

Water masses in the Atlantic Ocean: water mass ages and ventilation

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Abstract: The distribution of water masses and their characteristics, including ventilation, provides fundamental insights into large-scale oceanographic processes such as thermohaline circulation and marine biogeochemical cycles. The characteristics of main water masses in the Atlantic Ocean have been comprehensively documented in a companion study (Liu and Tanhua, 2021); this study presents quantitative assessments of water mass age characteristics and ventilation time scales through an analysis using the transient tracers chlorofluorocarbon-12 (CFC-12), sulfur hexafluoride (SF₆), and argon-39 (³⁹Ar). We use two distinct age concepts: mean-age as an integrative metric of water mass chronology, and mode-age as a proxy for advective time scales. Under our fixed transit-time distribution shape ($\Delta/\Gamma = 1$), the two ages are linearly related (mode-age $\approx 0.162 \cdot$ mean-age). Here we mainly report mode-ages; the corresponding mean-ages can be obtained by dividing with 0.162. Empirical results demonstrate systematic age progression with increasing depth and along water mass trajectories. Surface layer central waters exhibit mode ages up to ~ 30 years (mean-age ~ 100 years). In the intermediate layer, meridional age gradients characterize the Antarctic Intermediate Water (AAIW) reaching maximum mode-age ~ 80 years (mean-age ~ 300 years) at 30°N , whereas zonal variations manifest in Mediterranean Water (MW) with peak mode-ages ~ 100 years (mean-age ~ 400 years) observed in equatorial regions. As the dominant deep water component, North Atlantic Deep Water (NADW) exhibits extreme ages in the Antarctic Circumpolar Current (ACC) region at 50°S , achieving mode-age ~ 100 years (mean-age ~ 600 years). Bottom layer water masses display their oldest signatures: Antarctic Bottom Water (AABW) from the Weddell Sea reaches mode-age ~ 100 years (mean-age ~ 600 years) at equatorial latitudes, while its extension, Northeast Atlantic Bottom Water (NEABW), attains exceptional values of mode-age ~ 120 years (mean-age ~ 800 years) at 50°N . The age analysis reveals significant basin-scale asymmetries, with western basins exhibiting younger ages compared to eastern counterparts. Ventilation efficiency modulates these age distributions, as evidenced by lower mode-ages and reduced apparent oxygen utilization (AOU) in better-ventilated western basins. The calculated oxygen utilization rate (OUR) demonstrates spatial concordance with dissolved oxygen (DO) concentrations, corroborating enhanced oxidative processes in high-oxygen regimes. This integrated age framework provides novel insights into water mass ventilation dynamics and their biogeochemical implications through quantitative characterization of temporal-spatial age distributions across multiple oceanographic provinces.

Key words: Water Mass, Atlantic Ocean, Transient Tracer, Mean- and Mode-age, Ventilation, GLODAPv2 data product

31 1 Introduction

32 The Atlantic Ocean is composed by a complex vertical stratification of water masses characterized by differences in properties
33 such as salinity, temperature and oxygen. Liu and Tanhua (2021) have systematically characterized 16 main water masses that
34 have been extensively considered and studied due to their significant impacts on ocean circulation patterns and global climate
35 regulation (e.g., Hall & Bryden, 1982; Bryden et al., 2005; Kuhlbrodt et al., 2007; Clark et al., 2012). Ocean ventilation
36 constitutes a fundamental thermohaline process responsible for the vertical displacement of surface waters into the deep ocean,
37 effectively redistributing surface heat and salinity anomalies through density-driven stratification. This process inherently
38 obeys the principle of mass conservation, simultaneously enabling the upwelling of deep-sea waters to the surface ocean. The
39 resultant vertical exchange plays a critical role in maintaining global heat distribution patterns and sustaining the
40 thermodynamic equilibrium of the ocean system (Talley, 2013; Armour et al., 2016; Talley et al., 2016). Beyond its
41 thermodynamic significance, ocean ventilation emerges as a pivotal mechanism governing marine biogeochemical cycles. By
42 mediating the downward transport of atmospheric oxygen and carbon dioxide to abyssal depths, this process fundamentally
43 regulates dissolved oxygen concentrations and carbon sequestration rates in the deep ocean (Tanhua et al., 2006; Ziska et al.,
44 2013; Skinner et al., 2017). Related studies have further elucidated ventilation's deterministic role in controlling abyssal redox
45 conditions and organic carbon remineralization processes (van Heuven et al., 2011; Ito et al., 2016; Schmidt et al., 2017).

46 Early studies of the water mass ventilation are proposed by Sandström already in the early 20th century (Sandström, 1908;
47 1916). Sverdrup and Bjerknes also reported the impacts of ventilation and currents on the climate through the air-sea
48 interactions (Sverdrup, 1940; Bjerknes, 1964). Hereafter many studies have been focusing on the ventilation. For instance,
49 Dickson et al., (1994) estimated the formation and pathway of the North Atlantic Deep Water (NADW) based on the
50 hydrodynamics, Orsi et al., (1999) investigated the circulation of the Antarctic Bottom Water (AABW) by using
51 Chlorofluorocarbons (CFCs). Recent studies suggest that the intensity of ocean ventilation varies with environmental factors,
52 such as intensity of the westerly winds (e.g. Rahmstorf, 2010; Purkey et al., 2010; Morrison et al., 2015). Several studies
53 investigate the regional scale or specific water masses, for instance, the upwelling of Circumpolar Deep Water (CDW) in the
54 Southern Ocean (Tamsitt et al., 2017), or the ventilation in the South China Sea (Wang et al., 2021). In parallel, advances in
55 ocean modeling have provided new insights into water mass transformation and tracer transport on basin to global scales,
56 offering valuable perspectives on the climatic drivers of ventilation variability (e.g., Li et al., 2023). However, few studies
57 look at the ventilation over ocean basin scales, which is helpful to understand the effect of ocean-climate interactions, and in
58 addition can provide a basis for the biogeochemical studies.

59 The water mass age is an important parameter to evaluate the ventilation and refers to as the elapsed time on the pathway. The
60 CFCs and Sulfur hexafluoride (SF_6) are recognized as effective transient tracers for water masses and to estimate their ages,
61 i.e. the time since the water left the surface ocean (Fine, 2011). The concentrations of CFC-12 in surface sea water increased
62 continuously following the increase of the atmospheric concentration since their introduction in the early 20th century
63 (Gammon et al., 1982; Warner and Weiss, 1985) up to about the turn of the century. The use of CFCs as a tracer in the

64 oceanography is well established (e.g. Bullister and Weiss, 1983; Doney and Bullister, 1992; Fine, 2010; 2011), and transient
65 tracers is a core variable in the GO-SHIP repeat hydrography program. Similarly, the transient tracer is an Essential Ocean
66 Variable (EOV) in the Global Ocean Observing System (GOOS) framework. SF₆ has been used as a new tracer since mid-
67 1990s complementing the CFCs due to the reduction of CFCs in the atmosphere (Maiss et al., 1996; Bullister et al., 2002). The
68 application of SF₆ is generally appropriate for recently formed water masses in the upper ocean (Tanhua et al., 2008), i.e. for
69 well ventilated, or young, waters. The CFCs are used to trace the relative deeper water masses (before mid-1990s or partial
70 pressure of CFC-12 lower than about 450 parts per trillion, ppt), while the SF₆ is the better choice for recently formed shallow
71 water masses (after mid-1990s or partial pressure of CFC-12 higher than about 450 ppt, Tanhua et al., 2008). The
72 complementary nature of CFCs and SF₆ allows for more accurate tracing of water masses (Law and Watson, 2001; Vollmer
73 and Weiss, 2002; Watanabe et al., 2003; Tanhua et al., 2004, 2005; Bullister et al., 2006). In addition, the application of
74 transient tracers also provides support to the hydrographic and biogeochemical field in calculating the upwelling velocity
75 (Tanhua and Liu, 2015) or estimating the ventilation and anthropogenic carbon cycle (van Heuven et al., 2011; Tanhua et al.,
76 2013; Patara et al., 2021).

77 As two typical representatives of transient tracers, the CFC-12 and SF₆ are commonly employed for determining water mass
78 ages, verifying hydrological models, and estimating rates of marine biogeochemical processes, such as anthropogenic carbon
79 storage and oxygen consumption. They exhibit advantages in tracing upper and middle water masses with relatively younger
80 ages. However, challenges arise when attempting to obtain data from deep and bottom layers with old waters. The chemically
81 inert isotope argon-39 (³⁹Ar), which has a half-life of 269 years, addresses this gap. ³⁹Ar is particularly advantageous for tracing
82 deep and bottom water masses with ages exceeding 500 years (Broecker & Peng, 2000; Holzer & Primeau, 2010; Lu et al.,
83 2014). Initially, its application was constrained by the requirement for large sampling volumes—typically around 1000 liters—
84 and the limitations of low-level counting (LLC) detection methods (Loosli et al., 1983; Schlosser et al., 1994; Schlitzer et al.,
85 1985). Consequently, ³⁹Ar was not widely utilized in oceanographic tracer studies. With advancements in Atom Trap Trace
86 Analysis (ATTA) technology (Chen et al., 1999; Jiang et al., 2012), ³⁹Ar has gradually gained prominence as an effective tracer
87 for investigating deep and bottom waters (Lu et al., 2014; Ebser et al., 2018), the accuracy and stability have also been
88 significantly improved (Jia et al., 2025). Therefore, the multi-parameter combination of CFCs, SF₆, and ³⁹Ar can transcend the
89 temporal limitations associated with single-tracer dating methodologies while providing more comprehensive insights into
90 biogeochemical processes. In this study, we introduce ³⁹Ar to investigate water mass ages within the Atlantic Ocean, in
91 particular for the older waters.

92 The apparent oxygen utilization (AOU, $\mu\text{mol kg}^{-1}$) is an important indicator in characterizing the respiration of organic matter
93 consuming oxygen, and balancing the oxygen supply through ventilation and photosynthesis. The oxygen utilization rate (OUR,
94 $\mu\text{mol kg}^{-1} \text{ yr}^{-1}$), which can be calculated from AOU and water mass age, is a significant parameter in estimating the
95 biogeochemical processes such as primary productivity or remineralization (e.g. Jenkins, 1982; Bender, 1990). The OUR has
96 been recognized to be a function of pressure for a long time (e.g. Tseitlin, 1992). However, the OUR is the integrated oxygen

consumption rate along the pathway of a water mass and represents a regional view of export flux (e.g. Stanley et al., 2012; Koeve and Kähler, 2016), and thus an imperfect measure of local OUR. In this study, the OUR is calculated from water mass ages estimated from the CFC-12 and SF₆. The change of OUR over time in different water masses are presented and the impacts from the currents and topography are discussed.

In the study by Liu and Tanhua, (2021), the characteristics of main water masses in the Atlantic Ocean are defined by six key properties, using the Global Ocean Data Analysis Project version 2 the annual update 2023 (GLODAPv2.2023, Lauvset et al., 2016; Olsen et al., 2016; Lauvset et al., 2024) data product as a source of unbiased data. The static distribution of the water masses is estimated with the Optimal Multi-Parameter analysis (OMP analysis, Karstensen and Tomczak, 1997; 1998). As a continuation of that work, here we estimate the ventilation of water masses in the Atlantic Ocean with the combination of the above mentioned three transient tracers (CFC-12, SF₆ and ³⁹Ar). The water mass ages are estimated and the OURs are calculated. The goal of this study is to further improve the report of water masses in the Atlantic Ocean by adding the age as a measure of ventilation, and to provide a hydrographic assistant for the biogeochemical researches.

In this article, we start in Section 2 with an overview of the data sources and methodologies employed. Section 3 presents the core results, illustrating the spatial distributions of mean and mode ages for the principal Atlantic Ocean water masses, stratified from surface to bottom. In Section 4, the derived water mass ages are applied to estimate Oxygen Utilization Rates (OUR) and to examine associated biogeochemical implications. Section 5 offers a comparative analysis of water mass ages obtained from traditional transient tracers (CFC-12 and SF₆) and the radioisotope ³⁹Ar. Finally, Section 6 provides a comprehensive summary of conclusions, discusses the broader relevance of the findings, and acknowledges the limitations of the present study.

2 Data and methods

2.1 Transit Time Distribution (TTD) and the Inverse Gaussian Model

To quantify the ventilation timescales of water masses, we apply the Transit Time Distribution (TTD) framework. The TTD, denoted as $G(\tau)$, represents the distribution of transit times τ that water parcels take to travel from the surface formation region to an interior observation point (Hall and Plumb, 1994; Waugh et al., 2003). The concentration of a transient tracer in the ocean interior, $c(t)$, can be modeled as the convolution of its time-evolving atmospheric source function, $c_0(t)$, with the TTD:

$$c(t) = \int_0^\infty c_0(t - \tau) G(\tau) d\tau$$

A common and practical parameterization of the TTD is the one-parameter Inverse Gaussian (IG) distribution (Waugh et al., 2003, 2004), which represents a solution to the one-dimensional advection-diffusion equation for a steady-state flow. The IG-TTD is defined as:

$$G(\tau) = \sqrt{\frac{\Gamma^3}{4\pi\Delta^2\tau^3}} \exp\left(-\frac{\Gamma(\tau-\Gamma)^2}{4\Delta^2\tau}\right);$$

126 Where:

- 127 • τ is the transit time (in years);
- 128 • Γ is the mean-age (in years), representing the first moment of the TTD: $\Gamma = \int_0^\infty \tau G(\tau) d\tau$, which signifies the average
129 time elapsed since the water was last in contact with the atmosphere;
- 130 • Δ is the width (in years) of the TTD, which quantifies the spread of transit times around the mean age due to mixing and
131 dispersion processes.

132 The mixing ratio, defined as (Δ/Γ) , unitless), indicates the relative importance of mixing versus advection in the transport. A
133 low ratio ($\ll 1$) suggests advection-dominated transport with a narrow range of transit times, while a high ratio ($\gg 1$) indicates
134 strong mixing and a wide range of transit times.

135 The mixing ratio, defined as Δ/Γ (unitless) — note: this is a shape parameter of the TTD and is not to be confused with
136 atmospheric tracer mixing ratios (e.g., the concentration ratio of a gas in air) — indicates the relative importance of mixing
137 versus advection in the transport. A low ratio ($\ll 1$) suggests advection-dominated transport with a narrow range of transit
138 times, while a high ratio ($\gg 1$) indicates strong mixing and a wide range of transit times.

139 While other forms of TTD, for instance, a two-IG distribution to characterize a mix of two water masses with different TTs
140 was used by Stöven and Tanhua (2014) for the Mediterranean Sea, or non-steady-state circulations may better represent reality
141 in certain regions, the one-parameter IG-TTD remains a widely adopted and practical model. This is particularly true given
142 the typical scarcity of multiple, independent tracer observations needed to constrain more complex distributions across large
143 ocean basins. In this study, we apply the one-parameter IG-TTD framework as a robust approach for basin-scale analysis.

144 **2.2 Definitions of Water Mass Age: Tracer-age, Mean-age, and Mode-age**

145 The fundamental concept of water mass age is the time elapsed since a water parcel was last in equilibrium with the atmosphere
146 at the surface (Thiele and Sarmiento, 1990). From the TTD framework, we derive and utilize three specific age definitions,
147 each with its own physical interpretation and application (see conceptual diagram in Fig. 1).

148 **Tracer-age** is the simplest age estimate, obtained by matching the observed tracer partial pressure in a water sample directly
149 to the historical atmospheric record (Fig. 1 a). It assumes the transport along a purely advective pathway (i.e., no mixing),
150 corresponding to a TTD that approaches a delta function as $\Delta/\Gamma \rightarrow 0$. This concept systematically underestimates the “true”
151 age in the presence of mixing, as it ignores the distribution of transit times/pathways (Sonnerup et al., 2001).

152 **Mean-age (Γ)** is the first moment of the TTD. The mean-age represents the average value of the transport times of all water
153 parcel and represents an integrative timescale (Fig. 1 b), particularly applicable to the study of biogeochemical accumulation
154 processes, such as calculating integrated oxygen consumption rates (e.g., Jenkins, 1987; Sonnerup et al., 2013).

155 **Mode-age (τ_{mode})** is the transit time at which the TTD, $G(t)$, reaches its maximum (Fig. 1 b). For the IG-TTD, it is calculated
 156 as $\tau_{\text{mode}} = \Gamma \cdot \left(\sqrt{1 + (3 \Delta/\Gamma)^2} - 3 \Delta/\Gamma \right)$, with the mixing ratio fixed at $\Delta/\Gamma = 1$ in this study, this simplifies to $\tau_{\text{mode}} =$
 157 $\Gamma \cdot (\sqrt{10} - 3) \approx 0.162 \cdot \Gamma$. The mode-age represents the most probable transit time of the dominant water fraction in the
 158 sample, making it a better proxy for the advective timescale along the core pathway of a water mass.

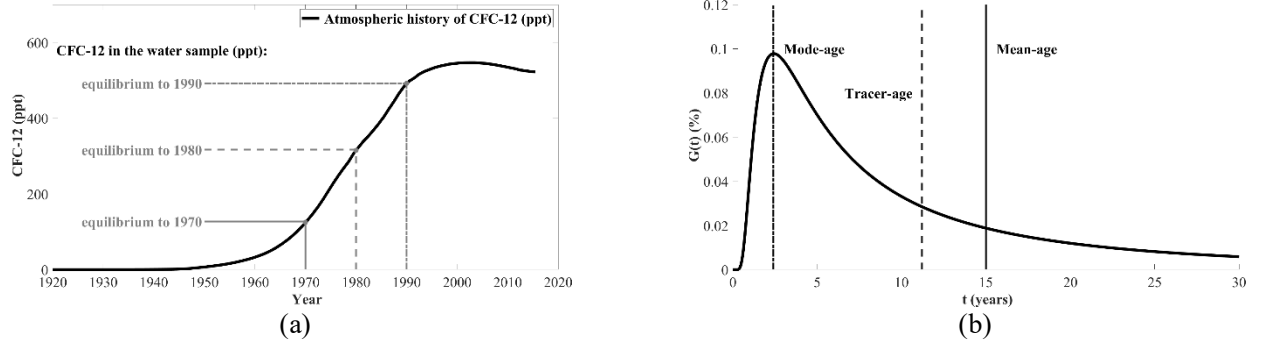


Figure 1. Schematic diagram illustrating three concepts of water mass ages.

- (a) The tracer age: The transient tracer (such as CFC-12) in the water sample in equilibrium with the atmospheric at a certain year. This approach assumes that the ocean is a purely advective process (without mixing);
- (b) The concepts of mean- and mode-ages: Based on the inverse Gaussian transport time distribution model, assuming the mixing ratio ($\Delta/\Gamma = 1$) and observed tracer partial pressure CFC-12 = 300 ppt, as an example. Shown are the positions of the mean-age (Γ), tracer age, and the mode-age (τ_{mode}).

159 To calculate tracer partial pressures from measured concentrations, the saturation of transient tracers (CFC-12 and SF₆) in
 160 surface seawater must be determined. This saturation is a function of potential temperature (θ) and practical salinity (S),
 161 following established solubility relationships (Warner and Weiss, 1985; Bullister et al., 2002).

162 2.3 Determination of the Mixing Ratio and age calculation procedure

163 A critical step in applying the IG-TTD is specifying the shape parameter, the mixing ratio (Δ/Γ). The atmospheric
 164 concentrations of these tracers provide the essential input for age calculations: the CFC-12 increased monotonically until mid-
 165 1990s, while the SF₆ is still increasing (Fig. 2a). Ideally, the mixing ratio is determined by fitting the TTD model to observations
 166 of two or more transient tracers with different input histories. However, the similar atmospheric growth curves of CFC-12 and
 167 SF₆ often preclude a robust, independent determination of Δ/Γ across large ocean basins (Waugh et al., 2004), and the short
 168 atmospheric history of SF₆ reduce this method to recently ventilated waters. Furthermore, the limited number of collocated
 169 transient tracer observations restricts the application of more complex models. In this context, understanding how the mixing
 170 ratio influences age estimates is crucial. For instance, under given tracer partial pressure (e.g., CFC-12 = 300 ppt) and sampling
 171 year (e.g., 1990), an increase in the Δ/Γ ratio of the IG-TTD leads to a corresponding increase in the calculated mean-age,
 172 while the mode-age decreases. This behavior arises from the shape of the Inverse Gaussian distribution: a higher mixing ratio
 173 extends the distribution's tail, elevating the mean-age, while simultaneously shifting its peak earlier, thereby reducing the

mode-age (Fig. 2b). Observational evidence from hydrographic sections further supports this theoretical relationship, demonstrating how the resulting age fields vary systematically under different assumed mixing ratios (Fig. 3)

To ensure a consistent and widely comparable analysis across the whole Atlantic Ocean, we adopt the standard mixing ratio $\Delta/\Gamma = 1$. This value represents a balance between advective and diffusive processes and is commonly used in large-scale oceanographic studies (Vaugh et al., 2004; Stanley et al., 2012; Stöven and Tanhua, 2014; Thomas et al., 2020). We acknowledge this as a limitation but note it is a necessary and widely accepted approach for basin-scale analyses where tracer data are limited.

The mean-age (Γ) and mode-age (τ_{mode}) for each sample was calculated by finding the value that minimizes the difference between the observed tracer partial pressures and those predicted by convolving the IG-TTD (with $\Delta/\Gamma = 1$) with the atmospheric histories of CFC-12 and SF₆. The atmospheric histories for the Northern Hemisphere were taken from Bullister et al., (2002). Where both CFC-12 and SF₆ data were available, they were used jointly in the inversion, but a switch criterion based on the observed CFC-12 partial pressure was applied. Following Tanhua et al. (2008), we note that the atmospheric growth rate of CFC-12 slowed down significantly around the turn of the century, making it less reliable as a tracer for recently ventilated waters. Consequently, when the in-situ CFC-12 partial pressure exceeded ~450 ppt (indicative of young, well-ventilated waters), the inversion preferentially relied on SF₆, whose atmospheric concentration continues to increase. Conversely, when the CFC-12 partial pressure fell below ~450 ppt (typical of older, deeper waters), CFC-12 was used as the primary tracer, owing to its longer atmospheric history and higher signal-to-noise ratio at low concentrations. This threshold-based selection ensures that the most appropriate tracer is employed for each water mass age regime. The corresponding mode-age was then calculated from the derived mean-age and the fixed mixing ratio. For the radioactive tracer ³⁹Ar, the convolution integral incorporates an additional exponential decay term, $e^{-\lambda t}$, where λ is the decay constant corresponding to its 269-year half-life.

Because this study adopts a fixed mixing ratio of $\Delta/\Gamma = 1$, the mean-age and mode-age become linearly related with specifically $\tau_{mode} \approx 0.162 \cdot \Gamma$. Under this fixed ratio, the spatial patterns of the two ages are identical, differing only by a constant scale factor. To maintain logical consistency with the OUR calculations (Section 4), which use the mode-age, we present the mode-age distributions in the main figures (Figs. 5–13, 15, 17). The corresponding mean-age sections are provided in the Supplement (Figs. S1–S9) for readers interested in the integrative timescale.

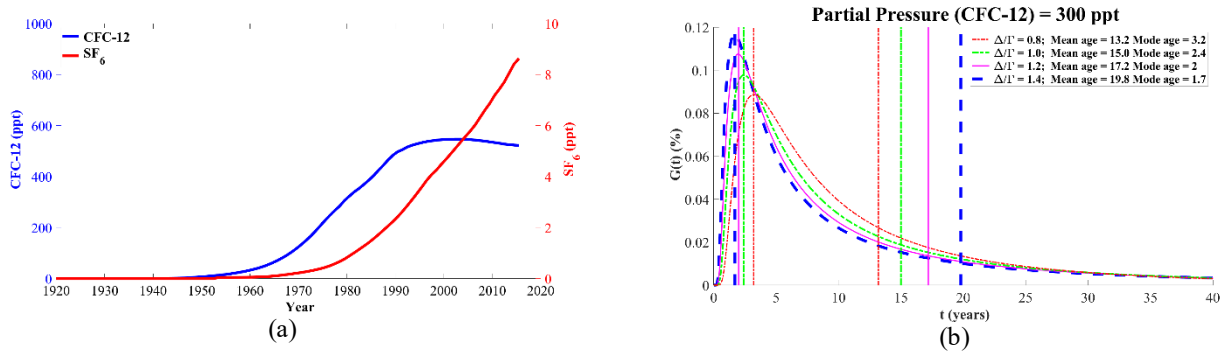


Figure 2. The relationships between mean-age, mode-age and mixing ratio (Δ/Γ).

(a) Historical atmospheric partial pressure data of transient tracers over time (CFC-12 and SF₆) in the Northern Hemisphere (data source: Bullister, 2002);

(b) Relationship between mean-ages and mode-ages under same partial pressure (CFC-12 = 300 ppt) but different mixing ratios (Δ/Γ) assuming the sampling year is 1990.

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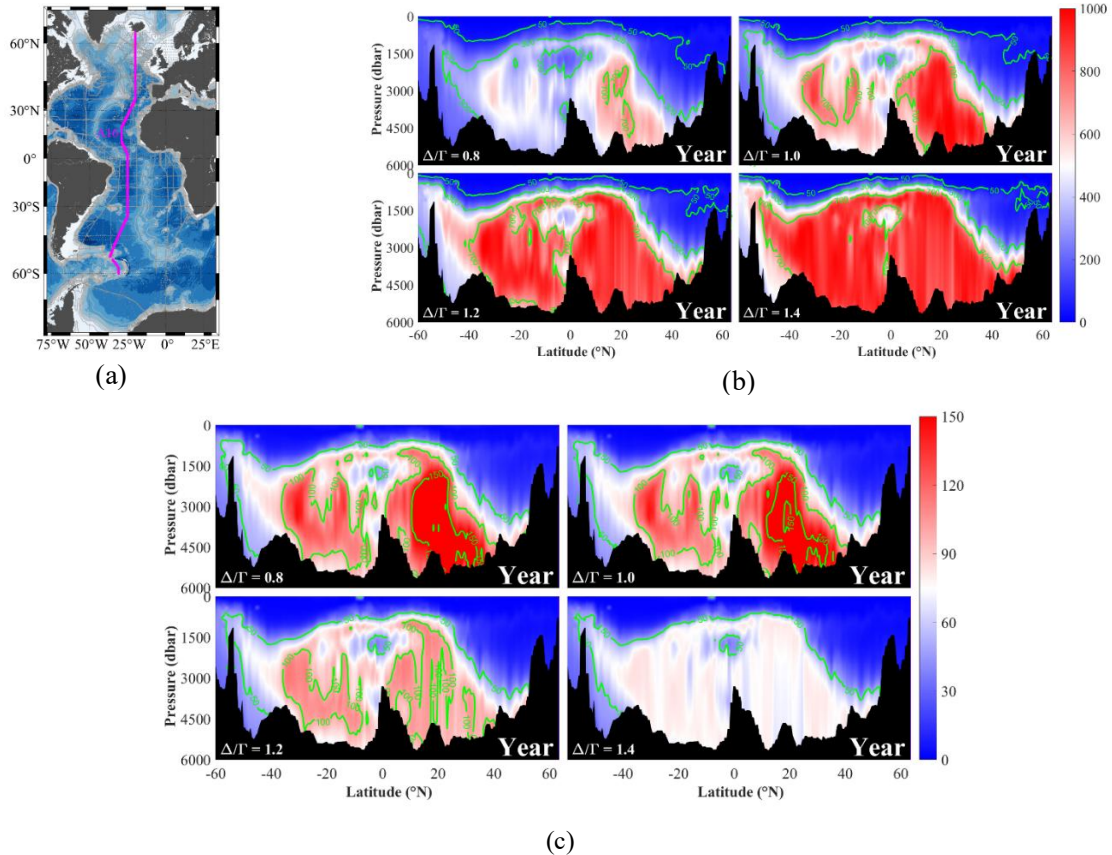


Figure 3. (a) Map of the A16 section under the framework of the WOCE/GO-SHIP project in 2013.

(b) Distributions of mean-ages and (c) mode-ages under varying mixing ratios ($\Delta/\Gamma = 0.8, 1.0, 1.2, \text{ and } 1.4$) derived from GLODAPv2 observational data. Green contour lines correspond to mean-ages of 50, 300, and 700 years and mode-ages of 50, 100, and 150 years, respectively.

201 3 Results: Ages of water masses in the Atlantic Ocean

202 In this study, we follow the division of vertical layers and distributions of water masses presented by Liu and Tanhua (2021),
 203 but now focus on the transient tracers and water mass ages. Three hydrographic sections are selected from the WOCE/GO-
 204 SHIP sections in this work to represent the mode-ages of the main water masses (Table 1). The A16 sections in 2013 (Expo-
 205 code 33RO20130803 and 33RO20131223) show the meridional distribution across the whole Atlantic Ocean, while the A05
 206 section in 2010 (Expo-code 74DI20100106) and A10 section in 2011 (Expo-code 33RO20110926) show the zonal distributions
 207 in the North and South Atlantic Ocean respectively (Fig. 4). The partial pressures of CFC-12 and SF₆ along the above sections
 208 are shown in Fig. 4 (Only CFC-12 data are displayed along A05 section due to the lack of SF₆ data in this section). Both tracers
 209 show similar distributions with high values in the shallow layers, and a general decrease with increasing pressure. As detailed
 210 in Section 2.3, under our fixed mixing ratio ($\Delta/\Gamma = 1$) the mean- and mode-ages are linearly related; we therefore show only
 211 mode-age sections in the following figures (mean-age sections are provided in the Supplements).

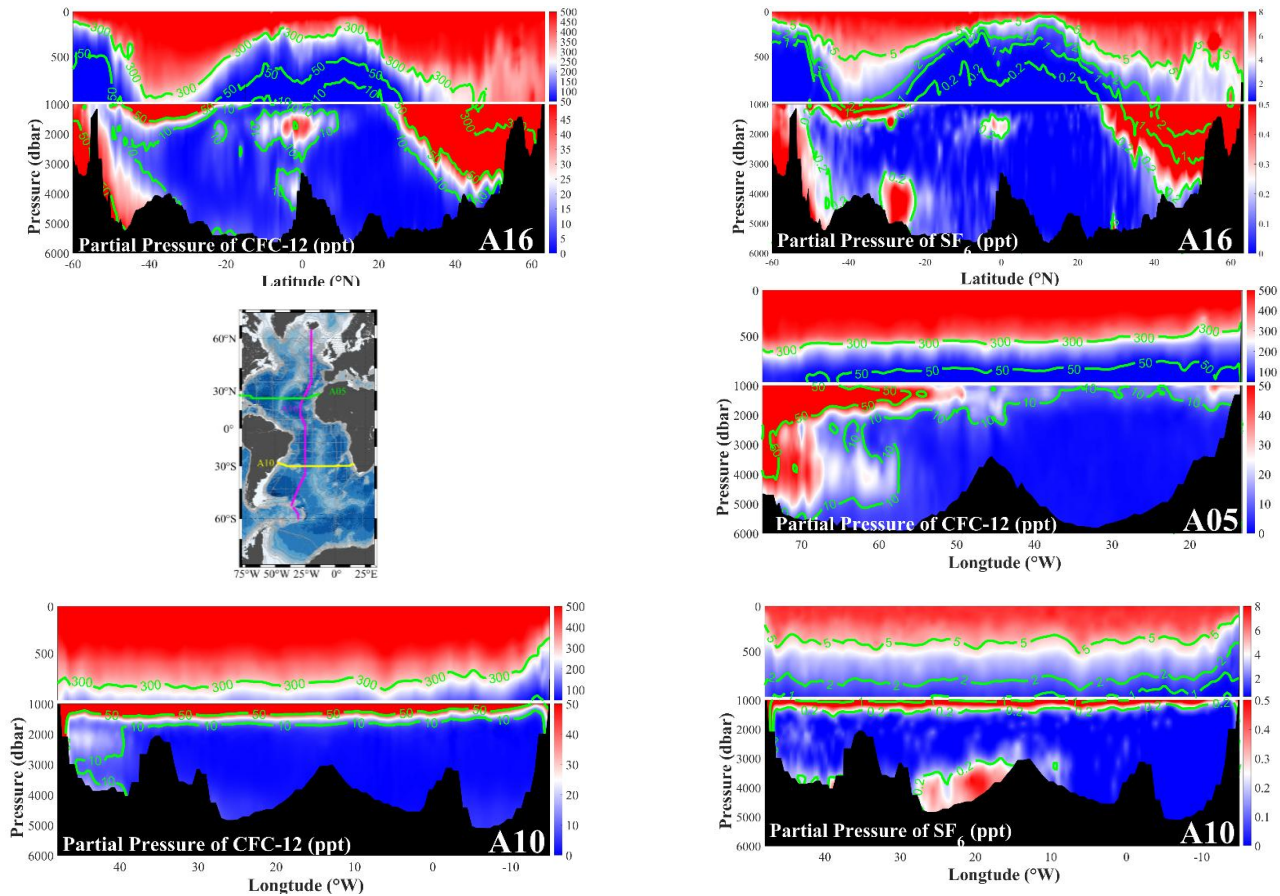


Figure 4. Map of three selected WOCE/GO-SHIP sections to represent the ages of the main water masses in the Atlantic Ocean (middle left) and partial pressures of CFC-12 and SF₆ along the three selected sections. Note the different colorbars between upper and lower parts divided at pressure of 1000 dbar. The green contour lines indicate the partial pressures of CFC-12 at 10, 50 and 300 ppt and SF₆ at 0.2, 1, 2 and 5 ppt, respectively.

213 **Table 1:** Summary of hydrographic cruises in the Atlantic Ocean selected for the oceanographic sections in this study

WOCE/GO-SHIP section	Year	Expo-code	Ship	Chief Scientist	PI of transient tracers
A16N	2013	33RO20130803	Ronald H. Brown	John Bullister & Molly Baringer	John Bullister
A16S	2013	33RO20131223	Ronald H. Brown	Rik Wanninkhof	John Bullister
A05	2010	74DI20100106	Discovery	Brian King	Marie-José Messias
A10	2011	33RO20110926	Ronald H. Brown	Molly Baringer	John Bullister

214 The upper water layers have the lowest mode-ages within ~30 years (Fig. 5), the surface water has, by definition, zero age.
 215 The distributions of mode-ages also show spatial differences in the horizontal direction. Relatively low ages are found in the
 216 high latitude regions. This indicates that the deep water masses are newly formed here, and thus have low ages. In the region
 217 between 20 and 40 °N, the mode-age reaches the peak value up to ~150 years, suggesting the sluggish water exchange and
 218 long residence time of old water masses (Fig. 5 a). In the zonal direction, the mode-ages are significantly lower in the west in
 219 the general region below 1500m dbar, suggesting that these regions are better ventilated due to the western boundary current
 220 (Fig. 5 b and c).

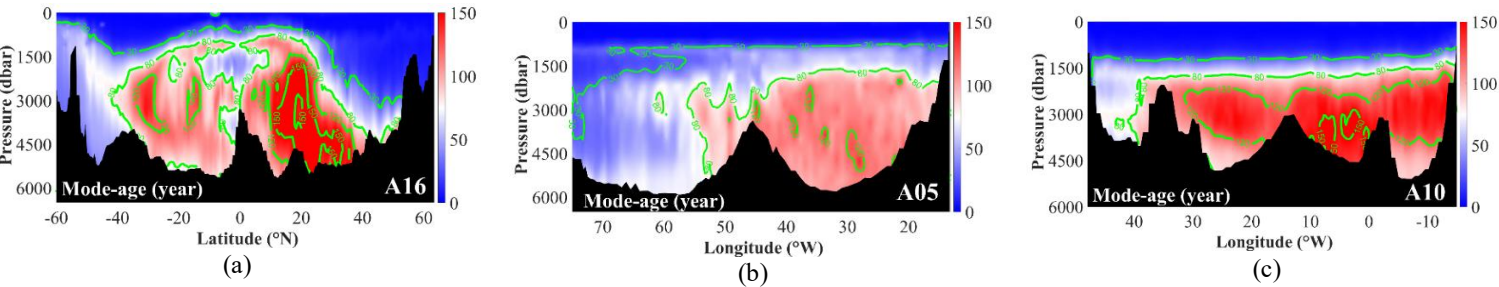


Figure 5. Mode-ages along the three selected WOCE/GO-SHIP sections

221 In the analysis of water mass ages in each layer, maps with mode-ages in each station are firstly presented as a qualitative
 222 overview. The average values of the mode-ages at the core pressure of each water mass are plotted at certain station. In addition,
 223 the quantitative calculations are made along the three selected sections to represent the distributions of water mass ages in
 224 detail and estimate the impacts from currents and mixing.

225 3.1 The upper layer

226 The spreading of central waters can be traced by the distributions of ages. After leaving their formation areas in the mid-
227 latitude regions, the West North Atlantic Central Water (WNACW) and West South Atlantic Central Water (WSACW) spread
228 generally in zonal direction towards the formation areas of the East North Atlantic Central Water (ENACW) and East South
229 Atlantic Central Water (ESACW) with Azores Current and South Atlantic Current, respectively (Fig. 6, a). The mode-ages
230 show that the “mixed” age of all the compositions in the western central waters is ~8 years (Fig. 6, b and Fig. 7). When focusing
231 on the main body of WNACW and WSACW, the mode-ages show that the advective time-scale is ~15 years for both water
232 masses (i.e. Both WNACW and WSACW take ~15 years from the formation area to the equator, Fig. 6, b and Fig. 7).

233 At greater depths, the ENACW and ESACW are transported from their formation areas and spread with the main currents
234 towards the equator (Fig. 6, a). In the northern hemisphere, the ENACW spreads southward with the Canary Current, while
235 the ESACW in the southern hemisphere is transported northward with the Benguela Current and the South Equatorial Current.
236 The difference in mode-age is ~16 years between the formation areas and the equatorial region (Fig. 6, b and Fig. 7). The
237 mode-ages indicate that the main body of ENACW and ESACW takes approximately 30 years to be transported from formation
238 areas to the equator (Fig. 6, b and Fig. 7).

239 The central waters occupy the upper layer above the neutral density isoline of 27.10 kg m^{-3} . The core pressures of western and
240 eastern central waters are around the neutral density (γ) of 26.5 and 26.9 kg m^{-3} respectively (Liu and Tanhua, 2021). The ages
241 of central waters at their core neutral densities are shown in Fig. 6. The eastern central waters have mode-ages up to
242 approximately 30 years, while the western central waters are younger with mode-ages of ~15 years (Fig. 6). The relatively low
243 ages in the upper layer indicate that these water masses are newly formed.

244 In the meridional direction along the A16 section, the mode-ages show low values of around 20 years in the mid-latitude
245 regions that are close to the formation areas of central waters (Fig. 6 and Fig. 7). In contrast, the ages are higher (~30 years) in
246 the tropical region. In the zonal direction, the ages in the eastern basin are slightly higher than in the western basin for the
247 central waters, but the difference is only within ~10 years for mode-ages, indicating that the influence from western boundary
248 current is not significant in the upper layer (Fig. 7).

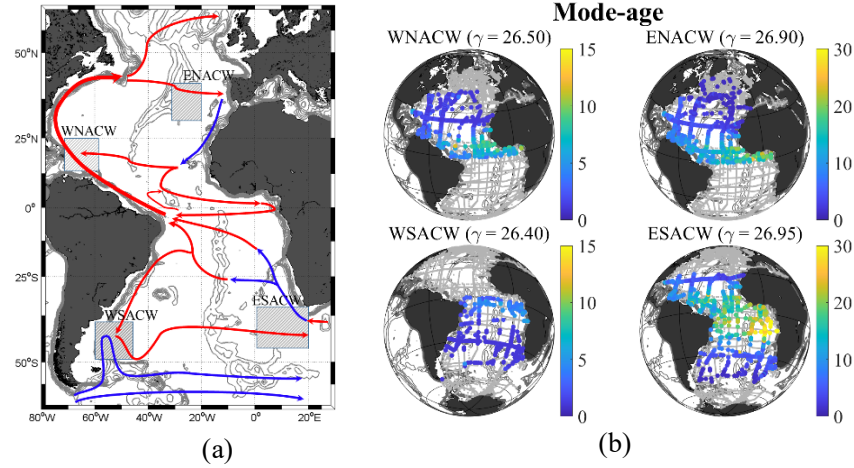
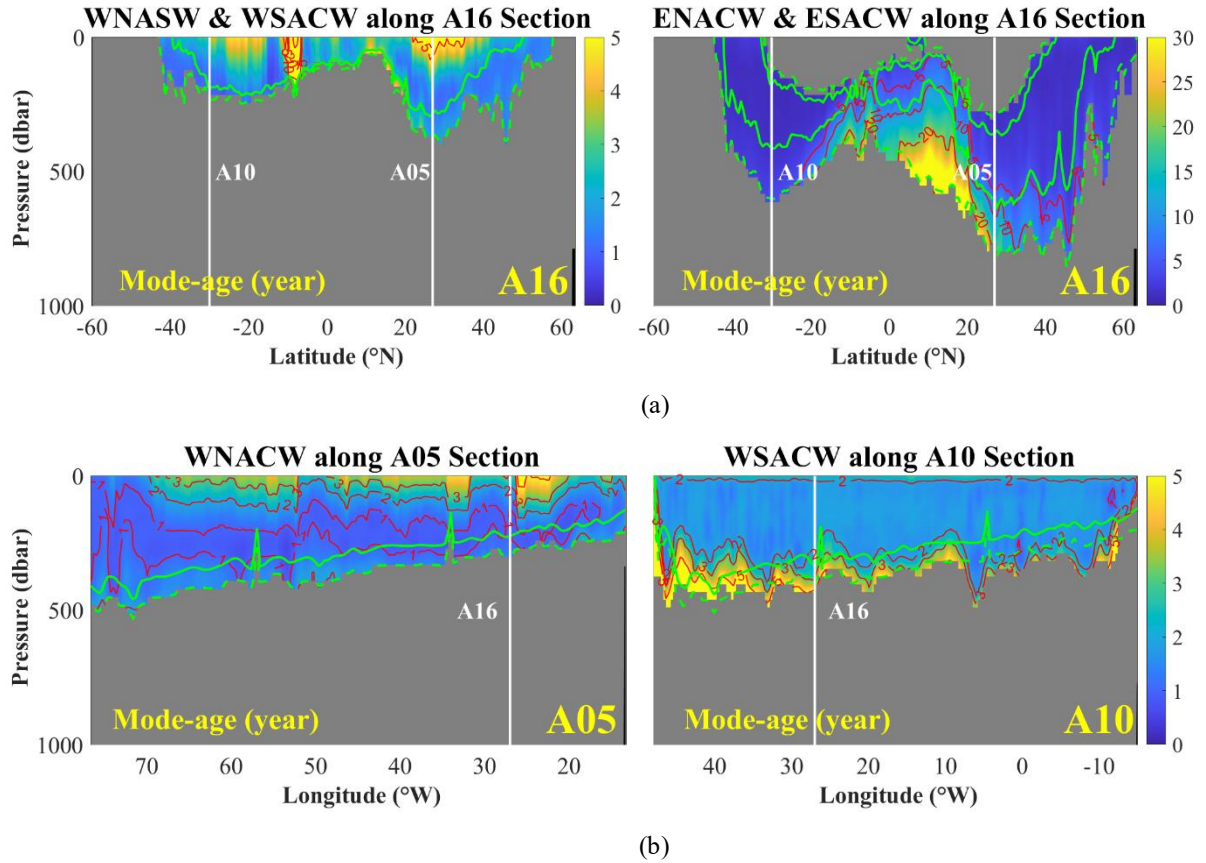
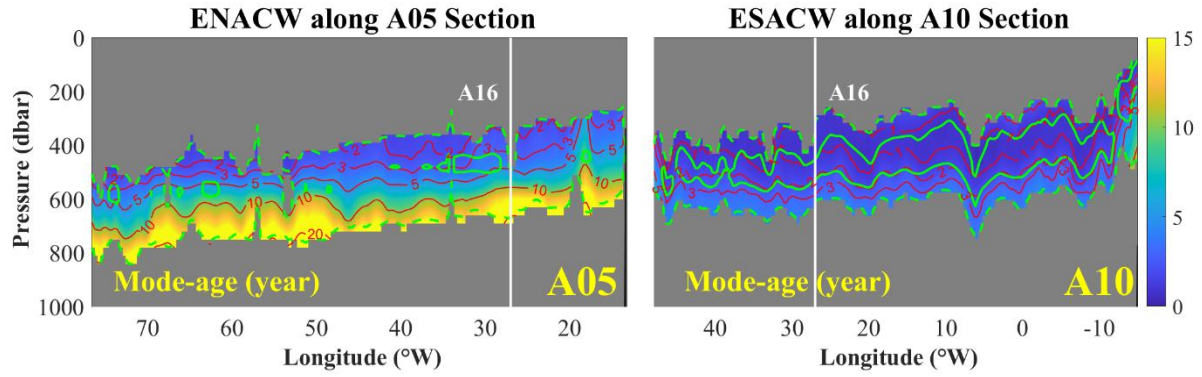


Figure 6. Mode-ages of central water masses in the Atlantic Ocean.

- (a) The main currents in the upper layer with warm (red) and cold (blue) currents and the approximate formation areas (rectangular shadows) of central water masses.
- (b) Mode-ages of central water masses at their core neutral densities (kg m^{-3}). The colored dots show the ages in each station. The grey dots show all the GLODAPv2 stations that have less than 20% contribution of the water mass in question or lack of transient-tracer data.





(c)

Figure 7. The mode-ages (years) of central waters along the three sections for the upper 1500 dbar. The solid green isolines show the 50% fractions of water masses and the dashed green lines show the 20 % fractions. The grey color indicates areas with less than 20% contribution from central waters. White vertical lines show cross overs with other sections.

3.2 The Intermediate Layer

Three main water masses belong to the intermediate layer (γ between 27.10 and 27.90 kg m⁻³), including the Antarctic Intermediate Water (AAIW), the Subarctic Intermediate Water (SAIW) and the Mediterranean Water (MW). The ages of SAIW are not displayed in this study due to the lack of tracer data with a complete section through the distribution region of this water mass.

Antarctic Intermediate Water (AAIW): The AAIW spreads from the formation area between 40 and 50 °S northward to approximately 30 °N along the Western Boundary Current (WBC) (Fig. 8, a). The northward flow of this water mass is part of the Atlantic western boundary current system (Talley, 1996) and the upper limb of AMOC (Kirchner et al., 2009).

Compared to the central waters, AAIW has significantly higher ages. In principle, the AAIW is supposed to get higher ages towards the north, as the transport time increases with the distance from the formation area in the south. In the meridional direction, the mode-ages increase from 0 to ~80 years from formation area to the equator, and further increase up to ~100 years to 20 °N (Fig. 8 and 9). However, both mean- and mode-ages decrease (the mean-age follows the fixed proportion) in the further north region between 20 °N and 30 °N (Fig 8, b), giving the impression of the AAIW being younger in the north Atlantic Ocean. The observed decrease in ages likely results from mixing with younger surrounding water masses, although other factors such as limitations of the steady-state IG-TTD assumption (e.g., Waugh et al., 2004) or regional variations in the mixing ratio (e.g., Thomas et al., 2020) may also contribute. The maximum distance of AAIW to the north can reach 30 °N. However, the mixing is speculated between 20 °N and 30 °N, since the ENACW and upper NADW comes into contact with AAIW from the upper and lower (Liu and Tanhua, 2021) (Fig 7, a, Fig. 9, a and Fig. 11, a). Both ENACW and upper NADW are newly formed in this region, so the “mixed” AAIW obtains younger ages. Age differences are also found in the zonal direction. In better ventilated western basins, the AAIW transports northward with the western boundary current (WBC), so the mode-ages are lower. By contrast, the ages are significantly higher in the eastern basins with poor ventilation. The mode-

270 ages reach up to ~100 years in the east between 0 and 20 °S (Fig. 8, b). The mode-ages along the A05 section also show the
271 zonal difference (Fig. 9, b). In the eastern basin (east of the Mid-Atlantic-Ridge, MAR), the mode-age appears up to ~80 years,
272 in contrast only ~50 years in the west. This distribution confirms that the AAIW is transported by the WBC and takes a longer
273 time to cross the MAR. Generally lower ages are found along the A10 Section in contrast to the A05 Section due to closer
274 distance to the formation area of AAIW (Fig. 9, c). Similar zonal difference exists with mode-ages of ~10 years in the west
275 and ~20 years in the east.

276 The mode-ages of AAIW also vary in the vertical direction. These variations are consistent with the influence of mixing with
277 adjacent water masses along its boundaries. Near the upper boundary, the AAIW is characterized by relatively lower ages,
278 which is consistent with mixing with the generally younger eastern central waters above. Conversely, at the lower boundary
279 with the deep and overflow layer, AAIW encounters the southward upper NADW. Along the A10 section, the upper NADW
280 exhibits high mode-ages (~80 years) due to the long southward transport. The resulting composite water mass at this interface
281 shows intermediate mode-ages (~60 years), higher than those in the AAIW core (~20 years) (Fig. 9 c).

282 A contrasting pattern is observed along the A05 section. In the northern Atlantic sampled by A05, the AAIW obtains higher
283 water mass ages, while the locally encountered upper NADW is relatively younger. Consequently, the mode-ages at the
284 interface (~50 years) are lower than those in the AAIW core at this location (~60 years) (Fig. 9 b). These observed patterns at
285 the boundaries are interpreted as reflecting the mixing of AAIW with water masses of contrasting ages, and the resulting
286 composite age signals are consistent with the independent age estimates for NADW presented in the following section.

287 **Mediterranean Water (MW):** The MW is referred to as the product of the Mediterranean Overflow Water (MOW) that flows
288 across the Strait of Gibraltar and mixes with the ENACW (Liu and Tanhua, 2021). This water mass is formed in the Gulf of
289 Cadiz where the MOW exits the Strait of Gibraltar as a deep current and turns into two branches (Fig 8, a). The northward
290 branch spreads into the West European Basin until 50 °N, while the westward branch spreads across the MAR into the west
291 basin of the North Atlantic Ocean (Price et al., 1993; Carracedo et al., 2016).

292 The spreading of MW can also be traced by the transient tracers (Fig 9, a), in addition to the water mass variables that goes
293 into the OMP analysis (Liu and Tanhua, 2021). The mode-ages of MW have an increasing trend towards the south (Fig. 8, b).
294 The northward flow spreads faster so the mode-age is ~20 years at 50 °N. In contrast, the southward flow shows a higher
295 mode-age of ~100 years at 25 °N along the A05 Section (Fig 9, b). In the zonal direction, the ages decrease to the west (Fig 9,
296 b), since the fraction of MW is only between 20% and 30% along this section (green dash contour lines in Fig 9, b), and is
297 therefore influenced by the lower ages from the NADW.

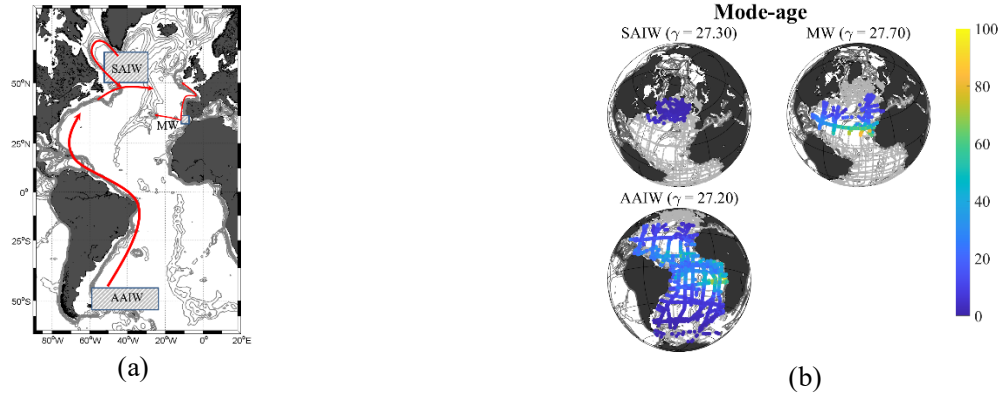
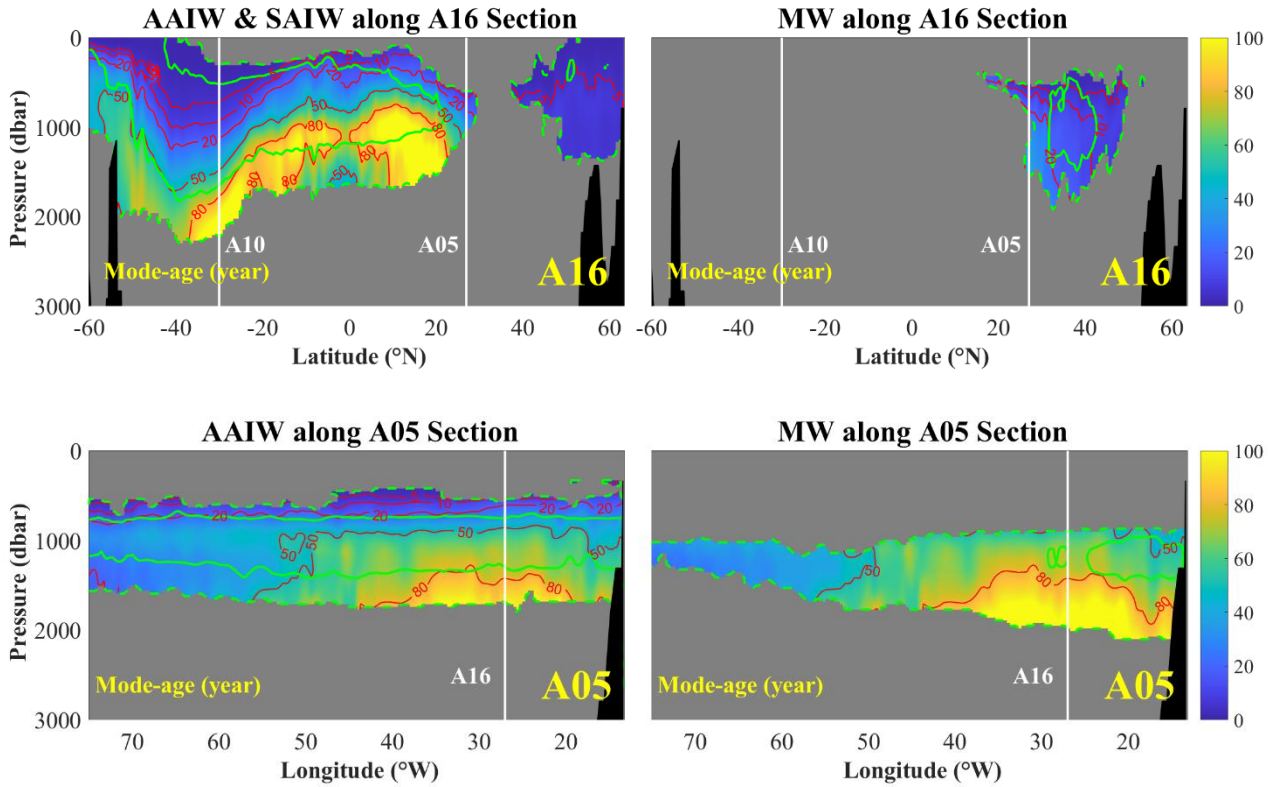


Figure 8. Mode-ages of intermediate water masses in the Atlantic Ocean.

- (a) Main currents in the intermediate layer. The currents (arrows) and the formation areas (rectangular shadows) of water masses in the intermediate layer.
- (b) The mode-ages of intermediate water masses at their core neutral densities (kg m^{-3}). The colored dots show the ages in each station. The grey dots show all the GLODAPv2 stations that have less than 20% contribution of the water mass in question or lack of transient-tracer data.



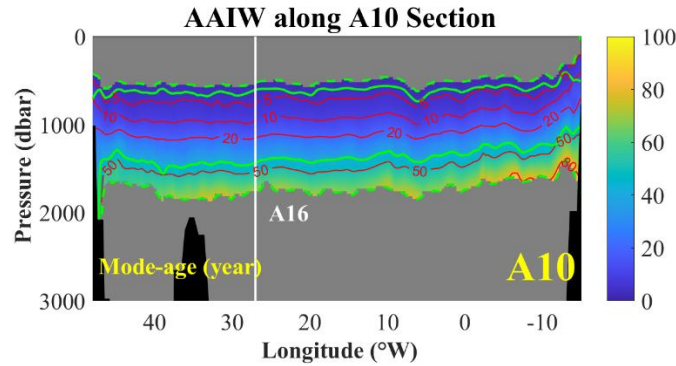


Figure 9. Mode-ages (years) of intermediate water masses along the three sections for the upper 4000 dbar. The solid green isolines show the 50% fractions of water masses and the dashed green lines show the 20 % fractions. The grey blank indicates less than 20% contribution from intermediate water mass. White vertical lines show cross overs with other sections.

3.3 Deep and Overflow Layer

The North Atlantic Deep Water (NADW), including its upper and lower portions, dominates the deep and overflow layer (γ between 27.90 and 28.10 kg m^{-3}) of the Atlantic Ocean. This water mass is formed in the high latitude region in the North Atlantic and spreads southward at pressures between 2000 and 4000 dbar until it meets AAIW and AABW in the Antarctic Circum-polar Current (ACC) region. The southward spreading pathway is along the Deep Western Boundary Current (DWBC, Fig. 10, a). Meanwhile, the NADW also extends eastward with the eddies and finally covers the deep and overflow layer (Dickson and Brown, 1994).

Upper and lower North Atlantic Deep Water (NADW): The NADW is a water mass that is transported far southward from its formation area, with a mode-age of ~ 100 years at the southern limb in the Antarctic Circum-polar Current (ACC) region (Fig. 10, b). The increasing ages during the pathway to the south is shown along the A16 section (Fig. 11, a). After leaving the formation area, the mode-ages increase ~ 20 years from 60 °N to 40 °N. Afterwards, a sharp increase in mode-ages appears in the region between 30 °N and 10 °N. In this interval, the highest mode-age reaches up to ~ 150 years. This is because the A16 section at this latitude range passed through the east part of the Atlantic Ocean (east of MAR) and the distance from the Deep Western Boundary Current (DWBC, Fig. 4 and Fig. 10, a) was high. High ages in this region are due to the sluggish water exchange over the ridge. This result can also be confirmed by the observation along the A05 Section (Fig. 11, b): The NADW near the DWBC region has a low mode-age of ~ 40 years, while it reaches up to ~ 100 years east of the MAR. After passing the equator, when the A16 section returns to the west of the MAR, the mode-ages of NADW are ~ 80 years in the latitude range between 0 and 20 °S, which further increase to ~ 100 years along the pathway southward until 40 °S. In the ACC region between 40 °S and 60 °S, the mode-ages of upper and lower NADW decrease to ~ 80 years. Similar to the situation of AAIW in the north Atlantic, this is also the result of mixing with newly formed AAIW and AABW. The spreading of NADW in the zonal direction is slower eastward, so the ages are generally higher in the eastern basin of the Atlantic Ocean (Fig. 10, b). The tracer data along the A05 and A10 sections also reflect the above result. In the A05 section, the mode-ages are ~ 50 years in the west and ~ 100 to ~ 120 years in the east (Fig. 11, b). In the A10 section, the mode-ages are ~ 80 years in the west and ~ 120 years in

321 the east (Fig. 11, c).

322 In the vertical direction, the upper NADW mixes with the AAIW from above and the lower NADW mixes with the AABW
323 from below. As the NADW is transported in the opposite direction to these two water masses, the "mixed" ages in the north,
324 which is closer to the formation area of NADW, are higher than the age at the core density of NADW, while the opposite
325 situation exists when spreading to south. After entering the ACC region, both mean- and mode-ages decrease (the mean-age
326 follows the fixed proportion) since the fractions of newly formed AAIW and AABW are high.

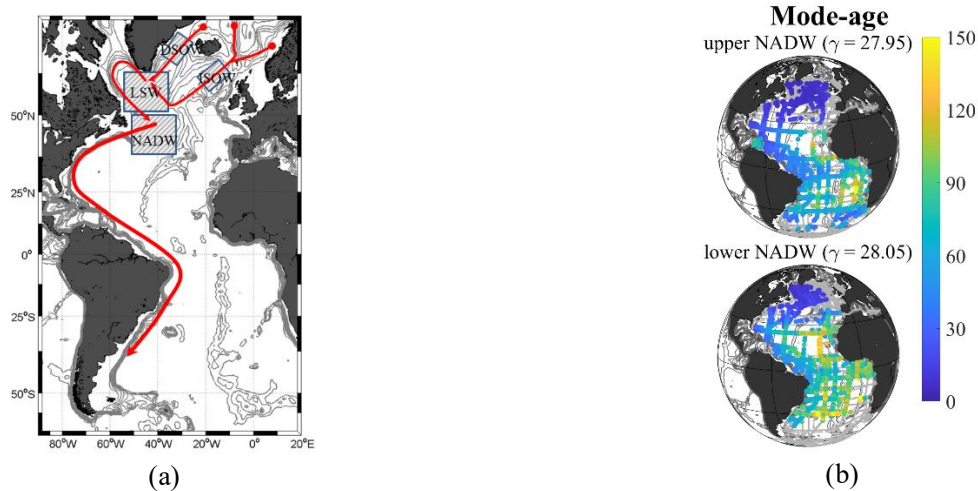


Figure 10. Mode-ages of upper and lower NADW in the Atlantic Ocean.

(a) Main currents in the deep and overflow layer. The currents (arrows) and the formation areas (rectangular shadows) of water masses in the deep and overflow layer.

(b) The mode-ages (years) of upper and lower NADW at their core neutral densities (kg m^{-3}). The colored dots show the ages in each station. The grey dots show all the GLODAPv2 stations that have less than 20% contribution of the water mass in question or lack of transient-tracer data.

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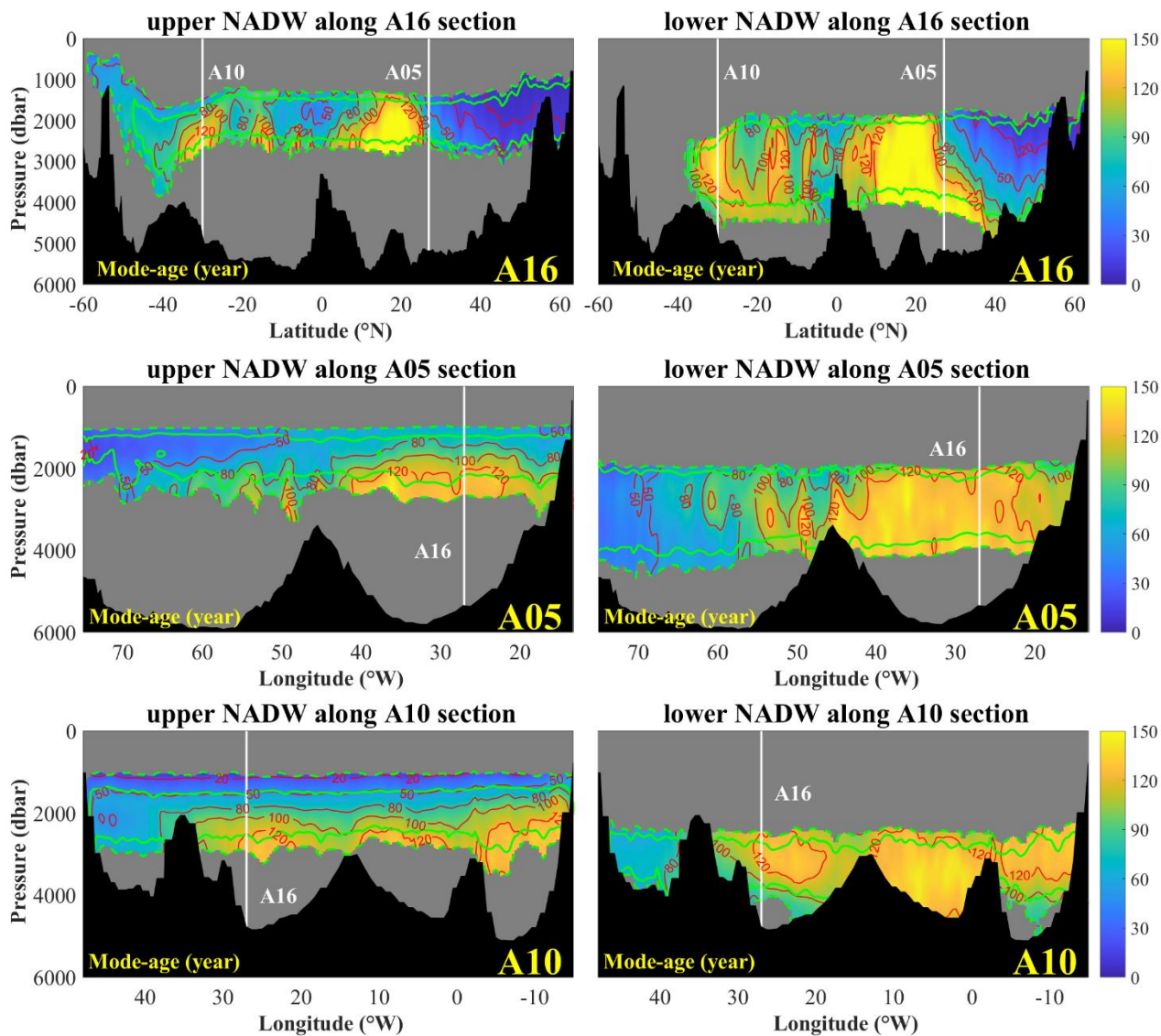


Figure 11. Mode-ages (years) of upper and lower NADW along the A16 (a), A05 (b) and A10 (c) section lines. The green contour lines show fractions of water masses in 50 % and the dashed green lines show the 20 % fractions. The grey blank indicates less than 20% contribution from NADW. The white vertical lines show cross overs with other section lines.

329 3.4 Bottom Layer

330 The bottom layer is noticed already by Wüst, (1933) as it is filled with water from the south. Sverdrup, (1940) further pointed
331 out the role of ACC plays in the formation and spreading of Antarctic Bottom Water (AABW). On this basis, Mantyla and
332 Reid, (1983) elaborated that the most original dense bottom water is restricted near the Antarctic region by topography and the
333 northward flow of bottom water is the mixture of original dense water and the overlying warm water. Gordon and Huber (1990)
334 and Orsi et al., (1999) illustrated that the pathway of AABW to the north is through the Drake Passage sill into the Argentina
335 and Brazilian Basin (Fig. 12, a). In the Atlantic Ocean, the AABW dominates the bottom layer ($\gamma > 28.10 \text{ kg m}^{-3}$), which
336 originates from the Weddell Sea and spreads to the north. After passing the equator, the AABW is redefined as Northeast Atlantic
337 Bottom Water (NEABW, Liu and Tanhua, 2021).

338 **Antarctic Bottom Water (AABW) & Northeast Atlantic Bottom Water (NEABW):** The transport of bottom water is a long
339 process going from the south towards the north. In the meridional direction, the AABW is transported towards the equator with
340 a mode-age of ~100 years at the equator, and further northward as NEABW to 40 °N with a mode-age of ~150 years (Fig. 12,
341 b). Combined with the A16 Section, more specific segments/details can be seen. In the early stage from the surface of Weddell
342 Sea to the bottom (below the pressure 4000 dbar) at 40 °S, the mode-age of AABW is ~50 years. In the range between 10 and
343 30 °S, high values of ~120 years (mode-age) appear due to the mixing with the lower NADW with high ages (Fig. 13, a). After
344 passing the equator, the redefined NEABW contains a mode-age up to ~150 years at 40 °N, indicating that the bottom water,
345 including the AABW and NEABW, takes about 150 years (mode-age) for the transport from the formation area (Weddell Sea)
346 to 40 °N. North of 40 °N, the ages decrease due to the mixing with newly formed lower NADW in this region (Fig. 12, b and
347 c, Fig. 13, a). The distribution of mode-ages also shows a difference in the zonal direction. As shown in Fig. 12 b, the general
348 age distribution presents a trend with lower ages in the west, and higher in the east. Along the A10 section at 30 °S, the AABW
349 has a mode-age of ~50 in the west and ~100 in the east. A similar situation is recorded along the A05 section; the mode-age of
350 NEABW is ~80 years in the west and ~120 years in the east (Fig. 13, b). An additional factor that affects the age distributions
351 is the mixing between AABW and NADW. In the south hemisphere, mode-ages are relatively low (~80 years) at the pressures
352 below 4000 dbar, where the AABW has a fraction higher than 50%, while they are significantly higher when the AABW mixes
353 with lower NADW at pressure of ~3000 dbar, especially in the region between 0 and 10 °W, reaching ~120 years. This is
354 because the "old" NADW takes up a fraction for more than 50% to 80%. The situation in the north hemisphere is the opposite,
355 the mode-ages of NEABW are higher in the bottom layer below 4000 dbar but lower in the deep layer at 3000 dbar due to the
356 mixing with the young NADW. In the western basin, the mode-age is ~80 years in the bottom, while ~50 years when mixed
357 with NADW. Similar in the east basin, the mode-age is ~120 years below 4000 dbar, while ~100 years at 3000 dbar (Fig. 13,
358 b).



Figure 12. Mode-ages of bottom water masses in the Atlantic Ocean.

- (a) Main currents in the bottom layer. The currents (arrows) and the formation areas (rectangular shadows) of water masses in the deep and overflow layer.
- (b) The mode-ages (years) of bottom water masses at their core neutral densities (kg m^{-3}). The colored dots show the ages in each station. The grey dots show all the GLODAPv2 stations that have less than 20% contribution of the water mass in question or lack of transient-tracer data.

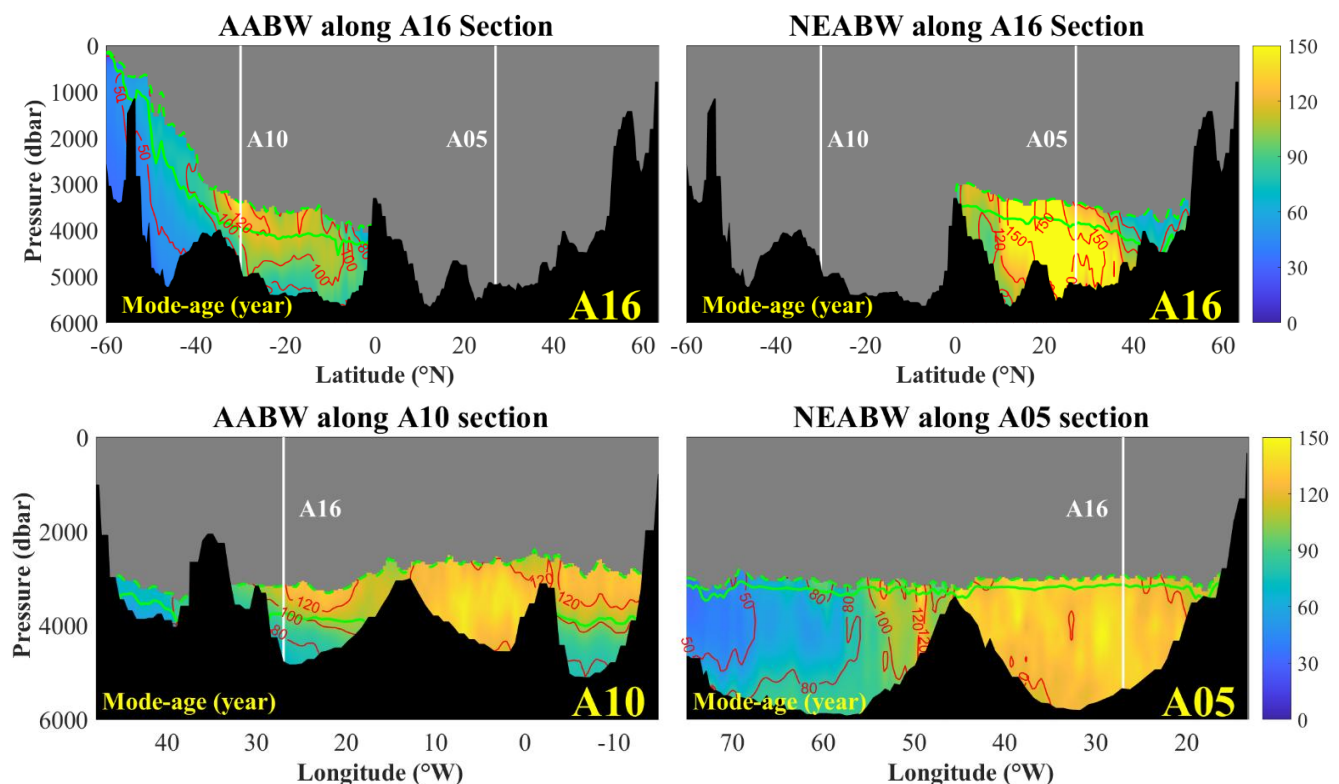


Figure 13. Mode-ages of AABW and NEABW along the A16 (a), A05 and A10 (b) section lines.

The green contour lines show fractions of water masses in 50 and the dashed green lines show the 20 % fractions. The grey blank indicates less than 20% contribution from bottom water masses. The white vertical lines show cross overs with other section lines.

359 4 The application of water mass ages in estimating the oxygen utilization rate (OUR)

360 The transit time distribution (TTD) provides a framework for using transient tracers to estimate the time scales governing
361 ventilation, which in turn can be applied to quantify biogeochemical rates. For instance, based on the TTD determined from
362 the tracer observations, Wang et al., (2021) estimated the OUR in the northern South China Sea.

363 The oxygen concentration in seawater under steady state is maintained by the balance of supply from oxygen-rich surface
364 water and consumption by the respiration of organic matter. The Apparent Oxygen Utilization (AOU) shows the accumulated
365 oxygen consumption in a water mass since isolated from the atmosphere, and is defined as the difference between saturated
366 and measured oxygen concentration (i.e. $\text{AOU} = \text{Oxy}_{\text{saturated}} - \text{Oxy}_{\text{measured}}$, Redfield, 1942; Redfield et al., 1963; Pytkowicz,
367 1971). The Oxygen Utilization Rate OUR ($\mu\text{mol kg}^{-1} \text{ yr}^{-1}$) can be calculated from AOU and water mass age and shows the
368 integrated oxygen utilization rate (oxidation rate) during the pathway of a water mass from surface of the formation area to the
369 sampling location (Jenkins, 1987; Doney and Bullister, 1992; Sonnerup et al., 2013). In addition, the OUR is also an important
370 indicator in characterizing the combination of ventilation and respiration of the interior ocean (Koeve and Kähler, 2016;
371 Thomas et al., 2020).

372 The true local OUR is difficult to determine because water samples contain a mixture of paths with different transit times
373 (Tomczak, 1999; Koeve & Kähler, 2016). Researchers have tried a variety of ways to define the age of a water mass. For
374 instance, the tracer-age (CFC age) is used in Karstensen et al., (2008), while the mean-age (partial pressure ages) is used in
375 Sonnerup et al., (2015) in estimating the “integrated” water mass age that effects the oxygen consumption. There is no direct
376 evidence that one age definition is theoretically superior for OUR calculations; both the mean-age and the mode-age are
377 approximations. The mean-age (Γ) averages over all pathways, including old water fractions that have experienced very low
378 respiration rates in the deep ocean; using it would tend to lower the estimated OUR. The mode-age, in contrast, represents the
379 most probable transit time of the dominant water fraction and is often interpreted as the advective timescale. As a working
380 assumption, we adopt the mode-age to calculate OUR, reasoning that it may better reflect the respiration accumulated along
381 the core pathway of the water mass. However, we emphasize that the resulting OUR values are indicative rather than absolute.
382 If the mean-age were used instead, the computed OUR would be roughly six times lower (given $\tau_{\text{mode}} = \Gamma \cdot (\sqrt{10} - 3) \approx$
383 $0.162 \cdot \Gamma$), bracketing the plausible range. Our choice of mode-age likely provides an upper-end estimate of the respiration rate
384 for the dominant water mass. For example, if we use the mean-age instead of the mode-age for the NADW core in the western
385 basin, the OUR would decrease from $\sim 2 \mu\text{mol kg}^{-1} \text{ yr}^{-1}$ to $\sim 0.3 \mu\text{mol kg}^{-1} \text{ yr}^{-1}$. Both values are within the range reported in the
386 literature (e.g., Jenkins, 1987; Stanley et al., 2012), highlighting the uncertainty associated with the choice of age metric.

387 Water masses with mode-ages larger than ~ 50 years in some parts of their distribution, including the AAIW, NADW, AABW
388 and NEABW in the Atlantic Ocean, are defined here as wide-spread water masses. The transports of these wide-spread water
389 masses are significantly influenced by the currents and topography. Therefore, their mode-age shows regional differences.
390 Such differences also exist in DO and AOU, and lead to the spatial differences in OUR, especially between the eastern and

391 western basins (Fig. 14). In the meridional direction, the mode-age increases with the distance from formation area. In the
392 zonal direction, significant differences exist between the east and west of MAR. The wide-spread water masses generally have
393 lower mode-ages in the western basin, which is better ventilated by the WBC and the DWBC (Fig. 14, left). In addition, surface
394 oxygen-rich water is transported by these water masses to the deeper layers and are gradually consumed during the path-way.
395 As a result, the DO is highest in the formation area and decreases with the distance. Meanwhile, the western basin displays
396 higher DO due to the better ventilation (Fig. 14, middle left). The OUR shows a similar distribution to DO indicating that
397 higher oxygen consumption rate, or oxidation rate is co-located with the regions with high oxygen concentration. At the same
398 time, Fig. 14 also shows that the OUR approaches 0 when the mode-age is larger than ~50 years.

399 In the intermediate layer, the AAIW show low mode-ages, high DO and low AOU in the formation area near the Antarctic
400 region. During the northward transport between 40 °S and 20 °N, the mode-ages increase and the DOs decrease with distance
401 from the formation area (Fig. 14, upper panel). In general, the mode-ages are lower in the west, while the zonal difference for
402 DOs is not significant except for some regions in the eastern basins. The mode-ages are mostly between ~60 and ~90 years
403 and reaches up to ~120 years in the poor ventilated eastern basin. The DOs are between 160 and 220 $\mu\text{mol kg}^{-1}$ in the south
404 hemisphere and mostly between 100 and 160 $\mu\text{mol kg}^{-1}$ in the north hemisphere. The OURs in the Antarctic region obtain the
405 highest value in the latitude region between 20 and 40 °S and decrease on the pathway to the north until 20 °N.

406 In the deep and overflow layer, the NADW is newly formed in the high north latitude. In most regions covered by the NADW
407 between 30 °N and 40 °S, the differences between west and east exist with obvious low mode-age and AOU (high DO) and in
408 the west, where is well ventilated by the DWBC (Fig. 14). The mode-age is between ~30 and ~60 years in the west and between
409 ~90 and ~150 years in the east. The AOU is lower in the west and higher in the east. The OUR, which shows the opposite
410 distribution as the AOU in the zonal direction, obtains the highest value in the formation area of NADW, maintains relative
411 higher in the west along the DWBC region and shows a lower value in the eastern basin.

412 The bottom layer is occupied by the AABW and NEABW in the south and north hemisphere respectively. The AABW is
413 formed in the Antarctic region with an original mode-age lower than ~30 years and increases on the way to the north. Similar
414 to the AAIW and NADW, the western basin is better ventilated, therefore the mode-ages are lower in the west at the same
415 latitude (Fig. 14, lower panel). In contrast to the AAIW and NADW, the AABW obtains relative lower DOs and higher AOU
416 when formed (see also Fig. 15, b). In the western basin of the South Atlantic, the DOs decrease to 220 $\mu\text{mol kg}^{-1}$ and the AOU
417 increase to 130 $\mu\text{mol kg}^{-1}$ on the pathway, while in the eastern basin, the DOs are 240 $\mu\text{mol kg}^{-1}$ and the AOU are 110 μmol
418 kg^{-1} . This is because the eastern basin is more influenced by the relative oxygen-rich NADW from the north. Similarly, the
419 DOs and AOU of NEABW in the North Atlantic also show similar states to the NADW. In west regions ventilated by the
420 DWBC, the DOs reach the values of higher than 260 $\mu\text{mol kg}^{-1}$ and AOU is lower than 70 $\mu\text{mol kg}^{-1}$. This result can also be
421 confirmed in Figure 15. Although the distributions of DOs and AOU are more complex in the bottom layer, the OURs still
422 match the distribution of mode-ages. The highest OURs appears in the formation area near the Antarctic region with values
423 higher than 4 $\mu\text{mol kg}^{-1} \text{ yr}^{-1}$. In the western Atlantic region with low mode-ages, the OURs show relative higher values (>

424 between 1 and 3 $\mu\text{mol kg}^{-1} \text{yr}^{-1}$) and low value (below 1 $\mu\text{mol kg}^{-1} \text{yr}^{-1}$) in the east (Fig. 14, lower panel).

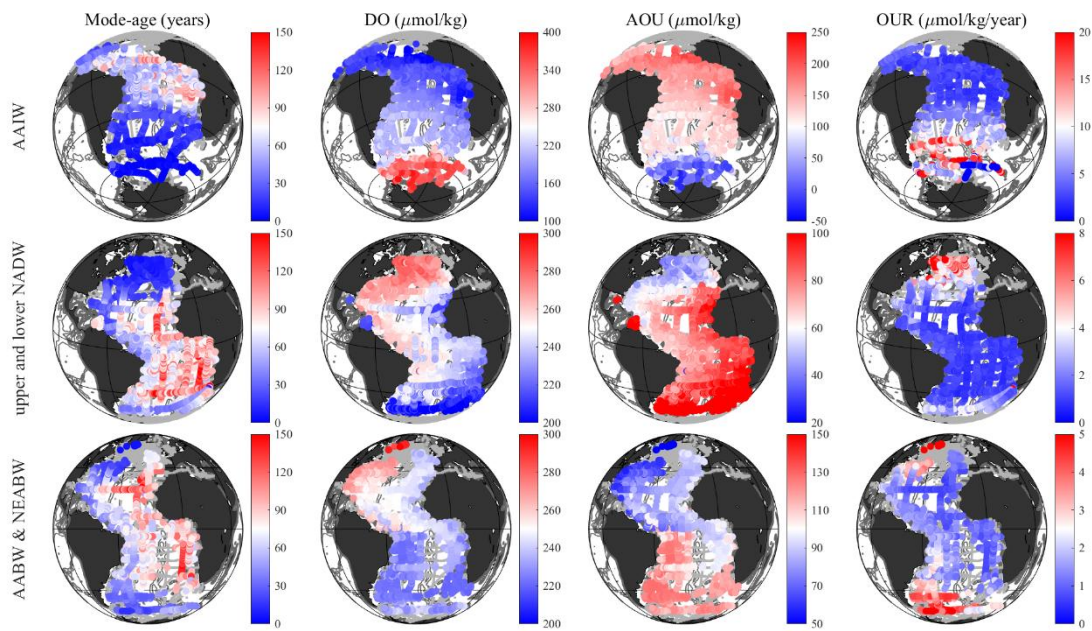


Figure 14. Mode-age, DO, AOU and OUR of wide-spread water masses in the Atlantic Ocean.

The colored dots show the values in each station at core neutral densities of each water mass (AAIW at 27.20, upper NADW at 27.95, lower NADW at 28.05, AABW at 28.15, NEABW at 28.10 kg m^{-3} , the upper and lower NADW, AABW and NEABW are shown in one plot). The grey dots show all the GLODAPv2 stations that have less than 20% contribution of the water mass in question. Note: Negative AOU values indicate oxygen supersaturation, which is typically observed in well-ventilated surface or near-surface waters.

425 The above four properties are also represented along the A16 section (Fig. 15). The AAIW has the lowest mode-age among
 426 the three water masses, but also the largest variations in the DO and AOU. The mode-age indicates that in total of ~40 to 60
 427 years is needed for the main body of AAIW (fractions > 80%) to be transported to the equator (Fig 15, a). The DO decreases
 428 rapidly from 260 to below 180 $\mu\text{mol kg}^{-1}$ at the oxygen minimum zone at 20 °S, while the AOU increases from 60 to 150 μmol
 429 kg^{-1} during the northward pathway (Fig 15, b and c). The OUR is higher than 8 $\mu\text{mol kg}^{-1} \text{yr}^{-1}$ at the beginning and decreases
 430 to 3 $\mu\text{mol kg}^{-1} \text{yr}^{-1}$ close to the equator (Fig 15, d). The mode-age of NADW is ~140 years at 40 °S and reaches a peak of ~160
 431 years in the region of 20 °N due to the sluggish water exchange caused by the topography. The NADW contains the highest
 432 oxygen concentration and the lowest AOU and OUR in the abyssal Atlantic Ocean. The value of DO is 260 $\mu\text{mol kg}^{-1}$ at the
 433 starting point at 60 °N and still remains 220 $\mu\text{mol kg}^{-1}$ at 40 °S where it meets the AABW and AAIW. The AOU of NADW
 434 remains below 100 $\mu\text{mol kg}^{-1}$ during the entire pathway. The OUR of NADW is below 5 $\mu\text{mol kg}^{-1} \text{yr}^{-1}$ close to the formation
 435 area, and below 1 $\mu\text{mol kg}^{-1} \text{yr}^{-1}$ south of 60 °N. The transport of the main body of AABW takes in total of ~100 years from
 436 the Weddell Sea to the equator. The DO and AOU remain unchanged at 220 and 130 $\mu\text{mol kg}^{-1}$ respectively on the pathway to
 437 the equator, indicating very low OUR. Although derived from the AABW, the characteristics of the NEABW are more similar
 438 to the NADW in all the above properties.

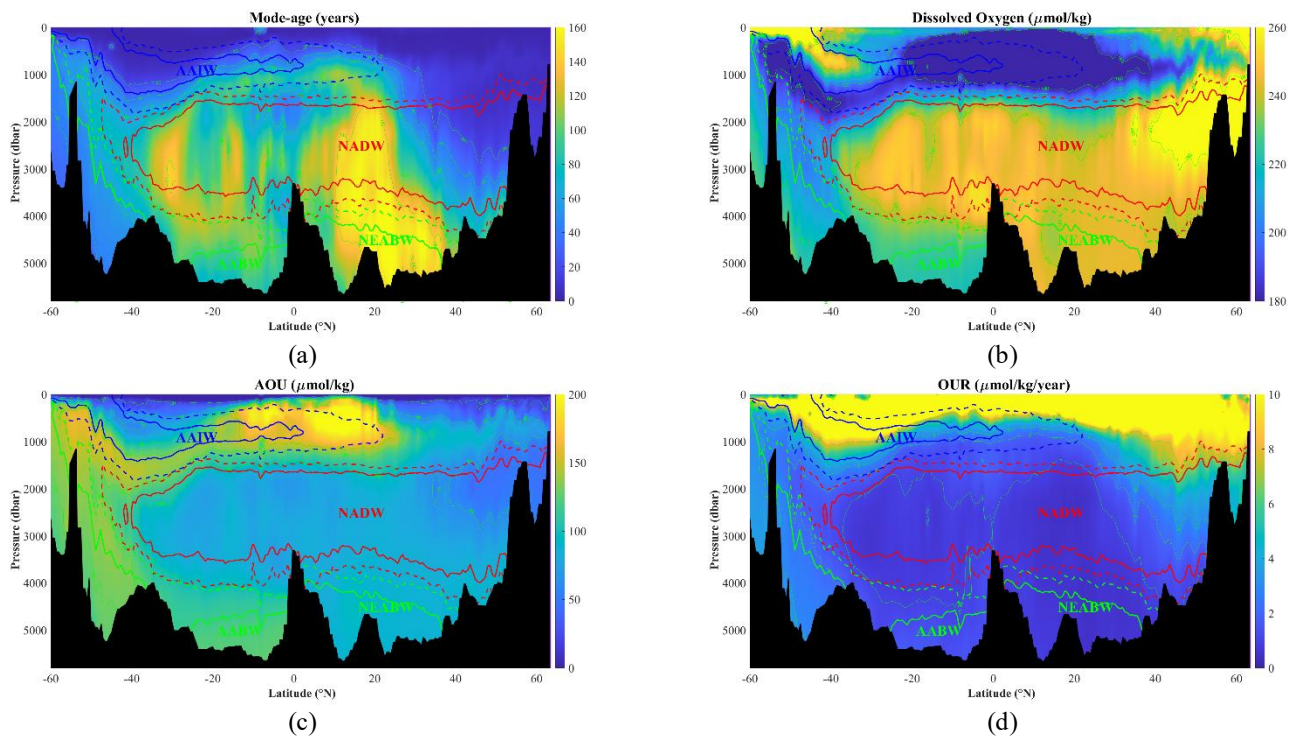


Figure 15. Distributions of (a) mode-age, (b) dissolved oxygen, (c) AOU and (d) OUR along the A16 section line. The color solid lines show the fractions of 80% in each water mass and the dashed lines show the fractions of 50%.

440 5 Estimation of water mass ages based on ^{39}Ar

441 The radioactive isotope ^{39}Ar (half-life 269 years) is a valuable tracer for dating old water masses. The ^{39}Ar data used in this
 442 study are compiled from the doctoral thesis of Rodriguez (1993), which tabulates 125 measurements collected during the 1980s
 443 and early 1990s in the Atlantic Ocean using low-level decay counting (Loosli et al., 1983; Schlosser et al., 1994). More recent
 444 ^{39}Ar data from ATTA technology (e.g., Lu et al., 2014; Ebser et al., 2018) are not yet available in sufficient numbers for
 445 basin-scale synthesis. Therefore, our analysis relies on the Rodriguez (1993) dataset. We note that these ^{39}Ar measurements
 446 were taken approximately two decades before most GLODAPv2 CFC-12/SF₆ data (which are from the 2000s and 2010s). This
 447 temporal mismatch, together with the assumption of steady-state circulation, introduces additional uncertainty into the
 448 comparison. Nevertheless, the paired analysis presented below (Fig. 18) is based on spatially collocated stations, recognizing
 449 that the ventilation state may have changed over the intervening decades. For ^{39}Ar we focus our attention on older water masses,
 450 as younger ones are better characterized by CFC-12 and SF₆. Because this study adopts a fixed mixing ratio ($\Delta/\Gamma = 1$), the
 451 ^{39}Ar -derived ages, which physically represent mean residence times, are converted to mode-ages by multiplying by 0.162 for
 452 consistent comparison with the CFC-12/SF₆-derived mode-ages. All ages reported in this section are therefore mode-ages.

453 For this purpose, we make a distinction between the western (green dots) and eastern (red dots) Atlantic, divided by the Mid-
 454 Atlantic-Ridge (Fig. 16a). The distribution of ^{39}Ar is shown in Fig. 16 b. Near the formation area of each wide-spread water
 455 mass, the concentration of ^{39}Ar approaches approximately 100%, indicating that the surface waters are in equilibrium with the
 456 atmosphere. As these water masses are transported over long distances and extended periods, the continuous decay of ^{39}Ar
 457 leads to concentrations decreasing to roughly 30% at their furthest locations. By integrating this information with its half-life
 458 of 269 years, the water mass ages can be estimated. For ^{39}Ar we focus our attention to the older water masses, as the younger
 459 ones are better characterized by CFC12 and SF₆.

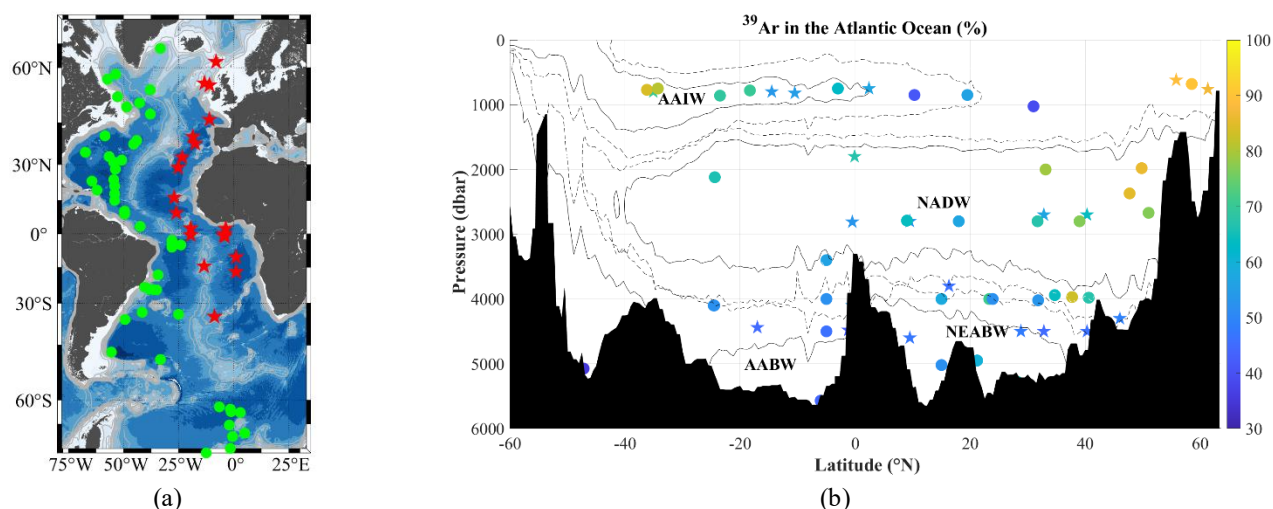


Figure 16. (a) Map of the ^{39}Ar sampling stations. The green circle dots show the stations in the west Atlantic Ocean (west of the Mid Atlantic Ridge), while the red stars show the stations in the east Atlantic Ocean. (b) Distributions of ^{39}Ar in the

Atlantic Ocean (% Modern). The circle dots show the data in the west, while the stars show the data in the east Atlantic Ocean. The solid isolines show the 80% fractions of water masses and the dashed lines show the 50% fractions.

460 Similar to the findings from CFC-12 and SF₆ observations, the mode-ages derived from measurements using ³⁹Ar increase
 461 with transport from the source of formation (Fig. 17). At a latitude of approximately 40°S, the AAIW exhibits mode-ages
 462 ranging between ~16 to ~32 years (the corresponding mean-ages are ~100–200 years). When transported northward to about
 463 20 °S, the mode-ages rise to ~60 years (mean-age ~300 years). Upon reaching approximately 10 °N during further northward
 464 transport, these values escalate to around ~100 years for mode-age (~700 years for mean-age). However, near the region at
 465 about 20°N, the terminus for northward flow, the mode-age decreases back down to roughly ~65 years (mean-age ~400 years)
 466 due to mixing with younger upper NADW. In the deep and overflow layer, the newly formed NADW initiates at high northern
 467 latitudes, exhibiting both mean- and mode-ages approaching zero. During southward transport at around 30 °S, the
 468 concentration data indicates corresponding mode-ages of approximately ~65 years (mean-age ~400 years). Similarly, the
 469 bottom water masses (AABW and NEABW) display mode-ages of around ~80–100 years (mean-age ~500–600 years) at the
 470 farthest end of the transport distance approximately 50 °N.

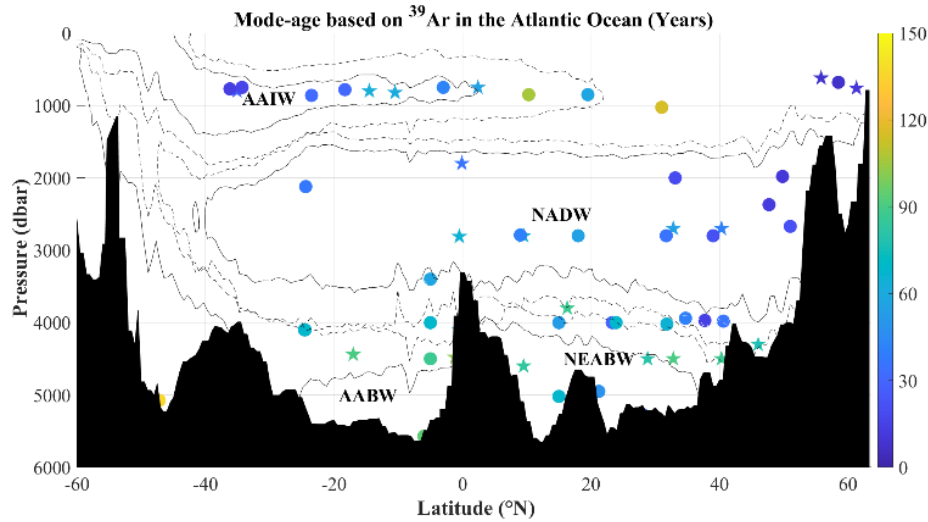


Figure 17. Mode-ages of the wide-spread water masses estimated by ³⁹Ar in the Atlantic Ocean. The circle dots show the data in the west, while the stars show the data in the east Atlantic Ocean. The solid isolines show the 80% fractions of water masses and the dashed lines show the 50 % fractions.

471 The contrast between the western and eastern Atlantic Ocean is distinctly evident (Fig. 16, b). Generally, higher concentrations
 472 of ³⁹Ar are observed in the west compared to the east at equivalent latitudes. For instance, in the vicinity of the AAIW formation
 473 area at 40 °S, the ³⁹Ar concentration is approximately 100% in the west and around 70% in the east. In the intermediate layer
 474 near the equator, these concentrations decrease to about 70% (west) and 50% (east). These findings suggest that mode-ages are
 475 close to zero at 40 °S. The mode-age is estimated to be ~50 years in the west and ~80 years in the east north of 40 °S. The
 476 mean-age is estimated to be ~300 years in the west and ~500 years in the east north of 40°S. Similar results are also observed

477 in the deep water and overflow layers. Based on elevated levels of ^{39}Ar near its formation area, it can be inferred that both
 478 mean- and mode-ages of NADW are close to zero at 60 °N across both West and East Atlantic Oceans. Between 20 °N and the
 479 equator, the the mode-age is estimated to be ~50 years in the west and ~80 years in the east, respectively. For bottom water
 480 masses below a depth of 4000 m, concentrations of ^{39}Ar generally fall below 50%, with higher levels observed in the west
 481 (50%-60%) compared to lower levels seen in the east (40%-50%) between 20 °S and 40 °N, indicating the mode-ages range
 482 from ~65–80 years in the west, and ~80–100 years in the east.

483 **Table 2. Comparison of mode-ages (years) estimated from CFC-12/SF₆ and ^{39}Ar for selected Atlantic water masses.**

Water Mass	Location	Latitude/Longitude Range	Neutral Density (kg m ⁻³)	CFC-12/SF ₆ mode-age	^{39}Ar mode-age
AAIW (40°S)	West	35–45°S, 60–20°W	27.15–27.25	11 ± 2	24 ± 3
AAIW (40°S)	East	35–45°S, 20°W–10°E	27.15–27.25	32 ± 4	28 ± 4
AAIW (20°S)	West	15–25°S, 40–20°W	27.20–27.30	49 ± 6	49 ± 6
AAIW (20°S)	East	15–25°S, 20°W–10°E	27.20–27.30	81 ± 10	81 ± 10
NADW (20°N)	West (A05)	15–25°N, 70–40°W	27.95–28.05	32 ± 5	21 ± 3
NADW (20°N)	East (A05)	15–25°N, 40–10°W	27.95–28.05	97 ± 13	65 ± 8
AABW (30°S)	West (A10)	25–35°S, 50–30°W	>28.10	65 ± 8	81 ± 10
AABW (30°S)	East (A10)	25–35°S, 10°W–10°E	>28.10	97 ± 13	97 ± 13

484 Note: Uncertainties for CFC-12/SF₆ are derived from mixing ratio sensitivity tests ($\Delta/\Gamma = 0.8 - 1.2$); uncertainties for ^{39}Ar
 485 are from measurement statistics (1σ). Location definitions follow the neutral density ranges given in Liu and Tanhua (2021),
 486 “West” = west of Mid-Atlantic Ridge (MAR); “East” = east of MAR. A full version of this table including mean- and
 487 mode-ages and additional locations is provided in the Supplement (Table S1). The ages are core values at the specified neutral
 488 density ranges. For NADW, “A05” refers to the zonal section at ~24 °N; for AABW/NEABW, “A10” refers to the zonal section
 489 at ~30 °S. Uncertainty ranges are rounded to nearest 5 years. ^{39}Ar data are from stations not exactly coincident with CFC-

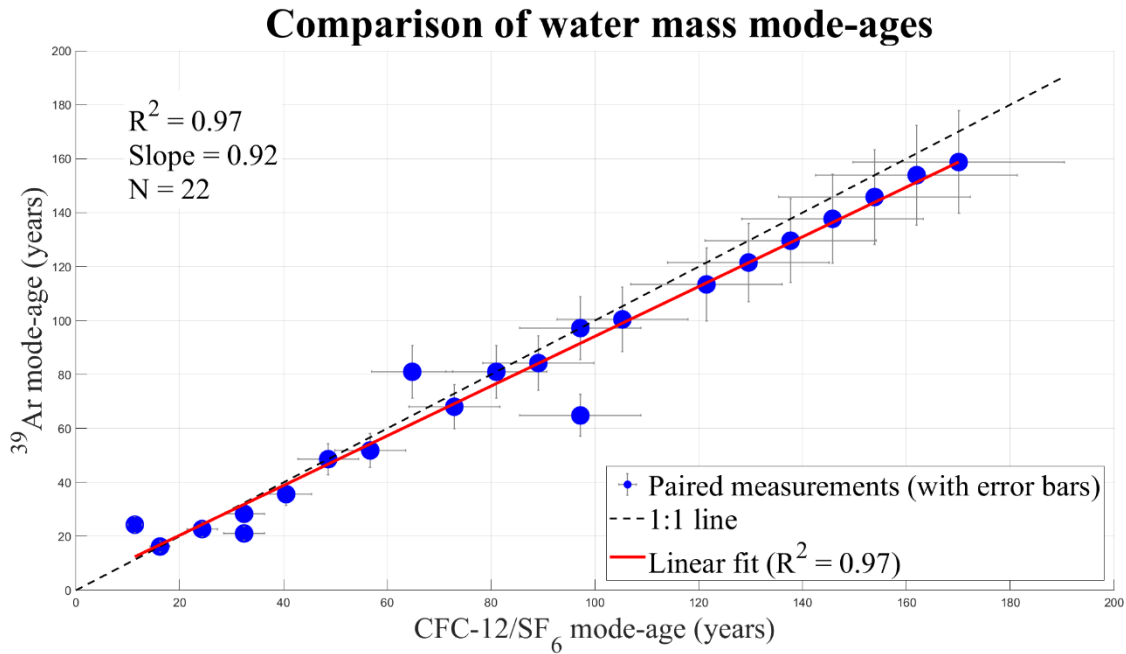


Figure 18. Scatter plot of ³⁹Ar-derived mode-ages versus CFC-12/SF₆-derived mode-ages for paired Atlantic water mass samples (AAIW, NADW, AABW/NEABW). Each point represents an ³⁹Ar measurement (Rodríguez, 1993; collected in the 1980s–1990s) paired with the nearest GLODAPv2 station (2000s–2010s) within 2° in latitude/longitude and 200 dbar in pressure, sampling the same water mass. Error bars: horizontal = CFC-12/SF₆ age uncertainty estimated from mixing ratio sensitivity tests ($\Delta/\Gamma = 0.8 - 1.2$); vertical = ³⁹Ar age uncertainty from measurement statistics (1 σ). Dashed line: 1:1 reference. Solid red line: linear regression (major axis) with $R^2 = 0.98$ ($N = 22$). The comparison assumes steady state ventilation over the ~20 years temporal mismatch between the two datasets.

491 Linear regression of the 22 paired points yields: $^{39}\text{Ar mode-age} = 0.92 \times (\text{CFC-12/SF}_6 \text{ mode-age}) + 2.5 \text{ years}$ ($R^2 = 0.98$). The
492 1:1 reference line (dashed) is crossed at a CFC-12/SF₆ mode-age of ~31.25 years, corresponding to a ³⁹Ar mode-age of the
493 same value. This crossing point (~30 years) marks the threshold below which the two methods agree within uncertainties. For
494 mode-ages > ~30 years, CFC-12/SF₆ ages are systematically higher than ³⁹Ar ages, as indicated by the slope of 0.92, suggesting
495 overestimation by CFC-12/SF₆ in deep, old waters due to tracer concentrations approaching detection limits. For younger
496 waters (<30 years mode-age), the two methods agree within uncertainties, although for such young waters ³⁹Ar is not an ideal
497 tracer. The high R^2 confirms overall consistency, while the offset at high ages demonstrates the complementary value of ³⁹Ar
498 for dating waters on centennial-to-millennial timescales. We note that the comparison assumes steady-state ventilation over
499 the ~20-year temporal mismatch between the two datasets; changes in circulation over that period may contribute to the bias.

As expected, the inferred ages of water masses differ depending on the tracer employed. These discrepancies arise not only from potential limitations in the assumed IG-TTD, but also from the distinct temporal validity ranges and detection limits of each tracer. Our comparative analysis (as summarized in Table 2 and Fig. 18) reveals a systematic pattern: mean- and mode-ages derived from ^{39}Ar are generally lower than those estimated from CFC-12 and SF_6 for deeper/older water masses. This is exemplified by the NADW in the eastern basin near 20°N , where CFC-12/ SF_6 yield a mean-age of ~ 1000 years and a mode-age of ~ 150 years, whereas ^{39}Ar suggests younger ages of ~ 600 and ~ 90 years, respectively. Similarly, for AABW/NEABW near 50°N , CFC-12 and SF_6 based estimates (~ 800 years mean, ~ 120 years mode) exceed those from ^{39}Ar (~ 500 years mean, ~ 100 years mode). A likely explanation for this systematic tendency towards higher age estimates from CFC-12 and SF_6 in deep and bottom layers might be attributed to their extremely low concentrations at great depths and for older waters. These concentrations often approach or fall below analytical detection limits, which can amplify uncertainties and appear to skew age calculations towards higher values. In contrast, the ^{39}Ar with 269-year half-life appears to be better suited for dating waters with transit times on the order of centuries to a millennium, and thus could provide relative more robust age estimates for these older water masses, as its slower decay rate remains measurable over such longer timescales.

6 Conclusions and discussion

The geographic distributions of main water masses in the Atlantic Ocean are reported in Liu and Tanhua (2021). In the present work, the transports of water masses are traced by transient tracers (CFC-12, SF_6 and ^{39}Ar) and their ventilation timescales (water mass ages) are quantified. Three distinct concepts of water mass age are discussed within the analytical framework. While the tracer-age concept (assuming purely advective transport) is introduced for context, the quantitative analysis and results presented herein are based on the mean-age and mode-age derived from the Transit Time Distribution (TTD) model. The application of a standard mixing ratio ($\Delta/\Gamma = 1$) and 100% surface saturation is followed. The mean-age, representing the average transit time of all water parcels in a sample and providing an integrative timescale, is used to characterize the overall ventilation status. The mode-age, representing the most probable transit time and serving as a proxy for advective timescales, is applied to trace the dominant transport pathways and to estimate process rates like the oxygen utilization rate (OUR).

In general, the water mass age increases with the depth and distance from the formation area, so the central waters in the upper layer obtain the lowest ages of within ~ 100 years. In the intermediate layer, the AAIW and MW are two conspicuous water masses and both take ~ 300 years to spread from south to north (AAIW) and east to west (MW) respectively. As the dominant water mass in the deep and overflow layer, the NADW takes ~ 500 years to spread along the DWBC in the west Atlantic Ocean, and takes ~ 900 years eastward to cover the east part. The bottom waters origin from south of Antarctic Circumpolar Current region take ~ 600 years to spread northward and cover the bottom layer. Similar as the NADW, the mean-age of both AABW and NEABW show spatial difference between the west (low mean-age) and the east (high mean-age). This is because the west Atlantic Ocean is well ventilated by the DWBC while the water exchange is sluggish in the east.

532 The transient tracers (CFC-12, SF₆ and ³⁹Ar), serve as effective tools for identifying and quantifying oceanographic ventilation,
533 each tracer possessing distinct advantages and limitations in determining and tracing the depths and mean ages of water masses.
534 CFC-12 and SF₆ exhibit optimal applicability for water masses range within 2000 dbar pressure (or mean-ages < ~500 years).
535 However, their utility becomes constrained in deeper aqueous regimes (below 2000 dbar or mean-age > ~500 years) due to
536 inherent limitations in tracer concentration sensitivity, which may induce systematic overestimations of water mass ages when
537 applied to deep or bottom layers. The chemically inert radioisotope ³⁹Ar, characterized by a 269-year half-life, effectively
538 addresses this observational gap through its unique temporal resolution. Consequently, the synergistic application of SF₆, CFCs,
539 and ³⁹Ar in multi-tracer frameworks overcomes temporal constraints inherent to single-tracer approaches, enabling
540 comprehensive investigation of coupled biogeochemical-climate interactions. This integrated methodology facilitates novel
541 insights into climate-driven water mass reorganization, carbon cycle dynamics, and other critical marine processes essential
542 for understanding oceanic responses to global change.

543 The ventilations and respiration rates (oxidation rates, OUR) of wide-spread water masses are estimated by the mode-ages.
544 The currents and topography have significant impacts on the mode-ages and furthermore the apparent oxygen utilization rate
545 (OUR). The western basins of the Atlantic Ocean are better ventilated with lower the mode-ages, while the OURs show the
546 opposite distribution, suggesting that the mode-age is a more important determinant of OUR than AOU, and that higher
547 respiration rates exist in better-ventilated regions.

548 The quantitative age framework and ventilation rates established in this study provide a crucial baseline dataset. This baseline
549 is essential for detecting future changes in ocean circulation and biogeochemical cycles, and will serve to validate the next
550 generation of climate and ocean models.

551 This study provides a quantitative, basin-scale assessment of Atlantic water mass ages using a multi-tracer approach. However,
552 several methodological assumptions should be considered when interpreting the results. First, the calculated ages rely on the
553 assumption of a steady-state circulation and an Inverse Gaussian Transit Time Distribution (IG-TTD) with a fixed mixing
554 ratio ($\Delta/\Gamma = 1$). While this is a necessary and common simplification for basin-scale analysis, regional deviations from this
555 idealized transport are likely. Consequently, the mean-age and mode-age presented in this study are linearly related ($\tau_{\text{mode}} \approx$
556 $0.162 \cdot \Gamma$) and do not provide independent statistical information; they are shown together solely for conceptual comparison of
557 their different physical interpretations (integrative vs. advective timescales). Second, the complementary use of tracers
558 (CFC-12, SF₆, ³⁹Ar) mitigates but does not fully eliminate the inherent uncertainties associated with each method, particularly
559 in deep layers where tracer concentrations are very low. Third, the apparent oxygen utilization rates (OUR) derived here
560 represent integrated rates along flow paths and are subject to the same age estimation uncertainties. Moreover, the choice of
561 using mode-age for OUR calculations is a working assumption; the true local OUR remains difficult to constrain. Fourth, the
562 ³⁹Ar data used for comparison originate from the 1980s–1990s (Rodriguez, 1993), whereas the CFC-12/SF₆ data are from later
563 cruises (predominantly 2000s–2010s). The comparison thus implicitly assumes a steady-state circulation and ventilation,
564 which is a simplification given observed multidecadal changes in the Atlantic Ocean (e.g., Talley et al., 2016). This temporal

565 mismatch introduces additional uncertainty into the age comparisons, particularly for older water masses where ventilation
566 may have varied over the intervening decades (e.g. Guo et al. 2026).

567 Notwithstanding these limitations, the comprehensive age and ventilation maps presented here establish a baseline of the
568 contemporary Atlantic Ocean ventilation structure. This dataset can serve to validate and constrain the next generation of ocean
569 circulation and biogeochemical models. Furthermore, by quantifying current ventilation timescales and their spatial patterns,
570 this work provides a reference frame against which future observational studies can detect changes in ocean circulation and
571 ventilation in response to climatic forcing.

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573 wrote the paper.

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