OCEANIC RADIOCARBON BETWEEN ANTARCTICA AND SOUTH AFRICA ALONG WOCE SECTION I6 AT 30°E

Viviane Leboucher¹ • James Orr² • Philippe Jean-Baptiste² • Maurice Arnold¹ • Patrick Monfray¹ Nadine Tisnerat-Laborde¹ • Alain Poisson³ • Jean-Claude Duplessy¹

ABSTRACT. Accelerator mass spectrometry (AMS) radiocarbon measurements were made on 120 samples collected between Antarctica and South Africa along 30°E during the WOCE-France CIVA1 campaign in February 1993. Our principal objective was to complement the Southern Ocean's sparse existing data set in order to improve the ¹⁴C benchmark used for validating ocean carbon-cycle models, which disagree considerably in this region. Measured ¹⁴C is consistent with the θ-S characteristics of CIVA1. Antarctic Intermediate Water (AAIW) forming north of the Polar Front (PF) is rich in ¹⁴C, whereas surface waters south of the PF are depleted in ¹⁴C. A distinct old ¹⁴C signal was found for the contribution of the Pacific Deep Water (PDW) to the return flow of Circumpolar Deep Waters (CDW). Comparison to previous measurements shows a ¹⁴C decrease in surface waters, consistent with northward displacement of surface waters, replacement by old deep waters upwelled at the Antarctic Divergence, and atmospheric decline in ¹⁴C. Conversely, an increase was found in deeper layers, in the AAIW. Large uncertainties, associated with previous methods for separating natural and bomb ¹⁴C when in the Southern Ocean south of 45°S, motivated us to develop a new approach leads to inventories for CIVA1 that are equal to or higher than those calculated with previous methods. Differences between old and new methods are especially high south of approximately 55°S, where bomb ¹⁴C inventories are relatively modest.

INTRODUCTION

The Southern Ocean extends south to Antarctica and north to the Subtropical Front (STF), which meanders between 40°S and 50°S. As such, the Southern Ocean covers about 21% of the world's ocean surface. Within the Southern Ocean, large vertical exchanges take place between water masses. Important processes include ventilation of deep waters at the Antarctic Divergence Zone (ADZ, ca. 65°S), production of ventilated bottom waters in the Weddell Sea and around the Antarctic continental slope, and formation of Antarctic Intermediate Water (AAIW) within the Polar Frontal Zone (PFZ) between the Polar Front (PF) and the Subantarctic Front (SAF).

The Southern Ocean is considered to be a major sink for anthropogenic CO_2 (Sarmiento et al. 1992). However, for this region, there is substantial disagreement between carbon-cycle simulations from different ocean general circulation models (OGCMs), which offer the only means to estimate future oceanic CO_2 uptake (Orr 1996). Radiocarbon is routinely used to evaluate the performance of such models (Toggweiler et al. 1989a, 1989b; Maier-Reimer 1993; Taylor 1995; Orr 1996). However, few ¹⁴C data are available in the remote Southern Ocean, where both the spatial and temporal changes in ¹⁴C are important. Here, we provide ¹⁴C data to complement the sparse established dataset. Previous ¹⁴C measurements include those from GEOSECS (Geochemical Ocean Sections Study) during 1973–1978 (Östlund and Stuiver 1980; Stuiver and Östlund 1980; Stuiver et al. 1981), from the Winter Weddell Sea Experiment (WWSP) in 1986 (Schlosser et al. 1994), and from INDIGO (Indian Gas Ocean project) in 1985–1987 (Östlund and Grall 1988). The ¹⁴C data presented here were collected during CIVA1 (Circulation et Ventilation dans l'Antarctique) of WOCE-France taken in February– March 1993. Measurements of ¹⁴C were made by AMS in Gif-sur-Yvette, France.

¹Laboratoire des Sciences du Climat et de l'Environnement (LSCE), Avenue de la Terrasse, F-91198 Gif-sur-Yvette Cedex, France

²Laboratoire des Sciences du Climat et de l'Environnement (LSCE), DSM/CEN Saclay/CEA, L'Orme, Bât. 709, F-91191 Gif-sur-Yvette Cedex, France

³Laboratoire de Physique et Chimie Marines (LPCM), Université Pierre et Marie Curie, 4, place Jussieu, F-75252 Paris Cedex 05, France

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Oceanic ¹⁴C must be separated into its natural and anthropogenic components if it is to be useful for model validation. Natural ¹⁴C is produced by cosmic radiation and can be considered at present to be in steady state with respect to the vertical turnover time of the ocean (ca. 1000 yr). Bomb ¹⁴C was injected in the atmosphere nearly exclusively by atmospheric thermonuclear weapons tests during the 1950s and 1960s. At its peak in 1963, the nuclear contribution nearly doubled prenuclear atmospheric levels of ¹⁴C. Since then, atmospheric ¹⁴C has decreased. At present, levels are roughly 10% higher than in the prenuclear period. Both ¹⁴C components are transferred to the ocean by air-sea gas exchange as ¹⁴CO₂. With its steady-state production and 5730 yr half-life, the natural component of ¹⁴C is useful for evaluating modeled deep-ocean circulation fields. Bomb ¹⁴C has had much less time to invade the ocean. Because its boundary conditions have changed rapidly since the late 1950s, bomb ¹⁴C is useful for evaluating near-surface circulation fields predicted by ocean models.

To separate natural and bomb ¹⁴C, we first applied the method developed by Broecker et al. (1995), which relies on the near linear correlation between natural Δ^{14} C and dissolved silica throughout much of the ocean. However, for the stations situated south of 45°S (~90% of samples collected here), uncertainties associated with Broecker et al.'s method are large. Since the GEOSECS campaign in the 1970s, it has become less certain that deep waters remain uncontaminated with bomb ¹⁴C, as supposed by Broecker et al. (1995) for GEOSECS era samples. To take this possibility into account, we modified the bomb versus natural separation methodology for our samples taken south of 45°S. Our new method takes advantage of CFC-11 and CFC-12 measured during CIVA1 and uses a simple ventilation model, similar to that used in previous tracer studies (Jenkins 1980; Andrié et al. 1986; Haine 1996).

METHODS

The ¹⁴C data (Table 1) were obtained following the procedure described by Bard et al. (1987) and Arnold et al. (1987). Seawater was collected in 12-L Niskins. Once aboard ship, seawater was transferred to 250 mL borosilicate bottles and poisoned with 1 mL of saturated HgCl₂ solution. The borosilicate bottles were then sealed. Back in the laboratory, CO₂ was extracted by adding 2 mL of 15N H₃PO₄ to a 100 mL seawater aliquot in a vacuum-tight system sparged by He gas (flow rate 80 mL min⁻¹). Extraction required 1 h. Water was removed by traps at -80° C; CO₂ was trapped at -180° C. Subsequently, CO₂ was reduced to graphite using hydrogen in the presence of iron powder (6–8 h). The carbon-iron mixture was divided into 3 parts, then pressed into an aluminum target for accelerator mass spectrometry (AMS) analysis. Usually, 2 or 3 targets per sample were analyzed to obtain a precision close to ±5%. The 3 isotopes, ¹²C, ¹³C and ¹⁴C, were measured directly in the AMS system to calculate Δ^{14} C normalized to a δ^{13} C of -25%. The data are expressed as Δ^{14} C in per mil (Stuiver and Polach 1977).

RESULTS

To establish the level of agreement of our ¹⁴C CIVA1 measurements with previous data, Figure 1 provides ¹⁴C profiles from station 442 of GEOSECS (1978) and from station 1 of JADE1 (Java Australia Dynamic Experiment 1989) cruise, both taken at the same location (1°S, 91°E) in the equatorial Indian Ocean. The ¹⁴C measurements from JADE1 were made in Gif-sur-Yvette by AMS just prior to those of CIVA1. JADE1 and CIVA1 samples received identical processing. The deep portions of GEOSECS and JADE1 profiles, where bomb ¹⁴C has not penetrated, agree to within the precision of the measurements. Therefore, our ¹⁴C measurements by AMS are consistent with those from GEOSECS, determined by β -counting. Furthermore, the methodology used at Gif-sur-Yvette has been previously checked by directly comparing β -counting and AMS measurements performed on the same samples from INDIGO (Bard et al. 1988).

Table 1	Data for	the stat	ions oc	cupied dur	ing the CIV	VA1 cruise					
Depth (m)	Potential temp. (°C)	Salinity (S, psu)	Sig 0	SiO ₂ (µmol kg ⁻¹)	SCO ₂ (mmol kg ⁻¹)	CFC-11 (pmol kg ⁻¹)	CFC-12 (pmol kg ⁻¹)	Num. of meas.	Δ ¹⁴ C (%c)	Nat. Δ ¹⁴ C (‰)	Bomb Δ ¹⁴ C (%e)
Station 6 10.2 19.9 40 100.6 150.5 199.7 324.4 698.8 998.8 1198.8 2247.1 2497.6 2997.6 3495.9 3681.3	$\begin{array}{c} 68^\circ S, \ 30^\circ E \\ 0.2009 \\ 0.1145 \\ -0.1182 \\ -1.7636 \\ -1.6967 \\ -0.5536 \\ 0.7518 \\ 0.7136 \\ 0.7136 \\ 0.4484 \\ 0.2899 \\ -0.2104 \\ -0.2729 \\ -0.378 \\ -0.4769 \\ -0.5667 \end{array}$	33.915 33.919 33.933 34.35 34.371 34.468 34.642 34.683 34.683 34.665 34.665 34.655 34.655	27.218 27.225 27.249 27.651 27.665 27.708 27.773 27.846 27.829 27.833 27.847 27.847 27.847 27.851 27.851	$\begin{array}{c} 62.25\\ 62.25\\ 63.06\\ 68.77\\ 70\\ 83.5\\ 99.93\\ 115.4\\ 124.7\\ 130.9\\ 136.3\\ 135.6\\ 136.3\\ 137.1\\ 137.1\\ 137.1 \end{array}$	2149 2152 2160 2217 2210 2240 2247 2254 2266 2264 2260 2258 2253 2253 2254 	6.76 6.72 6.7 5.68 5.59 3.59 1.14 0.58 0.46 0.41 0.56 0.64 0.91 1 0.99	$\begin{array}{c} 2.92\\ 2.93\\ 2.86\\ 2.47\\ 2.45\\ 1.52\\ 0.46\\ 0.23\\ 0.18\\ 0.17\\ 0.21\\ 0.24\\ 0.36\\ 0.41\\ 0.4 \end{array}$	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	$\begin{array}{c} -97 \pm 4 \\ -97 \pm 4 \\ -98 \pm 4 \\ -114 \pm 4 \\ -165 \pm 4 \\ -157 \pm 4 \\ -157 \pm 4 \\ -157 \pm 4 \\ -158 \pm 4 \\ -158 \pm 4 \\ -153 \pm 4 \\ -145 \pm 7 \\ -145 \pm 7 \\ -157 \pm 4 \\ -162 \pm 4 \end{array}$	-132 -132 -133 -139 -140 -154 -163 -163 -163 -163 -159 -153 -167 -171	35 36 35 25 23 13 11 6 5 4 6 6 9 10 10
Station 8 10.6 20.2 40 80.8 124.7 175.9 322.1 598.4 795 997.1 1195.8 1795.5 2242.4 2740.9 3193.9 3697.2 4065.7 4065.9	$67^{\circ}S, 30^{\circ}E$ 0.646 0.6406 -1.505 -1.7041 -1.142 0.4412 1.1479 1.0146 0.7838 0.5854 0.3901 0.0023 -0.795 -0.3279 -0.4247 -0.5206 -0.6255 -0.6265	33.866 33.866 33.869 34.249 34.356 34.412 34.564 34.679 34.708 34.708 34.708 34.667 34.667 34.667 34.667 34.652 34.653 34.655 34.656 34.652 34.653	27.155 27.155 27.568 27.653 27.682 27.729 27.778 27.812 27.823 27.834 27.834 27.844 27.844 27.846 27.855 27.855 27.855 27.856	$\begin{array}{c} 60\\ 60\\ 60.77\\ 65.5\\ 71.02\\ 86.01\\ 92.69\\ 102.5\\ 110.1\\ 115.4\\ 118.4\\ 124.8\\ 128.6\\ 127.8\\ 126.3\\ 127.1\\ 129.3\\ 130.9\\ \end{array}$	2158 2164 2197 2216 2228 2255 2266 2263 2267 2250 2267 2270 2263 2260 2258 2260 2258 2260 2258	$\begin{array}{c} 6.58 \\ 6.57 \\ 5.7 \\ 5.43 \\ 4.62 \\ 2 \\ 0.74 \\ 0.39 \\ 0.32 \\ 0.27 \\ 0.33 \\ 0.53 \\ \hline \\ 0.69 \\ 0.94 \\ 0.98 \\ 0.91 \\ 0.94 \\ \end{array}$	$\begin{array}{c} 2.89\\ 2.84\\ 2.46\\ 2.36\\ 1.95\\ 0.85\\ 0.29\\ 0.15\\ 0.13\\ 0.09\\ 0.12\\ 0.12\\ 0.28\\ 0.39\\ 0.42\\ 0.36\\ 0.36\end{array}$	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	$\begin{array}{c} -97 \pm 4 \\ -97 \pm 5 \\ -104 \pm 4 \\ -103 \pm 4 \\ -111 \pm 4 \\ +130 \pm 5 \\ -155 \pm 5 \\ -155 \pm 5 \\ -175 \pm 4 \\ -179 \pm 5 \\ -175 \pm 4 \\ -154 \pm 4 \\ -154 \pm 4 \\ -166 \pm 8 \\ -160 \pm 4 \\ -167 \pm 4 \\ -150 \pm 4 \\ -150 \pm 4 \\ -154 \pm 4 \end{array}$	$\begin{array}{c} -130\\ -130\\ -131\\ -136\\ -141\\ -152\\ -152\\ -152\\ -152\\ -157\\ -159\\ -160\\ -179\\ -160\\ -176\\ -159\\ -163\end{array}$	33 33 27 33 30 26 7 4 4 0 3 4 6 7 9 9 10 9 9
Station 12 10.6 20.1 39.4 99.8 149.7 200.1 324.5 499.8 698.9 998 1197.1 2000 3000 4495.9	2 65°S, 30° 1.004 1.0006 0.6738 -0.9583 0.9763 1.2006 1.2749 1.0755 0.749 0.5752 -0.6169	E 33.829 33.829 33.842 34.411 34.57 34.615 34.675 34.715 34.718 34.706 34.699 34.674 34.663 34.652	27.1 27.1 27.15 27.673 27.721 27.74 27.799 27.813 27.827 27.831 27.856	54.24 53.44 54.24 70.95 85.3 89.28 95.66 102 	2164 2155 2168 2225 2259 2261 2255 2264 2266 2257 2259 2264 2261 2253	$\begin{array}{c} 6.45\\ 6.44\\ 6.46\\ 4.37\\ 1.84\\ 1.29\\ 0.72\\ 0.3\\ 0.24\\ 0.2\\ 0.2\\ 0.2\\ 0.2\\ 0.4\\ 0.63\\ \end{array}$	$\begin{array}{c} 2.79\\ 2.79\\ 2.81\\ 1.87\\ 0.75\\ 0.49\\ 0.26\\ 0.07\\ 0.05\\ 0.04\\ 0.03\\ 0.04\\ 0.11\\ 0.22 \end{array}$	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	$\begin{array}{c} -117\pm 5\\ -97\pm 4\\ -97\pm 5\\ -125\pm 5\\ -125\pm 4\\ -144\pm 4\\ -148\pm 4\\ -152\pm 4\\ -158\pm 4\\ -158\pm 4\\ -161\pm 4\\ -164\pm 5\\ -155\pm 4\\ -164\pm 5\end{array}$	-125 -123 -124 -141 -155 -159 -155 -152 -158 -152 -161 -164 -158 -170	$ \begin{array}{c} 8\\ 26\\ 27\\ 16\\ 16\\ 15\\ 7\\ 0\\ 0\\ 0\\ 0\\ 0\\ 4\\ 6\\ \end{array} $
Station 1 11.3 20.9 40.7 80 126.5 175.2 252.1 599.1 799.7 998.9 1197.6 2244.5 2995.5 4495.7	$\begin{array}{c} 8 & 61^{\circ}\text{S}, 30^{\circ}\\ 1.7759 \\ 1.6254 \\ -0.0172 \\ -1.62 \\ -1.213 \\ 0.7429 \\ 1.4748 \\ 1.2922 \\ 1.0488 \\ 0.8452 \\ 0.6426 \\ 0.0032 \\ -0.2888 \\ -0.6071 \end{array}$	2E 33.686 33.721 33.891 34.122 34.198 34.482 34.635 34.718 34.714 34.709 34.7 34.673 34.664 34.655	26.938 26.973 27.211 27.463 27.509 27.646 27.72 27.799 27.812 27.823 27.828 27.828 27.843 27.849 27.857	44.3 44.72 45.56 50.6 57.35 82.22 90.64 100.3 107.1 112.1 116.4 127.3 127.3 129	2136 2140 2152 2176 2196 2243 2250 2260	$\begin{array}{c} 6.32 \\ 6.33 \\ 6.41 \\ 6.11 \\ 5.56 \\ \hline \\ 0.85 \\ 0.36 \\ 0.31 \\ 0.24 \\ 0.21 \\ 0.22 \\ 0.24 \\ 0.26 \end{array}$	2.83 2.85 2.85 2.74 2.44 0.13 0.15 0.11 0.1 0.09 0.07 0.1 0.09	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	$\begin{array}{c} -74\pm 5\\ -85\pm 5\\ -80\pm 5\\ -80\pm 5\\ -91\pm 5\\ -124\pm 5\\ -146\pm 5\\ -164\pm 5\\ -164\pm 5\\ -162\pm 5\\ -156\pm 5\\ -157\pm 5\\ -154\pm 5\end{array}$	-114 -116 -121 -127 -152 -155 -146 -169 -164 -162 -156 -157	$\begin{array}{c} 40\\ 30\\ 36\\ 34\\ 36\\ 28\\ 8\\ 4\\ 3\\ 0\\ 0\\ 0\\ 0\\ 0\\ 0\\ 0\\ 0\\ 0\\ 0\\ 0\\ 0\\ 0\\$
Station 2 10.4 19.8 40.7 99 150.5 199.3 320.7	24 57°S, 30 2.2178 2.2133 2.2012 -0.7262 -1.6046 -1.4569 0.5166	°E 33.626 33.625 33.633 33.92 34.141 34.234 34.596	26.853 26.853 26.862 27.267 27.476 27.549 27.751	40.6 40.6 50.32 68.97 73.03 99.78	2131 2139 2140 2202 2209 2256	6.27 6.27 6.26 6.91 6.62 6.08 1.6	2.84 2.82 2.84 3.06 2.91 2.68 0.7	2 2 2 2 2 2 2 2 2 2	$ \begin{array}{r} -83 \pm 5 \\ -77 \pm 5 \\ -74 \pm 5 \\ -81 \pm 5 \\ -87 \pm 5 \\ -92 \pm 5 \\ -141 \pm 5 \end{array} $	-111 -111 -120 -139 -143	

Table 1 Data for the stations occupied during the CIVA1 cruise

Table 1 Data for the stations occupied during the CIVA1 cruise (Continued)

	· · · · · · · · · · · · · · · · · · ·		nomb ot	Joupieu uu	8		Commu	<u></u>			
_	Potential			au -				Num.		Nat.	Bomb
Depth	temp.	Salinity		SiO ₂	SCO_2	CFC-11	CFC-12	of	$\Delta^{14}C$	$\Delta^{14}C$	$\Delta^{14}C$
(m)	(°C)	(S, psu)	Sig 0	(µmol kg ⁻¹)	(mmol kg ⁻¹)	(pmol kg ⁻¹)	(pmol kg ⁻¹)	meas.	(%0)	(%0)	(%c)
496.9	0.9574	34.695	27.803	108.3	2263	0.45	0.18	2	-153 ± 5	-157	5
696.1	0.7337	34.695	27.818	114.4	2267 2268 2268	0.38	0.15	2 2 2 2 2 2 2 2 2 2 2	-153 ± 5 -151 ± 5	-157	J 4
997.1	0.5577	34.695	27.829	120.9	2268			2	-151 ± 5	-153	4 3
1195.8	0.4318	34.69	27.833	125.3	2268	0.23	0.11	2	-175 ± 5	-177	3
1992.9	0.0367	34.675	27.842	129		0.19	0.07	$\tilde{2}$	-177 ± 5	~177	õ
2995.3	-0.3072	34.663	27.85	130.6		0.22	0.08	$\overline{2}$	-171 ± 4	-171	ŏ
4495.2	-0.6165	34.654	27.856	129.4	2258 2256	0.23	0.09	2	-171 ± 4 -156 ± 4	-156	ŏ
5422.2	-0.7349	34.65	27.858	129	2256	0.28	0.11	2	-150 ± 4	-153	3
Station 28	54.3°S, 3	О°Е									
10.9	2.8814	33.876	26.998	26.37	2147	5.91	2.72	2	-69 ± 6	-96	28
19.9	2.8829	33.877	26.998	26.37	2147	5.1	2.53	2	-81 ± 5	-96	16
19.9 39.5 101.2	2.7778	33.877 33.952	27.066	29.66	2155	5 94	2.72	2	-83 ± 5	-100	17
101.2	0.0656	34.096	27.378	48.6	2190	5.85	2.67	2	-97 ± 5	-119	17 21
151.1 197.2	0.6092	34.294	27.456	55.18	2210	4.81	2.09	$\overline{2}$	-80 ± 5	-125	45
197.2	1.3819	34.366	27.511	65.07	2210 2239 2262	2.54	1.08	$\overline{2}$	-109 ± 5	-135	45 26
323.1 496.7	1.9407	34.572	27.632	78.23	2262	0.81	0.33	2	-142 + 5	-148	7
496.7	1.8816	34.658	27.707	85.23	2262	0.47	0.17	2	-150 ± 5 -153 ± 5	-155	7 5 5 0
701.6	1.6213	34.689	27.752	91.4	2263	0.49	0.21	2	-153 ± 5	-158	5
995.2 1399.7	1.6213 1.5408 1.0201	34.737	27.795	91.81	2257	0.25	0.08	2	-147 ± 4	-147	0
2494.4	1.0201	34.68	27.823	105.4	2262 2263 2257 2261 2261	0.22	0.09	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	-156 ± 4	-156	0
	0.1553 -0.4818	34.68 34.658	27.841	126.4	2261	0.25	0.07	2	-164 ± 5	-164	0
3993.7 4996.3	-0.4818	34.658 34.657	27.853	130.1	2262	0.21	0.08	2	-164 ± 4	-164	0
	-0.6523	34.657	27.858	129.7		0.21	0.09	2	-156 ± 4	-156	0
			27.858	128.4	2260	0.19	0.08	2	-151 ± 4	-151	0
Station 38											
9.3 18.8	4.5363	33.874	26.835	3.03	2121	5.29	2.95	2	-52 ± 5	73	22
18.8	4.5366	33.874	26.836	3.89	2120	5.33	2.89	2	-50 ± 5	74	24 29 58 43 59
38.2	4.5302	33.876	26.836	5.19 15.12	2119	5.3	2.88	2	-47 ± 5	-75	29
79 125.9	3.139 2.6988	33.982	27.061	15.12	2150	5.3	2.84	2	-27 ± 5	-85	58
172.9	2.6988	34.011	27.123	18.15	2154	5.12	2.74	2	-45 ± 5	-88	43
325.4	2.6227	34.125 34.199	27.173	20.31	2162	4.11	2.2	2	-31 ± 6	-90	59
576.9	2.3437	34.199 34.36	27.28	34.17	2188	2.96	1.56 0.67	2	-61 ± 6	-104	43 26
795.4	2.3437 2.4431	34.50 34.525	27.433 27.556	56.5	2233 2254	1.34	0.67	2	-100 ± 5	-127	26
998	2.4451	34.525	27.556	70.07 75.32	2254	0.51	0.26	2	-130 ± 5	-140	10
1198	2.3759 2.3707	34.685	27.652	75.32	2258	0.23	0.09	1	-141 ± 8	-141	0
2498.4	1.1142	34.735	27.825	105.6	2258 2255 2272	0.16	0	2	-131 ± 5	-131	0
3994.5	-0.1069	34.672	27.848	132.2	2272	0.08	0	2	-158 ± 5	-158	0
5183.5	-0.4829	34.656	27.848	135.2	2231	0.25 0.21	0.12 0.08	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	-156 ± 5 -153 ± 5	-156	Õ
			21.055	155.2	_	0.21	0.08	2	-153 ± 5	-153	0
Station 52 9.8	44°S, 30°. 12.315		06 105	1.07							
9.8 19.4	12.315	34.475 34.472	26.125 26.128	1.96		3.45	1.82	2	43 ± 5	-72	115
39.7	12.297 12.045	34.472 34.45		1.96	2061	3.41	1.87	2	49 ± 5	-72	121
100.5	9.2985	34.45 34.596	26.156	1.96	2058	3.47	1.95	2	53 ± 5	-72	125
142.6	9.2985 8.7958	34.596 34.594	26.756	7.34	2116	3.62	1.95 1.89 1.79 1.75	2	55 ± 5	-77	132
142.0	0.7930	34.394 34.497	26.835 26.892	8.31	2123	3.43	1.79	2	34 ± 5	-78	112
323 3	7.916 6.3703	34.341	20.092	9.78 13.71	2135 2143	3.36 3.29	1.75	2	33 ± 5	-80	113
323.3 499 698.7	5 1576	34.341	26.983 27.128	23.10	2143		1.62	2	11 ± 5	-84	94 65 45
698.7	5.1576 4.0819	34.329	27.128	23.19	2100	2.2 1.62	1.09	2	-29 ± 5	-93	65
1001.2	3.0136	34.421	27.423	58.5	2196 2239	0.73	0.8 0.32	2	-61 ± 5	-106	45
1198.1	3.0136 2.7672	34 53	27 532	68.5	2239	0.73 0.28	0.32	2	-125 ± 5	-125	0
1198.1 2244.9	2.2081	34.793	27.532 27.789	23.19 35.84 58.5 68.5 72.65	2251 2236	0.28	0.16	2	-142 ± 4	-142	0
3496.4	2.2081 1.0783	34.753	27.841	104.5	2250	0.04	0	2	-137 ± 4	-137	0
4497.9	0.1862	34.69	27.846	132	2207	0.04	0.1	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	-153 ± 4 -160 ± 4	-153	0 0
						0.15	0.1	4	-100 ± 4	-160	0

Figure 2 shows a map including the WOCE/CIVA1 transect between 68°S and 44°S along 30°E, where ¹⁴C samples were collected. Also indicated are other stations in the same area where ¹⁴C has been measured on previous campaigns: GEOSECS, WWSP, and INDIGO.

The θ and S along the CIVA1 transect (Fig. 3A,B)⁴ exhibit signatures of the major fronts that characterize the Southern Ocean: the Antarctic Divergence Zone (ADZ) between 63°S and 67°S, the Weddell Gyre Front (WF) at 57°S, the Polar Front (PF) at 52°S, and the Subantarctic Front (SAF) at 48°S (Gordon 1971; Whitworth and Nowlin 1987; Orsi et al. 1993, 1995; Archambeau et al. 1998). The

⁴Note: Color versions of Figs. 3, 4, 7 and 9 are available at http://www.radiocarbon.org/Journal/v41n1/Leboucher/

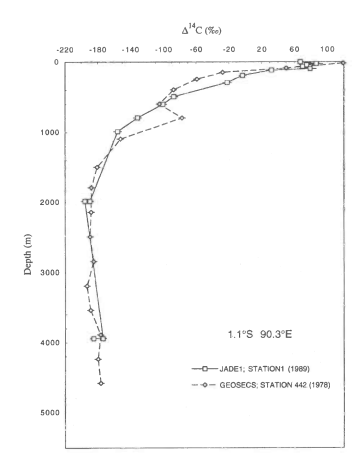


Figure 1 Profiles of measured Δ^{14} C from GEOSECS station 442 and JADE1 station 1

principal water masses are likewise apparent. From the Antarctic continent to the PF, the Summer Surface Water (SSW) overlies the near freezing Winter Water (WW). Below the WW lies a massive amount of Circumpolar Deep Water (CDW), which is relatively warm (>0.1 °C) and old (with respect to ventilation by contact with the surface). Beneath the CDW and along the bottom is found the Antarctic Bottom Water (AABW). In the AABW, one can trace the influence of the Weddell Sea Bottom Water (WSBW), which at its origin (the Weddell Sea) has a temperature <-0.7 °C. Another water mass, the Subantarctic Surface Water (SASW), lies between the PF and the SAF. The SASW has higher temperatures and salinities than the more southern surface waters and is remarkable for its south-to-north downward tilt in θ and S isolines, where AAIW is formed. In this northern part of the CIVA1 section, intermediate level CDW bears a high salinity signature (maximum 34.8 psu) from its contact with North Atlantic Deep Water (NADW). AABW north of the Atlantic-Indian ridge is warmer and less dense than AABW to the south, which is closer to the source of the cold WSBW. Further description of CIVA1 hydrography can be found in Archambeau et al. (1998).

Table 1 provides the CIVA1 ¹⁴C measurements. These data are presented as a composite latitude versus depth section (Fig. 4A) as well as individual profiles (Fig. 5A,B). The former reveals that the distribution of the measured ¹⁴C roughly follows CIVA1 hydrography. A striking contrast is found across the PF: north of the PF, surface waters are enriched in ¹⁴C (levels reach 55‰); south of the PF, the upper 200 m are depleted in ¹⁴C (levels from -70% to -140%). The Southern depletion is principally due to old CDW upwelling in the Antarctic Divergence Zone between 63°S and 67°S (Toggweiler and Wallace 1995). However, winter ice cover limits gas exchange and thus may play

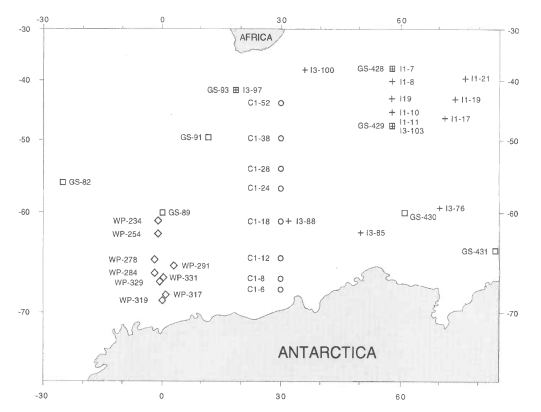


Figure 2 Stations occupied to collect samples for ¹⁴C during CIVA1 (C1) as well as adjacent ¹⁴C stations from GEO-SECS (GS), WWSP (WP), INDIGO (I1 and I3)

some role (Weiss et al. 1979; Poisson and Chen 1987). North of the PF, ¹⁴C-enriched surface waters sink below warmer subantarctic water and move toward the north along downward sloping isopycnal surfaces, where AAIW is formed.

A core of old water (Δ^{14} C less than -170%) is located near 57°S between 1000 and 2000 m (station 24). This corresponds to the oldest CDW. A shallow extension of old CDW is found near 66°S (station 8) with the oldest values (less than -175%) between 800 m and 1000 m.

End members for and mixing between intermediate, deep, and bottom waters ($\sigma_{\theta} < 27.7$) are more easily seen on CIVA1's θ -S diagram (Fig. 6A). The core of the CDW is identifiable by its salinity maximum induced by the contribution of NADW. Another influential water mass is the AABW, which forces most of the data points on the θ -S plot to lie on a straight line between the CDW and AABW end-member. The ¹⁴C data are helpful to pinpoint a third water mass component. Points from stations 8 and 24 reveal that the most ¹⁴C-depleted waters have slightly different θ -S characteristics (Figs. 6B, 6C). These tendencies point to the older ¹⁴C-depleted Pacific Deep Water (PDW) as an important contributor. The same trend was found from measurements of ³He in the Southern Ocean (Jean-Baptiste et al. 1991). The contribution of PDW to the CDW is also consistent with the return Pacific flow scheme described by Toggweiler and Samuels (1993) based on Pacific-Antarctic Δ^{14} C difference, on the silica distribution in sediments of the Pacific sector of the Southern Ocean (Worthington 1977), and on previous inverse model results (Wunsch et al. 1983). The WOCE sec-

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tion P6 along 32°S in the Pacific Ocean shows a substantially older 14 C signal for this return flow (Key et al. 1996).

Figure 4A further shows that AABW has higher levels of ${}^{14}C$ ($\Delta {}^{14}C = -150\%$ to -160%) than does CDW. The reason is that AABW originates principally in the Weddell Sea via mixing of newly formed WSBW with intermediate depth waters (Foster and Carmack 1976; Carmack and Foster 1977; Weiss et al. 1979; Mensch et al. 1996). The recent ventilation of the WSBW is clearly indicated by steady-state and transient tracer measurements (${}^{14}C$, ${}^{3}H$, CFC-11, CFC-12, ${}^{39}Ar$, ${}^{18}O$ and helium isotopes) in the Weddell Sea (Bayer and Schlosser 1991; Schlosser et al. 1991; Mensch et al. 1996). GEOSECS ${}^{14}C$ data provide further evidence of such formation and indicate an eastward spreading of recently ventilated bottom waters (enriched in ${}^{14}C$) from the Weddell Sea (Fig. 7).

DISCUSSION

Observed Change in Oceanic ¹⁴C

Few previously collected ¹⁴C data are available for ¹⁴C in the same region (Fig. 2). Those that are available include measurements from GEOSECS (1973–1978), ANTV in the Weddell Sea (1986) and INDIGO (1985–1987). Figure 8 provides details of the measured change of surface ocean ¹⁴C with time. Unfortunately, south of 65°S as well as between 55°S and 50°S, no adjacent GEOSECS era data exist. Between about 65°S and 55°S, surface ocean ¹⁴C decreased progressively from 1973–

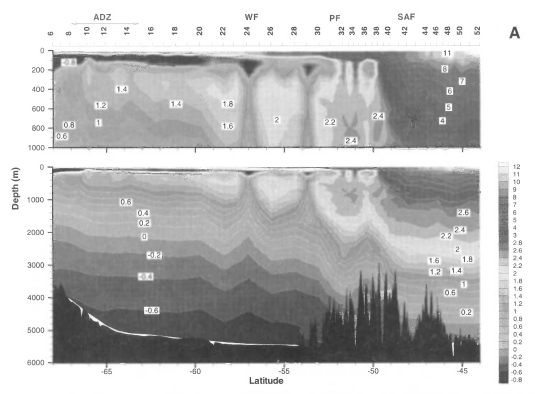


Figure 3 (A) Potential temperature θ , (B) salinity S, and (C) CFC-12 concentrations along the CIVA1 section. Also indicated are frontal zones mentioned in the text (ADZ, WF, PF and SAF).

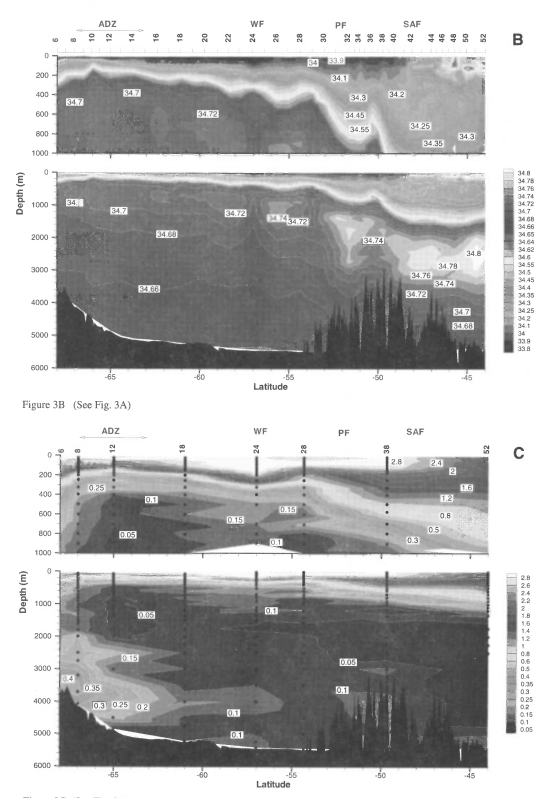


Figure 3C (See Fig. 3A)

1978 to 1993. This decreasing trend results from the decline in atmospheric ¹⁴C and the replacement of the surface waters south of the Polar Front with upwelled CDW, which remains largely uncontaminated by bomb ¹⁴C (Toggweiler and Wallace 1995). From 50°S to 44°S, there also appears to be a reduction with time, assuming a linear spatial trend between those 2 CIVA data points.

Figure 9 provides a more in-depth look at the CIVA1 minus GEOSECS temporal change. The GEO-SECS data in the Southern Ocean are sparse but were selected based on the proximity to the CIVA1 section. However, many of the chosen GEOSECS stations are far from the CIVA1 section, leading to substantial uncertainty about our calculated CIVA1-GEOSECS section. Nevertheless, natural variability must be substantially smaller than the large temporal changes in near-surface ¹⁴C that have occurred since GEOSECS. Within the upper 200 m, ¹⁴C has decreased substantially since GEO-SECS. Just below, ¹⁴C has increased. The increase extends deeper in the north than in the south. North of the CIVA1 Polar Front, the large subsurface increase in ¹⁴C since GEOSECS (up to +80‰) tags the formation of AAIW. The CDW remains largely uncontaminated by bomb ¹⁴C, a result that is consistent with the CIVA1 CFC data (Archambeau et al. 1998), which likewise reveal no significant recent contamination of the core of the CDW (Fig. 3C). Conversely, the AABW has been recently ventilated according to both the CIVA1-GEOSECS ¹⁴C difference and the CIVA1 CFC data.

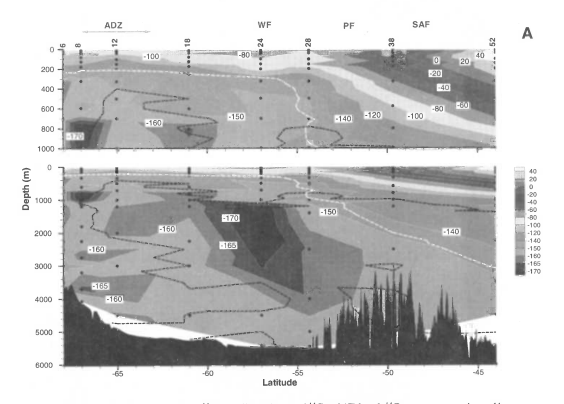


Figure 4 CIVA1 sections for (A) measured ¹⁴C as well as (B) natural ¹⁴C and (C) bomb ¹⁴C components estimated by our new CAS methodology; SAS was used for station 52 at 44°S. For SAS, the 90 μ mol kg⁻¹ isoline for dissolved silica (thick dashed line) provides the limit for penetration of bomb ¹⁴C. For CAS, that limit is provided by the 0.1 pmol kg⁻¹ isoline for CFC-12 (thin dashed line).

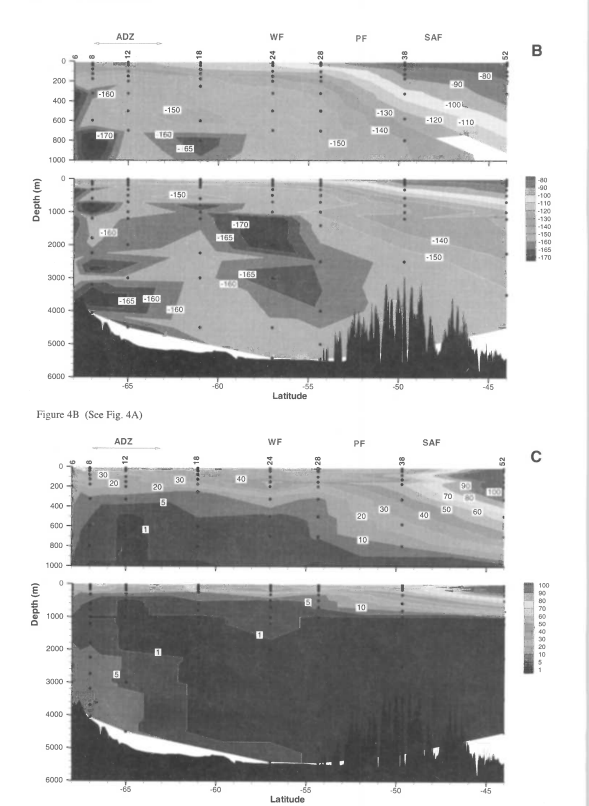


Figure 4C (See Fig. 4A)

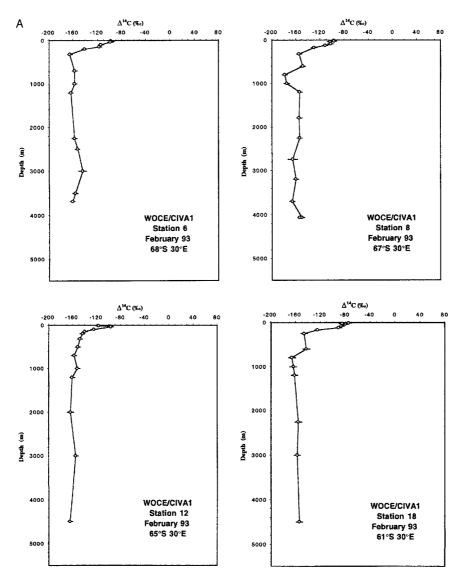


Figure 5A Profiles of measured Δ^{14} C for CIVA1 stations 6, 8, 12, and 18

Estimating Natural and Bomb ¹⁴C

Broecker et al. (1995) developed a general method to reconstruct the distribution of natural ¹⁴C in the ocean. They applied their method to the GEOSECS (1972–1978), TTO (1980–1983) and SAVE (1987–1989) data to distinguish natural from bomb ¹⁴C. Besides surface prenuclear ¹⁴C data, Broecker et al. (1995) used bomb tritium to delimit the penetration depth of bomb ¹⁴C, and the relationship between silica and natural ¹⁴C to determine the latter in the upper water column. For CIVA1, we employ the same method for our one station that fell north of 45°S (station 52 at 44°S). South of 45°S, Broecker et al. (1995) rely purely on SiO₂. In deep waters (SiO₂ > 90 µmol kg⁻¹), Broecker et al.

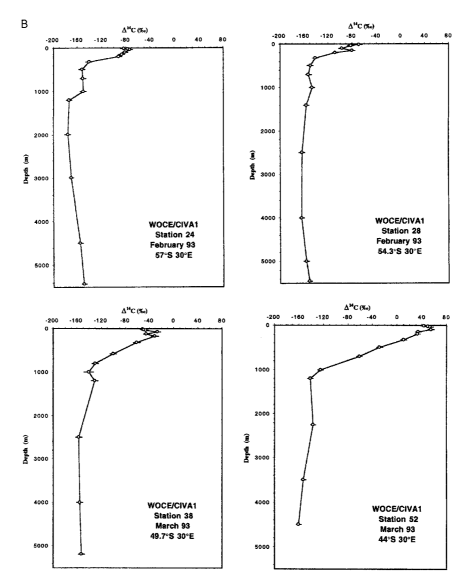


Figure 5B Profiles of measured Δ^{14} C for CIVA1 stations 24, 28, 38 and 52

(1995) consider that ¹⁴C is entirely natural in origin (i.e., that deep waters were entirely free from bomb ¹⁴C). Although this is a reasonable approximation for the time of GEOSECS, only about 10 yr after the peak in atmospheric ¹⁴C, such appears not to be the case for CIVA1 samples, collected about 20 yr later.

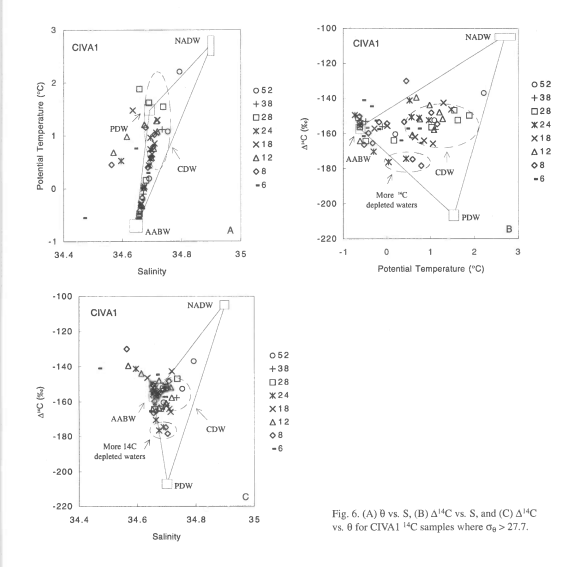
At the time of CIVA1, it is possible that bomb ¹⁴C had penetrated into deep waters of the Southern Ocean, where vertical mixing and advection are particularly active. To properly address such a possibility, we were forced to modify the method for the region south of 45°S. For this, we employ a simple ventilation model (see Appendix) that relates deep water concentrations to historical surface values. The ventilation parameters of such a model are calibrated by CFC data, measured during CIVA1,

because such gases have no natural component. Hence, this approach is termed the CFC Analogue Strategy (CAS). Then, with a data-based reconstructed history of surface ocean ¹⁴C in the same region, we computed the penetration of bomb ¹⁴C.

Silica Analogue Strategy and CFC Analogue Strategy

The Silica Analogue Strategy (SAS) from Broecker et al. (1995) improves on previous methodology (Broecker et al. 1985) to separate natural and bomb ¹⁴C in the ocean by exploiting the linear correlation between natural ¹⁴C and dissolved silica found to exist in ³H-free waters throughout most of the oceans:

$$\Delta^{14} C_{natural} = -70 - SiO_2 \tag{1}$$



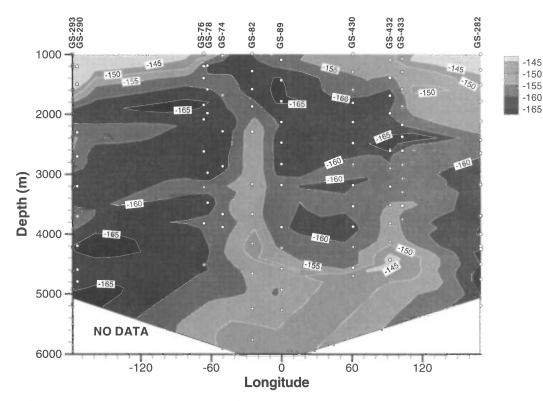


Figure 7 Longitudinal section of measured ¹⁴C in the Antarctic Circumpolar Current (ACC) from GEOSECS (1972–1978). Latitudes of stations vary between 52°S and 62°S.

where Δ^{14} C is in % and SiO₂ is in µmol kg⁻¹. Broecker et al. (1995) estimate the uncertainty of the natural ¹⁴C thus calculated as ±10‰. Figure 10 is a plot of ¹⁴C versus silica for GEOSECS, INDIGO3, and CIVA1 data from the Southern Ocean south of 50°S. Also given is the linear regression from equation (1). For waters south of 45°S, Broecker et al. (1995) apply equation (1) when SiO₂ < 90 µmol kg⁻¹. They assume that waters with SiO₂ > 90 µmol kg⁻¹ are uncontaminated with bomb ¹⁴C. However, south of 45°S, contamination of deep waters by bomb ¹⁴C is much more likely for CIVA1-era samples. Figure 10 suggests that bomb ¹⁴C has invaded deep waters since GEO-SECS: for waters with SiO₂ between 90 and 100 µmol kg⁻¹, ¹⁴C is greater during CIVA1 than during GEOSECS. Thus, for CIVA1, one cannot use the SAS method alone, which assumes that measured variability in deep water ¹⁴C is due to natural variation. Instead, south of 45°S, we modified the technique in defining 3 separate regions (Fig. 4A):

- 1. In the waters where $SiO_2 < 90 \ \mu mol \ kg^{-1}$, natural ¹⁴C is deduced from Broecker et al.'s (1995) SAS methodology.
- 2. In the core of the CDW, where Δ¹⁴C values are the lowest and have not changed since the time of GEOSECS, we consider the ¹⁴C to be entirely prenuclear in origin. Arbitrarily, we choose the weak CFC-12 concentrations of 0.1 pmol kg⁻¹ as a limit to any anthropogenic contamination. We are unable to employ the CFC-11/CFC-12 dating method near the surface, nor in deep waters. Indeed, the ratio CFC-11/CFC-12 is not usable to date young near-surface waters in CIVA1 because since the mid-1970s, growth rates of atmospheric CFC-11 and CFC-12 have

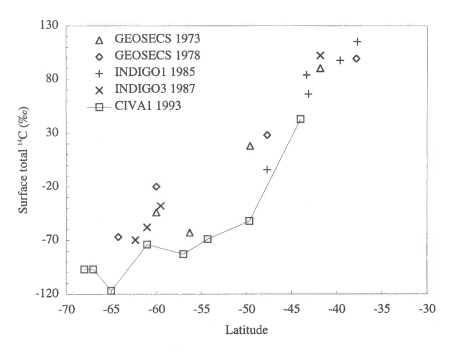


Figure 8 Measured surface ocean 14 C from GEOSECS, INDIGO and CIVA1 as a function of latitude for the region between 25°W to 85°E and 70°S to 30°S.

been the same. Furthermore, in CIVA's deep waters where the CFC concentrations are very low, uncertainty in the CFC-11/CFC-12 ratio is too large to be of any use.

3. Elsewhere (i.e., mainly in the ventilated southernmost part of the section), our vertical mixing model and CFCs from CIVA1 are used to calculate the annual ventilation rate of any water parcel, and subsequently, the bomb ¹⁴C contribution based on a reconstructed history of surface bomb ¹⁴C. By dealing directly with surface ocean concentrations, we avoid problems due to the roughly 100-fold difference in the time required for ¹⁴C versus CFCs to equilibrate between the atmosphere and surface ocean (Broecker and Peng 1974).

Natural Versus Bomb 14C

Figure 4B shows the computed natural ¹⁴C. Surface values of natural ¹⁴C increase from south to north from -132% to -72%. Our lowest surface values in the far south are consistent with the rare existing data set (Broecker et al. 1985; Bard 1988; Toggweiler and Samuels 1993; Berkman and Forman 1996), especially considering the uncertainties from analysis, the Suess effect (5‰ to 12‰ [Berkman and Forman 1996]), the silica-natural ¹⁴C relationship (±10‰), and natural spatial variability. Waters north of the PF, where AAIW forms, have higher natural ¹⁴C, largely because they rest longer at the surface and thus equilibrate more extensively with the atmosphere; also circulation patterns transport natural ¹⁴C to the north as described previously. South of the PF, a large portion of the intermediate and deep waters have natural ¹⁴C values lower than -160%, which is the average value of the waters of the Antarctic Circumpolar Current (Broecker et al. 1995).

The bomb ¹⁴C results (Fig. 4C) are consistent with the observed trends of the total Δ^{14} C values: waters are poor in bomb ¹⁴C south of the PF and richer in the north; intermediate, deep, and bottom

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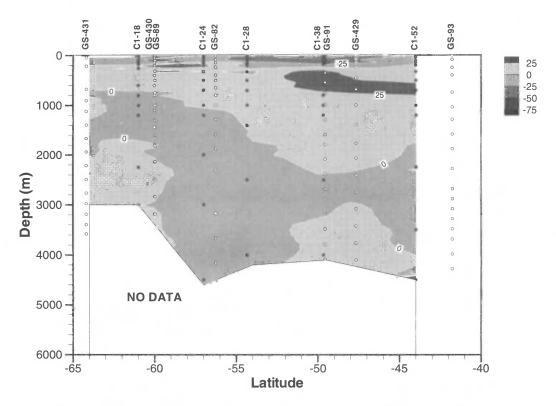


Figure 9 Section of the measured change in ¹⁴C between GEOSECS and CIVA1. GEOSECS stations are taken from longitudes between 25°W and 85°E. GEOSECS and CIVA1 station data were triangulated then interpolated to the same 1-degree latitudinal grid before taking differences.

waters contain little if any bomb ¹⁴C. The only exception is where substantial bomb ¹⁴C is detected between 2000 and 5000 m in the far south along the continental slope. Deep and bottom waters in the northern portion of the section contain insignificant levels.

Bomb ¹⁴C Inventories

Following Broecker et al. (1985), we calculated the vertical integral of bomb ¹⁴C (also known as the inventory) as the area between bomb and natural ¹⁴C profiles (Table 2, Fig. 11). With our method, inventories range from 1.7 to 10.9×10^9 atoms cm⁻². Inventories are larger north of the PF (stations 38 and 52) as well as in the far south, at stations 6 and 8. North of 45°S (station 52), the bomb ¹⁴C inventory is highest ($10.9 \pm 1.6 \times 10^9$ atoms cm⁻²) and its uncertainty is lowest: both the SAS and CAS methods give the same inventory. We were able to properly apply the SAS method at station 52, because our measured CIVA1 ³H (not shown) was well above background levels. Figure 11 compares CIVA1 bomb ¹⁴C inventories to those from GEOSECS and INDIGO. The temporal change is similar in structure to that seen for surface concentrations (Fig. 8). Around 60°S, the differences between inventories during CIVA1 (CAS) and GEOSECS (SAS) tend towards zero: the ¹⁴C loss near the surface is compensated by buildup at depth (Fig. 4C). Around 60°S, when the SAS methodology is used, CIVA1 inventories are only 20% to 50% of those determined using CAS. Further south, stations at 67°S and 68°S exhibit the largest differences between the SAS and CAS methods.

Uncertainties concerning the Southern Ocean's bomb ¹⁴C inventories are large but their magnitude is relatively small compared to the inventories throughout most of the rest of the ocean (Broecker et

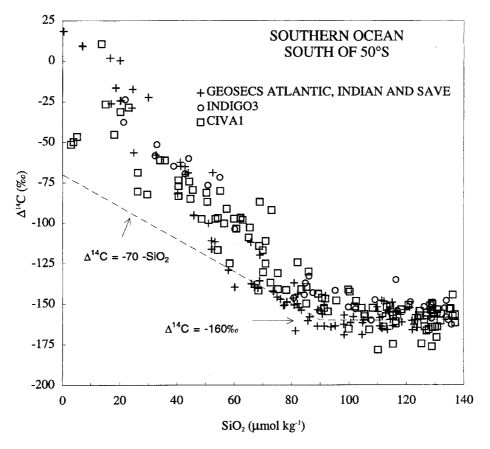


Figure 10⁻¹⁴C versus dissolved SiO₂ for stations taken south of 50°S from the GEOSECS, INDIGO3, and CIVA1 expeditions. Also provided is the linear correlation between the natural Δ^{14} C and silica for tritium-free waters (Broecker et al. 1995).

al. 1995). Thus, uncertainties in this region do not substantially alter estimates of the global ocean inventory of bomb ¹⁴C. Regional patterns of bomb ¹⁴C inventories are generally useful to help evaluate the behavior of OGCMs (Toggweiler et al. 1989b). However, they are less useful in the Southern Ocean, due to the large uncertainties there. On the other hand, levels of bomb ¹⁴C near the surface in this same region have much smaller uncertainties than do corresponding horizontal and vertical gradients, and are thus more useful for model validation.

CONCLUSION

To complement the sparse ¹⁴C data set in the Southern Ocean, AMS ¹⁴C measurements were made on recently collected samples taken along the CIVA1 cruise track. The measured ¹⁴C distribution reveals a dramatic contrast between samples taken south and north of the PF. Surface waters in the south are depleted due to upwelling of old CDW at the ADZ, which drives transport to the north; northern surface waters are rich in ¹⁴C due to transport from south to north and their more extensive equilibration with the atmospheric ¹⁴C. The oldest deep waters are found in the CDW (< -170%) and derive from the PDW.

Station	Surface total ¹⁴ C (%o)	Bomb ¹⁴ C inventories ^a	10 ⁹ atoms per cm ^{2b}		
6	-97	1	4.9		
8	-97	1.2	5.1		
12	-117	0.8	2.4		
18	-74	1.1	1.7		
24	-83	1.8	3.1		
28	-69	1.5	1.7		
38	-52	4.7	4.7		
52	43	10.9			

Table 2 CIVA1 bomb ¹⁴C inventories

^aFollowing the Broecker et al. (1995) method

^bFollowing the chlorofluorocarbon analogue strategy

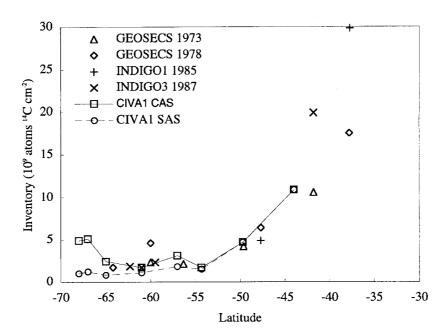


Figure 11 Bomb ^{14}C inventory versus latitude for GEOSECS, INDIGO3 and CIVA1 stations within 25°W–85°E and 70°S–30°S.

Bomb ¹⁴C has contaminated deep waters in the far south based on differences in measured ¹⁴C between CIVA1 and GEOSECS and the CIVA1 distribution of CFC-11 and CFC-12. Our new CAS method relies on measured CFCs, a simple mixing model, and the established history of surface ocean ¹⁴C. A previously developed method, SAS from Broecker et al. (1995), relies on the assumption that all deep waters of the Southern Ocean are free of bomb ¹⁴C. For CIVA1 as well as other WOCE-era samples, such an assumption is invalid. In terms of bomb ¹⁴C inventories, our new CAS methodology yields estimates identical to the SAS method for stations north of the PF, but substantially greater than SAS results when south of the PF (up to 5 times more when south of the ADZ). Although CAS offers an improvement for determining inventories south of PF, uncertainties remain high and are not ideal for validating ocean carbon-cycle models. CAS estimates in deep waters south of the PF would be improved if the history of surface ocean ¹⁴C there were better constrained. South

of the PF, we recommend that ocean modelers not concern themselves with inventories but rather with bomb and natural ¹⁴C distributions (Fig. 4B,C), weighing errors accordingly. For CIVA1 north of the PF, bomb ¹⁴C is found in surface and intermediate layers; in the far south, bomb ¹⁴C is found throughout the water column. Since GEOSECS, ¹⁴C has decreased in surface waters (roughly the upper 200 m) and has increased below, to as deep as 2000 m north of the PF.

The CIVA1 section began near Antarctica and was aborted prematurely at 44°S due to an emergency medical evacuation. Fortunately, the repeat section CIVA2 (February–March 1996) will allow the northern gap (where ¹⁴C inventories should be greatest) to be filled in, once ¹⁴C measurements are completed. The CIVA section along with other new WOCE ¹⁴C data will greatly improve coverage in the Southern Ocean, a key area for ocean CO₂ uptake. However, care will have to be exercised when estimating natural and bomb ¹⁴C components so that ocean modelers will be able to take full advantage of this unique data set.

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REFERENCES

- Andrié C, Jean-Baptiste P, Merlivat L. 1986. Tritium and Helium-3 in the northeastern Atlantic Ocean during the 1983 topogulf cruise. *Journal of Geophysical Re*search 93:12511–24.
- Archambeau A-S, Pierre C, Poisson A, Schauer B. 1998. Distributions of oxygen and carbon stable isotopes and CFC-12 in the water masses of the Southern Ocean at 30°E from South Africa to Antarctica: results of the CIVA1 cruise. Journal of Marine Systems 17:25–38.
- Arnold M, Bard E, Maurice P, Duplessy J-C. 1987. ¹⁴C dating with the Gif-Sur-Yvette Tandetron Accelerator: status report. *Nuclear Instruments and Methods in Physics Research* B29:120–3.
- Bard E. 1988. Correction of AMS ¹⁴C ages measured in planktonic foraminifera: paleoceanographic implications. *Paleoceanography* 3:635–45.
- Bard E, Arnold M, Maurice P, Duplessy J-C 1987 Measurements of bomb radiocarbon in the ocean by means of accelerator mass spectrometry: technical aspects. *Nuclear Instruments and Methods in Physics Re*search B29:297–301.
- Bard E, Arnold M, Östlund HG, Maurice P, Monfray P, Duplessy J-C. 1988. Penetration of bomb radiocarbon in the tropical Indian Ocean measured by means of accelerator mass spectrometry. *Earth and Planetary Science Letters* 87:379–89.
- Bayer R, Schlosser P. 1991. Tritium profiles in the Weddell Sea. *Marine Chemistry* 35:123–36.

- Berkman PA, Forman SL. 1996. Pre-bomb radiocarbon and the reservoir correction for calcareous marine species in the Southern Ocean. *Geophysical Research Letters* 23:363–6.
- Broecker WS, Peng TH. 1974. Gas exchange rates between air and sea. *Tellus* 26:21–35.
- Broecker WS, Peng T-H, Östlund G, Stuiver M. 1985. The distribution of bomb radiocarbon in the ocean. *Journal of Geophysical Research* 90:6953–70.
- Broecker WS, Sutherland S, Smethie W, Peng TH, Östlund G. 1995. Oceanic radiocarbon: separation of the natural and bomb components. *Global Bio*geochemical Cycles 9:263–88.
- Carmack EC, Foster TD. 1977. Water masses and circulation in the Weddell Sea. In: Dunbar MJ, editor. *Polar* oceans. Calgary: Arctic Institute of North America. p 151–65.
- Foster TD, Carmack EC. 1976. Frontal zone mixing and Antarctic bottom water formation in the southern Weddell Sea. *Deep-Sea Research* 23:301–17.
- Gordon AL. 1971. Antarctic polar front zone. In: Reid JL, editor. Antarctic Oceanology I, Antarctic research series. Washington: American Geophysical Union. p 205–21.
- Haine TWN. 1996. Combining passive tracer observations with ocean circulation models. *International WOCE Newsletter* 23:3–5.
- Jean-Baptiste P, Mantisi F, Mémery L, Jamous D. 1991.

³He and chlorofluorocarbons (CFC) in the Southern Ocean: tracers of water masses. *Marine Chemistry* 35:137–50.

- Jenkins WJ. 1980. Tritium and ³He in the Sargasso Sea. *Journal of Marine Research* 38: 533–69.
- Key RM, Quay PD, Jones GA, McNichol AP, Von Reden KF, Schneider RJ. 1996. WOCE AMS Radiocarbon I: Pacific Ocean results (P6, P16, P17). *Radiocarbon* 38(3):425–518.
- Maier-Reimer E. 1993. Geochemical cycles in an ocean general circulation model preindustrial tracer distributions. *Global Biogeochemical Cycles* 7:645–77.
- Mensch M, Bayer R, Bullister JL, Schlosser P, Weiss RF. 1996. The distribution of tritium and CFCs in the Weddell Sea during the mid-1980s. *Progress in Oceanography* 38:377–415.
- Orr JC. 1996. The ocean carbon-cycle model intercomparison project of IGBP/GAIM. In: Ormerod B, editor. International Energy Agency, ocean storage of CO₂ workshop 3: international links and concerns (ISBN 1 89 83 73 04 3). Southampton, England.
- Orsi AH, Nowlin WD, Whitworth T III. 1993. On the circulation and stratification on the Weddell Gyre. *Deep-Sea Research* 40:169–203.
- Orsi AH, Whitworth T III, Nowlin WD. 1995. On the meridional extent and fronts of the Antarctic Circumpolar Current. *Deep-Sea Research* 42:641–73.
- Östlund HG, Grall C. 1988. *INDIGO 1985–1987 Indian Ocean radiocarbon: tritium laboratory data report*. University of Miami: Rosenstiel School of Marine and Atmospheric Science.
- Östlund HG, Stuiver M. 1980. GEOSECS Pacific radiocarbon. Radiocarbon 22(1):25–53.
- Poisson A, Chen C-TA. 1987. Why is there little anthropogenic CO₂ in the Antarctic bottom water? *Deep-Sea Research* 34:1255–75.
- Sarmiento JL, Orr JC, Siegenthaler U. 1992. A perturbation simulation of CO₂ uptake in an ocean general circulation model. *Journal of Geophysical Research* 97: 3621–45.
- Schlosser P, Bullister JL, Bayer R. 1991. Studies of deep water formation and circulation in the Weddell Sea using natural and anthropogenic tracers. *Marine Chemistry* 35:97–122.
- Schlosser P, Kromer B, Weppernig R, Loosli HH, Bayer R, Bonani G, Suter M. 1994. The distribution of ¹⁴C and ³⁹Ar in the Weddell Sea. *Journal of Geophysical Research* 99:10275–87.

- Stuiver M, Östlund HG. 1980. GEOSECS Atlantic radiocarbon. Radiocarbon 22:1–24.
- Stuiver M, Östlund HG. 1983. GEOSECS Indian Ocean and Mediterranean radiocarbon. *Radiocarbon* 25(1): 1–29.
- Stuiver M, Östlund HG, McConnauqhey TA. 1981. GE-OSECS Atlantic and Pacific ¹⁴C distribution. In: Bolin B, editor. *Carbon cycle modeling*. New York: John Wiley & Sons. p. 201–9.
- Stuiver M, Polach HA. 1977. Reporting of ¹⁴C data. *Radiocarbon* 19(3):355–63.
- Taylor NK. 1995. Seasonal uptake of anthropogenic CO₂ in a ocean general circulation model. *Tellus* 47B:145–69.
- Toggweiler JR, Dixon K, Bryan K. 1989a. Simulations of radiocarbon in a coarse-resolution world ocean model. l. Steady state prebomb distributions. *Journal of Geophysical Research* 94:8217–42.
- Toggweiler JR, Dixon K, Bryan K. 1989b. Simulations of radiocarbon in a coarse-resolution world ocean model. 2. Distributions of bomb-produced carbon 14. *Journal* of Geophysical Research 94:8243–64.
- Toggweiler JR, Samuels B. 1993. New radiocarbon constraints on the upwelling of abyssal water to the ocean's surface. In: Heimann, M, editor. *The global carbon cycle*. Berlin: NATO ASI Series. p 333–66.
- Toggweiler JR, Wallace D. 1995. Transport capacity for passive tracers. *US WOCE Report 1995*. p 36–8.
- Warner MJ, Weiss RF. 1985. Solubilities of chlorofluorocarbons 11 and 12 in water and seawater. *Deep-Sea Research* 32:1485–97.
- Weiss RF, Östlund HG, Craig H. 1979. Geochemical studies of the Weddell Sea. *Deep-Sea Research* 26A: 1093–120.
- Weppernig R, Schlosser P, Khatiwala S, Fairbanks RG. 1996. Isotope data from Ice Station Weddell: implications for deep water formation in the Weddell Sea. *Journal of Geophysical Research* 101:25723–39.
- Whitworth T III, Nowlin WD. 1987. Water masses and currents of the southern ocean at the Greenwich Meridian. *Journal of Geophysical Research* 92:6462–76.
- Worthington LV. 1977. The case for near-zero production of Antarctic bottom water. *Geochimica Cosmochima Acta* 41:1001–6.
- Wunsch C, Hu DX, Grant B. 1983. Mass, heat, salt, and nutrients fluxes in the South Pacific Ocean. *Journal of Physical Oceanography* 13: 725–753.

APPENDIX: VENTILATION MODEL

Our ventilation model is inspired by Jenkins (1980) who used such a model to explain the penetration of ³H in the North Atlantic and to study the ³H/³He relationship. At any time t, our model determines the annual ventilation rate η , which is defined as the fraction of water at depth z replaced by water originating from the surface during the timestep Δt . A time-lag τ is defined to correspond to the travel time of a water parcel moving from the surface to depth z. Thus, at depth z

$$C_{z,t+1} = C_{z,t} + \eta \times [C_{0,(t-\tau)} - C_{z,t}]$$
(A1)

where $C_{0,(t-\tau)}$ is the surface concentration at time (t- τ) and $C_{z,t}$, $C_{z,t+1}$ are the concentrations at depth z in years t and t+1.

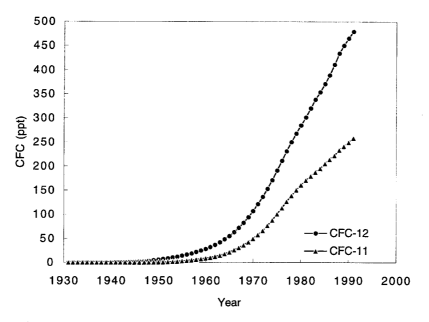


Figure A1 Partial pressure of atmospheric CFC-11 and CFC-12 in the Southern Hemisphere versus time (Walker, Salameh, and Weiss, personal communication 1995).

First, the annual ventilation rate and the transit time are adjusted using both CFC-11 and CFC-12 to match the measured concentrations at the time of CIVA1 (1993). For our calculations, we used the time evolution of atmospheric CFC-11 and CFC-12 concentrations (Fig. A1) given by Walker, Salameh, and Weiss (personal communication 1995) as well as solubilities from Warner and Weiss (1985). This simple method gives the best result when a transit time τ of < 1 yr is chosen. Results are essentially identical whether using CFC-11 or CFC-12. With our method ($\Delta t = 1$ yr), this means that τ is set to zero.

Second, we used the same model, now with the known η and τ from the CFC calibration, and a reconstructed ¹⁴C input function shown in Figure A2, to determine the evolution of the total ¹⁴C until the time of CIVA1 (see an example of the method in Fig. A3). For this, we again used equation (A1), where C is now ¹⁴C. The reconstructed ¹⁴C input function has large uncertainties due to the scatter of available data (Fig. A3). We have arbitrarily drawn minimum and maximum curves. We favor the lowest curve because it allows a close agreement around the transition at 90 µmol kg⁻¹

between the CAS and SAS methods; SAS is applied for waters where SiO_2 lies between 0 and 90 μ mol kg⁻¹.

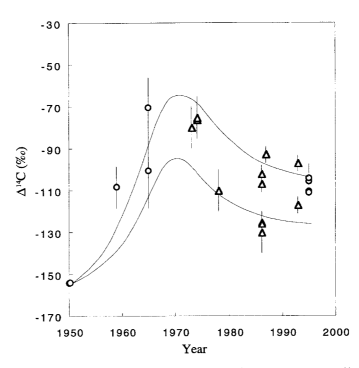


Figure A2 Time evolution of the surface ocean Δ^{14} C compiled from the 14 C record from marine molluscs, brachiopods, echinoderms and foraminifera (O) collected near the Antarctica (Berkman and Forman 1996), and 14 C measurements on surface samples from GEOSECS, INDIGO3 and CIVA1 stations (Δ) near Antarctica (Östlund and Stuiver 1980; Stuiver and Östlund 1980, 1983; Stuiver et al. 1981; Schlosser et al. 1994; Östlund and Grall 1988). Predicted prenuclear surface Δ^{14} C is ca. -150% (Berkman and Forman 1996). Minima and maxima curves are drawn arbitrarily. We chose the lower curve for our CAS methodology because with it CAS and SAS methods agree for waters where SiO₂ <90 µmol kg⁻¹.

