Extraordinary slow degradation of dissolved organic carbon (DOC) in a cold marginal sea

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Dissolved organic carbon (DOC) is the largest organic carbon reservoir in the ocean, and the amount of carbon in this reservoir rivals that in atmospheric CO₂. In general, DOC introduced into the deep ocean undergoes a significant degradation over a centennial time scale (i.e., ~50 μM to ~34 μM in the North Atlantic and Mediterranean Sea). However, here we show that high concentrations of DOC (58 ± 4 μM) are maintained almost constantly over 100 years in the entire deep East/Japan Sea (EJS). The degradation rate in this sea is estimated to be 0.04 μmol C kg⁻¹ yr⁻¹, which is 2–3 times lower than that in the North Atlantic and Mediterranean Sea. Since the source of DOC in the deep EJS is found to be of marine origin on the basis of δ¹³C-DOC signatures, this slow degradation rate seems to be due to low temperature (<1 °C) in the entire deep water column. This observational result suggests that the storage capacity of DOC in the world ocean is very sensitive to global warming and slowdown of global deep-water overturning.

Most of the DOC in the ocean is produced by marine organisms and removed by abiotic processes, such as photochemical oxidation¹⁻² and microbial degradation³. DOC is a highly complex mixture of compounds⁴, each having a different reactivity. Although DOC plays an important role in global carbon cycling⁵⁻⁷, the mechanisms underlying the production and removal of DOC have yet to be clarified⁴. The objective of this study was to examine the degradation rate of DOC in a very cold deep ocean (EJS), which may infer the sensitivity of DOC storage capacity in warming oceans.

The EJS is a semi-enclosed marginal sea and is divided into three major basins (Japan Basin, Ulleung Basin, and Yamato Basin), all of which are deeper than 2000 m (Fig. 1). The deep water masses (at depths greater than 1000 m) in the EJS are formed by deep convection or brine rejection in the Japan Basin⁸⁻¹¹. Using salinity and oxygen isotope budgets⁹, the rate of potential bottom-water formation due to brine rejection (at depths greater than 2500 m) was estimated to be approximately 4 × 10¹² m⁻³ yr⁻¹. This brine rejection accounts for 25–35% of the subsurface water formed in the EJS. The turnover time of the EJS deep water mass was determined to be ~100 years by using radionuclides such as ¹⁴C and ²²⁶Ra¹²,¹³ by assuming steady-state conditions.

Results and Discussion

Vertical profiles of DOC concentration were determined on several cruises covering the main EJS during the course of three years (Fig. 1). The average DOC concentrations in the surface layer (0–200 m) and in the deep (>200 m) layer over the entire area of the EJS were 68 ± 6 and 58 ± 4 μM, respectively (Fig. 2). In general, the DOC concentrations in the surface layer of EJS were comparable to those in the major world oceans (60–80 μM)¹⁴⁻¹⁰. However, the DOC concentrations in the deep layer of the EJS are...
significantly higher than those in the major oceans (34–43 μM)⁶,¹⁷,¹⁸ (Fig. 2) and slightly higher than or similar to those in the Arctic Ocean (54 ± 3 μM, up to ~4000 m)¹⁹ and the Nordic seas (50 μM, up to ~3500 m)²⁰, where deep water formation occurs. In the Greenland Sea, one of the Nordic seas, the concentrations of DOC decrease from the surface (60 μM) to the deep layer (53 μM, <1500 m depth) and reach a constant DOC concentration (50 μM) in the deeper layer (>1500 m depth)²⁰. Therefore, the concentration of DOC in the deep bottom water of the EJS seems to be the highest reported so far in the world oceans.

In the Arctic and Nordic Seas, the elevated concentrations of DOC are related to downwelling of waters that are enriched in terrigenous DOC¹⁹,²⁰. However, there is no large river that directly drains into the EJS. In addition, export of DOC in the deep EJS to the Pacific Ocean can be ignored since the EJS is connected to the Pacific Ocean only through the sills which are shallower than 150 m. The δ¹³C-DOC in the EJS ranged from −20.4 to −21.7% (avg.: −21.3 ± 0.4%), similar to that in the North Pacific (avg.: −21.2 ± 0.2%) and in the North Atlantic (avg.: −21.0 ± 0.3%) oceans²¹ (Fig. 2). The enriched δ¹³C values in these oceans are different from terrigenous C₃ plant material (−28 to −30%) indicating that the main fraction of the DOM is of marine origin (δ¹³C values of plankton organic matter = −18 to −20%). However, we cannot exclude the possibility that the Tsushima Current, which brings Yangtze River water into the EJS, may contribute some terrigenous DOC to the EJS. In addition, the DOC production from sinking particulate organic carbon (POC) could result in high DOC in the EJS. However, despite most of the sinking POC (>80%) being generally degraded between 200 and 1600 m²², the DOC profiles were uniform throughout the entire depths, down to 3500 m, indicating that DOC production in the deep
The ocean is not significant over the time scale of deep water mixing. The total export production (including sinking particles) through 200 m in the EJS has been estimated to be 99 Tg yr$^{-1}$ (8.25 mol C m$^{-2}$ yr$^{-1}$) by using $^3$H-$^3$He isotopes$^{22}$, which in general is not unusually higher than in other oceans.

The concentrations of DOC in the deep layer of the EJS in May 2007 decreased slightly from the north ($59 \pm 3 \mu M$) to the south ($55 \pm 2 \mu M$) along the deep-sea current flows, which was associated with biological degradation reflected in apparent oxygen utilization (AOU) (Fig. 3). Considering the fact that these DOC profiles include the entire East Sea through several cruises, including the southwestern EJS stations$^{23}$, over the last 10 years, the DOC concentrations in the deep EJS were remarkably stable, and neither systematic nor significant differences were observed among the stations.

In order to determine the characteristics of DOC degradation in the EJS, it was necessary to compare DOC degradation rates for the same DOM quality and the same age following deep water formation. In this connection, the degradation rate of DOC in the deep EJS was compared with those of the Atlantic Ocean and the Mediterranean Sea, which have similar deep water formation modes (thermohaline circulation). Because the turnover times of the EJS and Mediterranean Sea are about 100 years, we used the DOC difference in the entire regions of these marginal seas. In the case of the Atlantic Ocean, we used the difference in DOC concentration between the Greenland Sea water and the water mass around 50 °N, which was found to be approximately 100 years old based on $^{14}$C dating$^{24}$.

The slopes of DOC concentrations over the time scale of 100 years were obtained from the highest (59, 50, and 48 μM) to the lowest (55, 42, and 34 μM) concentrations in the EJS, Atlantic Ocean$^{25}$, and Mediterranean Sea$^{26}$, respectively. A major fraction of the DOC mineralization was assumed to have occurred during downwelling. Then, the degradation rate of DOC was estimated to be 0.04, 0.08, and 0.14 μmol C kg$^{-1}$ yr$^{-1}$ in the EJS, Atlantic Ocean, and Mediterranean Sea respectively. The degradation rate in the EJS is much lower than those of the Mediterranean Sea and the Atlantic Ocean.

In order to determine the contribution of DOC oxidation to oxygen consumption in deep EJS, DOC is plotted against AOU. The plotting was performed after converting AOU to carbon equivalents [AOU-C$\text{eq}$ (μM C) = AOU (μM O$_2$) × 0.72]$^{16}$. A ΔC/ΔO ratio of 0.72 was derived from the Redfield stoichiometry (C:O:N:P = 106:42:16:1)$^{27}$. The slope (~0.13) of the linear regression of DOC concentration versus AOU-C$\text{eq}$ in the deep EJS indicates that oxidation of DOC is responsible for 13% of the oxygen utilization in the deep EJS. This value in the deep EJS is similar to that (14 ± 3%) in the North Atlantic Deep Water$^{25}$, but much higher than that (approximately 32%) in the eastern Mediterranean Sea$^{28}$.

The reason behind the low degradation rates remains speculative at this point, but it is most likely associated with the difference in water temperature. The water temperature in the EJS is <1 °C (Fig. 2), which is lower than that in the Atlantic Ocean (3 °C, range from 2.2–3.5°C) and in the Mediterranean.
Sea (−13 °C, range from 12.8–13.7 °C). Carlson et al. (ref. 25) suggested that the degradation rates of DOC derived from single end-member and multiple linear regression models decrease exponentially as temperature decreases although temperature cannot be the sole control of DOC degradation. DOM quality other than temperature alone might control DOM degradation rates 29–31.

The water temperature increased by 0.1–0.5 °C in the upper 1000 m and by 0.01 °C below 2000 m from the 1950s to 1996 in the EJS because of the recent slowdown in deep-water formation 32. Similarly, the rate of anthropogenic CO₂ accumulation in the deep layer of the EJS decreased considerably from 1992 to 2007 owing to the considerable weakening of the overturning circulation 33. It is unknown whether this recent small change has resulted in DOC reduction in the deep EJS during the last few decades. However, our results from EJS in comparison with other seas suggest that global carbon cycles could be significantly affected by the warming and slow-down of water overturning in marginal seas.

**Methods**

The samples for DOC and δ¹³C-DOC were filtered onboard through a pre-combusted syringe glass-fiber filter (Whatman, 0.7 μm, 25 mm) into pre-combusted glass ampoules (550 °C for 5 h). The samples were then acidified to pH 2 with 6 M HCl and stored at 4 °C for preservation until further analysis. All sampling procedures were performed in a clean bench (class 100). The DOC concentrations were measured using a TOC-V CPH analyzer (Shimadzu, Japan), as described elsewhere 34. Briefly, the acidified seawater sample was bubbled with high-purity air gas (purity: 99.999) to completely purge inorganic carbon species in the injection system. Carrier gas was passed at a controlled flow rate of 150 mL min⁻¹. Three to five replicates (100 μL) for each sample were automatically injected into a combustion glass tube filled with a catalyst (Pt-coated Al) at 720 °C in a furnace. The organic matter present in the sample was oxidized on the catalyst to CO₂. The evolved CO₂ was measured by non-dispersive infrared detection (NDIR) by integrating peak area.

In order to determine DOC concentrations with high accuracy, it is necessary to reduce the system blank as low as possible. The blank run was continued until organic-free distilled water (DIW) was stable within the detection limit (<5 μM) of our DOC method. Six-point calibration curves for acetic anilide were used daily for DOC standardization. The reliability of the measurements was verified on a daily basis by comparing the measured values with a DOC-certified seawater sample (DSR: 44–46 μM for DOC, University of Miami) and procedural blanks. Most of our DSR measurement results (n = 34, 92%) were in good agreement with the DSR values (within 2%), although three measured values differed by 3–5% from the DSR values. These DSR measurements were performed for every 10 samples. The average standard deviation value for all DSR measurements was ±1.2 μM, which is similar to the value.
carbon (±1 μM) suggested by Carlson et al. (ref. 25). An inter-laboratory comparison between the two laboratories involved (Kim and Dittmar) confirmed the accuracy of measurements to be within 5%.

The stable isotope compositions of dissolved organic carbon (δ13C-DOC) were measured by using TOC-IR-MS (Isoprime IR-MS coupled with vario TOC cube), which is commercially available (Isoprime, Elementar). This instrument uses the common high temperature catalytic combustion method, which is the same principle as our DOC method. Briefly, 10 mL of filtered samples, following acidification to pH 2 with 6 M HCl, is placed into an auto-sampler. The instrument is programmed to purge the sample for 20–30 min with O2 gas to completely remove dissolved inorganic carbon species. Then, 1.5 mL of the sample is injected into Pt-impregnated silica spheres in a quartz tube, where DOC is fully converted to CO2 at 750 °C. The CO2 gas flows through a halogen trap and a water trap. After detecting DOC concentrations using an NDIR detector, CO2 gas enters the interface by O2 carrier gas. In the interface, a series of valves are set so that the flow from the TOC and the flow toward the IR-MS system can be separated. The CO2 gas, carried by O2 carrier gas, is trapped by CO2 column in the interface. After switching the valves, the trapped gas is released to IR-MS by helium carrier gas. After passing through a reduction furnace to remove any interfering species, including residual oxygen, CO2 gas is transferred to the IR-MS. The δ13C content of the CO2 is measured using Isoprime IR-MS.

The ion source parameters were fine-tuned every day, and the variations of the reference gas were checked to maintain good stability (standard deviation: ±0.01%). The blank was determined by running low carbon water (University of Miami), containing a DOC level lower than 2 μM before the analysis of each sample batch. Certified IAEA-CH6 sucrose (International Atomic Energy Agency, −10.45 ± 0.03%) dissolved in the low carbon water was used for standardization. For every 10 samples, one standard sample was measured to check the stability of measurements. The method used for blank determination and corrections followed the previous method. Our measurement result of δ13C-DOC for DSR was −21.5 ± 0.1%, which agree well with previous results reported by Lang et al. (ref. 37) (−21.7%) and by Panetta et al. (ref. 35) (−21.4 ± 0.3%). The samples were measured three times, and the values were reproducible within 0.3%.

At all the stations, dissolved oxygen (DO), potential temperature, and salinity were measured using a rosette system with a mounted CTD (SBE 911+). All DO data were from a CTD–DO sensor that was calibrated by the data from grab samples (measured on board using the Winkler titration method). AOU, determined from the same bottle as DOC concentration, was obtained by subtracting the measured DO concentration from the saturated DO concentration which is dependent on temperature and salinity. The temperature sensor used was a SBE–3/F thermometer (resolution 0.0003 °C). The probes were calibrated by the data from grab samples (measured on board using the Winkler titration method). Reported error margins were standard errors. Wherever appropriate, error propagation was applied.

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This study is part of T.H.'s Ph.D. thesis who performed sampling, chemical analysis and modeling under the supervision of G.K. and T.D., S.A. measured δ13C-DOC and wrote the method section for δ13C-DOC. The manuscript was written in a joint effort between the four authors.

Additional Information
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