in the ocean. We advocate three approaches to promote future research in this direction in accordance with Jiao *et al.* [48].

First, we recommend increased investigation of microbiomes in different natural environments, including a much better coverage of the deep ocean. These studies should integrate various omics...
Figure 4. Scenario models for the effects of upwelling on ocean carbon uptake/outgassing dynamics (adopted and modified from [46]). (1) Functioning of the BCP and the MCP in a non-upwelling region of the ocean. (2) Dominance of the MCP in scenario 1 where the total upward CO₂ flux exceeds downward POC export flux: nutrients are injected only into the lower layer of the euphotic zone; Prochlorococcus is dominant; microbial respiration is enhanced; CO₂ outgassing exceeds POC export; the MCP is the prevailing mechanism for carbon sequestration. (3) Dominance of the BCP in scenario 2 where the downward POC flux exceeds the total upward CO₂ flux: nutrients are injected into the upper layer of the euphotic zone; diatoms are dominant; POC export exceeds CO₂ outgassing; the BCP is the prevailing mechanism for carbon sequestration.

Figure 5. Schematic diagram of the MCP module (from [137]). The RDOC in the model is produced via two bacteria-related pathways: (1) direct exudation by bacteria and (2) passive release from viral lysis of microbial cells. The additional POC degradation pathway [9] is implicitly included by transforming from POC to labile/semi-labile organic carbon and then to RDOC via aforementioned two pathways (see [137] for detailed explanation).

approaches (i.e. metagenomics, metatranscriptomics, metaproteomics and metabolomics) at all levels of the microbial community (i.e. virus, bacteria, archaea, phytoplankton and zooplankton), as well as at selected time-series locations in the coastal and open ocean to identify how the metabolic capacity of the ocean’s microbiome responds to spatial and temporal changes in an environmental context (e.g. [27,133]).

The second proposed approach is to strengthen the understanding of the connections between microbial metabolism and the chemical structure of DOC compounds (e.g. [51]). Bioassays of DOC composition coupled with changes in bacterial communities can now be conducted integrating omics and FT–ICR MS and NMR technologies, which offers the potential for new insights into mechanisms responsible for the formation of RDOCt and RDOCc. In particular, efforts are needed to fully examine the fate of DOM under different trophic conditions and at the ecosystem level [53,134,142].

The third proposed approach is to establish and expand long-term incubation studies employing large-scale facilities, such as the existing Aquatron Tower Tank (Dalhousie University, Canada) and the planned Marine Environmental Chamber System (Shandong University, China) under controlled environmental conditions. Using such facilities provides a unique complement to field studies by seeking to mimic ocean-relevant physical, chemical and biological environmental conditions (e.g. vertical stratification) and their variations for long-term experiments. Such experiments are required to provide unique data and insight for testing hypotheses regarding the effects of global environmental change on the ocean carbon cycle [143,144].
We also highlight the need to examine the role of planktonic archaea in the carbon cycle. These archaea, such as *Thaumarchaeota*, have been recognized to play an important role in the ocean carbon cycle [145]. Yet, the claim made 7 years ago that ‘we are woefully unaware of DOM production (or assimilation) mechanisms in the Archaea’ [25] still holds true. The study of archaea is largely hampered by the difficulty of isolating strains from the ocean (e.g. MGII and MGIII). Hence, future efforts should include the development of new technologies for enrichment and isolation of these and other organisms, guided by genomic information [133,146].

The MCP has stimulated provocative and constructive discussions and studies on the processes and mechanisms of RDOC formation and preservation [26,47,111,112,147,148]. Increasing and synergistic efforts will continue to be made to gain further understanding of the ocean carbon cycle through an integration of the concepts of the BCP, ML, VS and MCP, particularly in the context of global ocean circulation (e.g. [111]).

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