

Natural and anthropogenic radionuclide distributions in the Nansen Basin, Artic Ocean: Scavenging rates and circulation timescales

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Abstract—Determination of the naturally occurring radionuclides 232 Th, 230 Th, 228 Th and 210 Pb, and the anthropogenic radionuclides 241 Am, 239,240 Pu, 134 Cs and 137 Cs in water samples collected across the Nansen Basin from the Barents Sea slope to the Gakkel Ridge provides tracers with which to characterize both scavenging rates and circulation timescales in this portion of the Arctic Ocean. Large volume water samples (~ 1500 l) were filtered in situ to separate particulate ($> 0.5 \mu$ m) and dissolved Th isotopes and 241 Am. Thorium-230 displays increases in both particulate and dissolved activities with depth, with dissolved 230 Th greater and particulate 230 Th lower in the deep central Nansen Basin than at the Barents Sea slope. Dissolved 228 Th activities also are greater relative to 228 Ra, in the central basin. Residence times for Th relative to removal from solution onto particles are ~ 1 year in surface water, ~ 10 years in deep water adjacent to the Barents Sea slope, and ~ 20 years in the Eurasian Basin Deep Water. Lead-210 in the central basin deep water also has a residence time of ~ 20 years with respect to its removal from the water column. This texture of scavenging is reflected in distributions of the particle-reactive anthropogenic radionuclide 241 Am, which shows higher activities relative to Pu in the central Nansen Basin than at the Barents Sea slope.

Distributions of ¹³⁷Cs show more rapid mixing at the basin margins (Barents Sea slope in the south, Gakkel Ridge in the north) than in the basin interior. Cesium-137 is mixed throughout the water column adjacent to the Barents Sea slope and is present in low but detectable activities in the Eurasian Basin Deep Water in the central basin. At the time of sampling (1987) the surface water at all stations had been labeled with ¹³⁴Cs released in the 1986 accident at the Chernobyl nuclear power station. In the ~1 year since the introduction of Chernobyl ¹³⁴Cs to the Nansen Basin, it had been mixed to depths of ~800 m at the Barents Sea Slope and to ~300 m in the central basin. "Pre-Chernobyl" inventories of ¹³⁷Cs (as well as ^{239,240}Pu) are 10 times those expected from global atmospheric fallout from nuclear weapons testing and are derived principally from releases from the Sellafield, U.K., nuclear fuel reprocessing facility on the Irish Sea. Based on the sources of ¹³⁷Cs to the Nansen Basin, mixing time scales are 9–18 years for the upper water column (to 1500 m) and ~40 years for the deep water. These mixing time scales, combined with more rapid scavenging at the basin margin relative to the central basin, produce residence times of particle-reactive radionuclides in the Nansen Basin comparable to other open ocean areas (e.g. north-west Atlantic) despite the presence of permanent ice cover and long periods of low-light levels that limit productivity in the Arctic.

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INTRODUCTION

The suite of naturally-occurring and anthropogenically-produced radionuclides contains tracers suitable for studying transport of both particles and water in the oceans. Several of the radionuclides of the uranium and thorium decay series, in particular the Th isotopes ²³⁰Th (half-life = 75,000 years) and ²²⁸Th (half-life = 1.9 years) and the Pb isotope ²¹⁰Pb (half-life = 22 years), are produced in seawater from decay of dissolved precursors (²³⁴U, ²²⁸Ra and ²²⁶Ra, respectively), become associated with particles, and are removed from the water column.

Deliberate and accidental releases of anthropogenic radionuclides to the oceans provide transient tracers that are also useful for studying scavenging and circulation processes (Kupferman et al., 1979; Bowen et al., 1980; Livingston et al., 1982, 1984, 1985). The Pu isotopes and ²⁴¹Am can associate with particle surfaces, and their distributions in the oceans are affected by scavenging and particle transport. Americium-241 in particular has a chemical behavior comparable to Th in its particle reactivity (Fisher et al., 1988; Cochran et al., 1987). Cesium-137, on the other hand, exhibits only weak particle reactivity in the open ocean, and its distribution is largely controlled by advection and mixing of water. Dumping of spent nuclear fuel, reactor parts and radioactive waste in the Kara Sea by the former Soviet Union has led to recent interest in the chemical behavior and fate of anthropogenic radionuclides in the Arctic Ocean. In this context, the natural U and Th series radionuclides can be used to provide clues to the processes controlling the distributions of their anthropogenic counterparts because the source functions of the natural nuclides are well characterized and their distributions tend toward a steady state.

Within the last decade, significant insights into the texture of scavenging of chemical species in the oceans have been gained from distributions of particle-reactive radionuclides (Nozaki et al., 1981; Bacon and Anderson, 1982; Mangini and Key, 1983; Bacon et al., 1985; Bruland and Coale, 1986; Cochran et al., 1987; Coale and Bruland, 1987; Huh and Beasley, 1987). Higher rates of particle production and flux in the surface ocean and at the ocean margins make these areas generally more effective scavenging regimes. Indeed transport of reactive radionuclides such as ²¹⁰Pb and ²³¹Pa to ocean margins followed by intense scavenging in these areas has been used to explain the distributions and gradients of these radionuclides away from the margins to the basin interior (Carpenter et al., 1981; Anderson et al., 1983a,b; Cochran et al., 1983, 1990; Bacon, 1988).

The Arctic Ocean has been regarded as an environment of low scavenging intensity relative to other open ocean areas because the permanent ice cover and extended periods of darkness during the Arctic winter serve to inhibit particle production and sinking. This possibility was first raised by Ku and Broecker (1967) and Finkel et al. (1977) to explain low activities and inventories of ²³⁰Th and ¹⁰Be, respectively, in Arctic Ocean sediments. More recently, Moore and Smith (1986) and Bacon et al. (1989) have shown that scavenging of ²¹⁰Pb and Th isotopes is slower in the waters of the Canada Basin relative to the open oligotrophic ocean. We present here distributions of natural and anthropogenic radionuclides in the Nansen Basin and show that removal of particle-reactive radionuclides is more rapid in this basin than in the Canada Basin. The difference is likely due to more rapid ventilation of the Nansen Basin coupled with enhanced scavenging in the slope and broad marginal Arctic shelf.

Sampling and analytical methods

Water and sediment samples were collected during the 1987 ARK IV/3 cruise of the R.V. Polarstern (Fig. 1). Water samples for Th isotopes and ²⁴¹Am were taken by in-situ pumps which filter large volumes of water through 0.5 µm polypropylene wound fiber cartridge prefilters and two similar cartridges loaded with manganese oxide. The former retain the fine particulate fraction, while the latter adsorb dissolved thorium (Cochran et al., 1987; Livingston and Cochran, 1987; Buesseler et al., 1992). Two modifications to previous sampling procedures with the pumps (Cochran et al., 1987) were made on the ARK IV/3 cruise. The manganese adsorber cartridges were not prepared in the laboratory before the cruise by soaking in hot KMnO₄ solution, but instead were loaded with freshly precipitated manganese oxide before each deployment. The MnO₂ was formed from a solution of MnCO₃ and KMnO₄ at pH = 8 and was filtered onto the cartridge. In addition, a few of the pump samples were taken with two prefilter cartridges in series, followed by the two manganese adsorber cartridges. The purpose of this modification was to check for possible artifacts caused by sorption of dissolved Th onto the prefilter cartridge.

The cartridges were returned to the laboratory where they were dried and ashed at 450°C. The ash was dissolved in HNO₃ in the presence of ²²⁹Th and ²⁴³Am yield tracers. Thorium

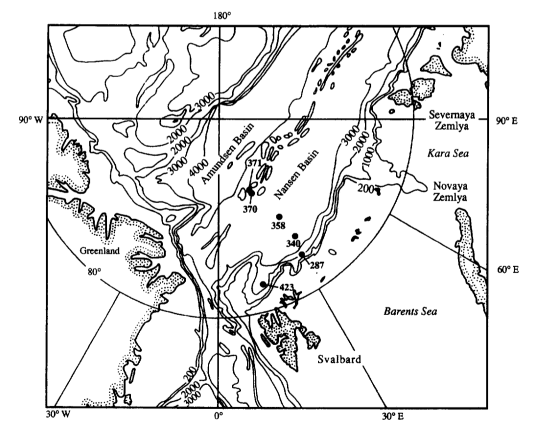


Fig. 1. Map of the Nansen Basin showing stations sampling for radionuclides during cruise ARK IV/3 of the R.V. Polarstern.

was separated from Am by ion exchange. The Am and Th fractions were further purified by ion exchange and subsequently electrodeposited for alpha spectrometry (Cochran *et al.*, 1987). Prefilters and cartridges deployed in pumps that failed to operate were used as blanks.

Water samples also were taken by hydrocast (Gerard barrels) for analysis of ^{239,240}Pu, ^{134,137}Cs and ⁹⁰Sr (Livingston *et al.*, 1975). These samples were acidified to pH 2, stored in 60 l delex containers and returned to the laboratory for processing. Aliqots (4 l) from two stations were taken from the 60 l samples for ²¹⁰Pb analysis. Polonium-209 tracer and Fe carrier were added to these samples and iron hydroxide was precipitated. Lead-210 was determined by plating and counting its ²¹⁰Po daughter (Flynn, 1968). Because the samples were processed and analyzed 1.5 years after collection, ²¹⁰Po should be within about 5% of equilibrium with ²¹⁰Pb.

Samples for Ra isotopes were collected from large volume (300 l) samples taken for radiocarbon. After the ¹⁴C had been converted to CO₂ by acidification and stripped from the samples, the pH was adjusted to 7 and radium was extracted by passing the sample through acrylic fiber coated with MnO₂ (Moore, 1976; Moore *et al.*, 1985). The ²²⁸Ra/²²⁶Ra activity ratio was measured on these samples. Because this procedure does not result in quantitative removal of Ra from solution, small volume (20 l) samples are generally taken for independent determination of ²²⁶Ra. This was not done on the ARK IV/3 cruise, and as a consequence only ²²⁸Ra/²²⁶Ra ratio data are available for those samples. The ²²⁸Ra/²²⁶Ra data are reported in detail by Rutgers van der Loeff *et al.* (1995).

RESULTS

The thorium isotope data are presented in Table 1. Extraction efficiencies for each Mn cartridge pair were calculated from

$$Efficiency = 1 - MnB/MnA$$
 (1)

where MnB is the activity on the second cartridge and MnA is the activity on the first cartridge.

Efficiencies generally ranged between 65 and 100% (based on 230 Th activities) with a mean ± 1 s.d. of 740 \pm 17.0%. A few samples had values less than 50%, and the uncertainties in the resultant dissolved activities were large. These data have not been included in Table 1. The mean extraction efficiency with the cartridges loaded with preformed MnO₂ is lower than that for cartridges impregnated with MnO₂ by soaking in hot KMnO₄ (Cochran *et al.*, 1987; Buesseler *et al.*, 1992), and we conclude that the latter is a better procedure.

The blank contribution to sample peaks was generally less than 40% for ²³²Th, 25% for ²³⁰Th and 5% for ²²⁸Th. Uncertainties reported in Table 1 were calculated from one sigma counting uncertainties propagated in the blank, sample activities and cartridge extraction efficiencies.

Some ²²⁸Ra is extracted from seawater onto the Mn cartridges, and then dissolved ²²⁸Th data must be corrected for ingrowth from ²²⁸Ra between sample collection and initial separation of Th and Ra. As the first step in making this correction, we calculated ²²⁸Ra activities from the measured ²²⁸Ra/²²⁶Ra activity ratios and ²²⁶Ra activities calculated from dissolved Si data. The estimation of ²²⁶Ra used the ²²⁶Ra–Si relationship obtained from TTO Stas 142–162 (Key et al., 1992). In effect these stations constitute the source water for

Table 1. Nansen Basin thorium isotope data

(m) Sta. 287 (81°40°N, 30°48°E, 2250 m) 6	230Th 0.0125 ± 0.0089 0.0130 ± 0.0110 0.0298 ± 0.0123 0.0670 ± 0.0100 0.102 ± 0.011	228Th	232TL			(dmm/1000 l)
727	0.0125 ± 0.0089 0.0130 ± 0.0110 0.0298 ± 0.0123 0.0670 ± 0.0100 0.102 ± 0.011	=	u 1	²³⁰ Th	²²⁸ Th	(r poor /mulin)
0 0 727	0.0125 ± 0.0089 0.0130 ± 0.0110 0.0298 ± 0.0123 0.0670 ± 0.0100 0.102 ± 0.011					
0 0 257	0.0130 ± 0.0110 0.0298 ± 0.0123 0.0670 ± 0.0100 0.102 ± 0.011	0.357 ± 0.041	0.0382 ± 0.0062	0.0919 ± 0.0172	7.26 ± 3.26	20.9 ± 1.1
0 0 27	0.0298 ± 0.0123 0.0670 ± 0.0100 0.102 ± 0.011	0.636 ± 0.071	0.0402 ± 0.0080	0.0626 ± 0.0141	7.28 ± 1.17	27.6 ± 1.6
0 r	$\begin{array}{c} 0.0670 \pm 0.0100 \\ 0.102 \pm 0.011 \end{array}$	0.917 ± 0.066	0.0156 ± 0.0074	0.0924 ± 0.0196	6.82 ± 0.69	18.9 ± 1.1
0 1	0.102 ± 0.011	0.841 ± 0.053	0.0549 ± 0.0102	0.129 ± 0.023	5.04 ± 0.73	19.4 ± 1.2
0 10		0.623 ± 0.036	0.0213 ± 0.0039	0.227 ± 0.021	2.80 ± 0.21	7.1 ± 0.5
227	0.238 ± 0.019	0.742 ± 0.038	0.0315 ± 0.0138	0.290 ± 0.048	2.92 ± 0.45	6.1 ± 0.6
0 10	0.254 ± 0.017	0.660 ± 0.026	0.0226 ± 0.0042	0.270 ± 0.026	1.77 ± 0.15	5.1 ± 0.7
0 0 1						
10 r	0.0125 ± 0.0079	1.19 ± 0.06	0.0132 ± 0.0044	0.0584 ± 0.0143	7.18 ± 1.24	26.2 ± 1.6
10 1	0.0258 ± 0.0122	0.828 ± 0.055	0.0194 ± 0.0039	0.144 ± 0.020	7.73 ± 0.94	10.6 ± 0.8
10 r	0.074 ± 0.0149	0.631 ± 0.045	0.0254 ± 0.0072	0.239 ± 0.047	3.92 ± 0.72	7.3 ± 0.7
10 r	0.0739 ± 0.0114	0.249 ± 0.020	0.0122 ± 0.0025	0.449 ± 0.021	1.66 ± 0.23	2.4 ± 0.4
10 r	0.109 ± 0.013	0.102 ± 0.012	0.017 ± 0.0046	0.522 ± 0.042	+1	3.2 ± 0.5
10 r	0.137 ± 0.017	0.307 ± 0.026	0.0215 ± 0.0065	0.583 ± 0.066	4.56 ± 0.49	6.1 ± 0.6
227	0.142 ± 0.019	0.464 ± 0.034	0.0112 ± 0.0080	0.510 ± 0.055	3.97 ± 0.39	7.2 ± 0.5
777	(E					
224	0.0204 ± 0.0084	1.61 ± 0.07	0.0298 ± 0.0047	0.097 ± 0.015	1	j
227	0.0096 ± 0.0096	1.22 ± 0.02	0.0189 ± 0.0038	0.136 ± 0.024	l	
227	0.0227 ± 0.0102	1.08 ± 0.02	0.0294 ± 0.0080	0.210 ± 0.034	I	
227	0.0545 ± 0.0076	1.34 ± 0.02	0.0123 ± 0.0055	0.355 ± 0.052	1	
222	0.0528 ± 0.0091	0.355 ± 0.012	0.0122 ± 0.0052	0.420 ± 0.055	l	
777	0.0725 ± 0.0106	0.325 ± 0.022	0.0100 ± 0.0035	0.489 ± 0.034	1	
224	0.233 ± 0.018	1.32 ± 0.06	0.0394 ± 0.0103	0.367 ± 0.046	1	
224	0.240 ± 0.019	1.38 ± 0.06	0.0334 ± 0.0072	0.229 ± 0.034	1	-
	2245 m)					
0.0179 ± 0.0038	0.175 ± 0.017	0.934 ± 0.037	0.0678 ± 0.0156	0.0895 ± 0.0347	-	1
0.033 ± 0.0052	0.0207 ± 0.0094	1.06 ± 0.04	0.0364 ± 0.0117	0.116 ± 0.035	1	I
70000 1 0000	0.064 ± 0.0090	0.943 ± 0.048	0.0178 ± 0.0047	0.147 ± 0.027	1	l
	0.0886 ± 0.0102	0.459 ± 0.025	0.0170 ± 0.0041	0.248 ± 0.029	l	J
	0.138 ± 0.015	0.557 ± 0.032	0.0377 ± 0.0052	0.310 ± 0.025	1	1
	0.232 ± 0.021	0.768 ± 0.040	0.0191 ± 0.0052	0.309 ± 0.034	1	l

Not detected.

	(Sta. 423)	
Depth (m)	²³² Th	²³⁰ Th
PFB/PFA (dpm	/dpm)	
6	$0.044 \pm .032$	$0.062 \pm .078$
1400	1.63 ± 0.41	$0.491 \pm .166$
1650	$0.387 \pm .140$	$0.091 \pm .067$
1900	$0.170 \pm .058$	$0.091 \pm .043$
PFB/(PFA + P	FB) (dpm/dpm)	
6	$0.042 \pm .031$	$0.059 \pm .074$
1400	0.620 ± 1.160	$0.329 \pm .113$
1650	$0.279 \pm .095$	0.083 ± 061
1900	$0.146 \pm .049$	$0.084 \pm .040$
PFB/Dissolved	(dpm/dpm)	
6	0.154 ± 118	0.122 ± 160
1400	$2.69 \pm .87$	$0.175 \pm .059$
1650	$0.257 \pm .082$	$0.040 \pm .029$
1900	$0.422 \pm .174$	$0.069 \pm .033$

Table 2. Retention of thorium isotopes on prefilters in series (Sta. 423)

PFB/PFA: ratio of activity on second 0.5 μm wound fiber cartridge prefilter to first prefilter in series. PFB/(PFA \pm PFB): ratio of activity on second prefilter to total activity retained on both prefilters. PFB/Dissolved: ratio of activity on second prefilter to dissolved activity calculated from manganese dioxide-loaded cartridges.

the Nansen Basin via the Fram Strait. Thus

226
Ra = $[(1.71 \pm 0.11)$ Si + $74.2 \pm 0.9]\frac{S}{35}$ (2)

Where 226 Ra is the 226 Ra activity (dpm/1000 kg); Si is dissolved Si (μ mol/kg); S is salinity (‰)

The constants $(1.71 \pm 0.11) \frac{\text{dpm}^{26} \text{Ra}}{\mu \text{molSi}}$ and $74.2 \pm 0.9 \frac{\text{dpm}}{1000 \text{kg}}$ were obtained from the best fit relationship of ²²⁶Ra vs Si at TTO Stas 142–162. Values of ²²⁸Ra calculated using equation (2) and measured ²²⁸Ra/²²⁶Ra ratios (reported in Rutgers van der Loeff *et al.*, 1995) are given in Table 1.

Following the initial Th separation, the Ra fractions from a few of the manganese cartridges were retained for determination of 228 Ra by the 228 Th ingrowth method (Moore et al., 1985). The mean Ra extraction efficiency ($\pm 1\sigma$) determined for these samples was 31.3 \pm 7.2%. This value, together with the calculated 228 Ra activities (Table 1), has been used to correct the 228 Th activities to sample collection. Stations 287 and 358 were analyzed promptly after sample collection and the corrections are minimal (generally $\leq 15\%$) for these samples. The dissolved 228 Th analyses at Stas 370 and 423 entail significant Ra ingrowth corrections and are not presented.

Table 2 gives 232 Tn and 230 Th data on the paired prefilters deployed in series on four pumps at Sta. 423. Relative to the total activity retained on the two filters (PFB/PFB + PFA), the fraction on the second prefilter generally was $\leq 33\%$. Greater proportions were

noted on the samples collected at 1400 m. The proportion of 232 Th retained on the second prefilter is consistently greater than that of 230 Th relative to either the total particulate (PFA + PFB) or "dissolved" activities. 232 Th has no in-situ source in the oceans and is added in association with particles; the observation of greater proportions of 232 Th than 230 Th on the second prefilter makes it likely that the Th activity on the second prefilter is due to particles which pass the first and are retained on the second prefilter. For 230 Th, however, this fraction (PFB/Dissolved; Table 2) is $\leq 18\%$ of the dissolved 230 Th activity calculated from the two manganese oxide cartridges. For 232 Th, the fraction is generally $\leq 40\%$, but is greater in the 1400 m sample. Thus "leakage" of particles through the prefilter does not contribute significantly to the dissolved 230 Th in a typical deployment consisting of one prefilter and two Mn-treated cartridges, but the "dissolved" 232 Th values must be viewed as upper limits.

Total ²¹⁰Pb activities measured on aliquots from large volume water samples (Table 3) show relatively little variation with depth. The anthropogenic radionuclide data are given in Tables 4 and 5. For ^{137,134}Cs and ^{239,240}Pu, total activities are determined from the 60 l samples. Americium-241 activities are determined from the in-situ pump samples and utilize

Table 3.	210 Pb data	for Nansen	Basin water samples

Depth (m)	Total ²¹⁰ Pb* (dpm/1000kg) kg)	²²⁶ Ra† (dpm/1000kg)
Sta. 340 (82 59'1	N, 31 58.6'E, 3800 m)	
100	32.8 ± 3.2	80.7
450	44.5 ± 4.2	84.4
750	30.9 ± 3.6	86.6
1100	30.8 ± 4.5	89.7
1500	36.9 ± 4.7	91.4
1800	52.4 ± 6.3	92.7
2200	36.5 ± 5.4	93.6
2800	25.9 ± 3.3	93.8
3400	47.9 ± 4.9	94.4
3750	32.2 ± 3.9	94.6
3800	29.2 ± 4.5	94.6
Sta. 371(86 7.9')	N, 22 4.3'E, 3945 m)	
6	87.1 ± 1.8	75.4
100	45.6 ± 5.4	78.2
250	34.1 + 4.8	83.5
450	27.5 ± 4.6	84.4
850	33.8 ± 4.7	85.7
1300	29.4 + 3.9	88.3
1900	31.5 ± 4.4	91.7
2500	215 ± 2.2	92.7
3100	39.7 ± 3.0	93.4
3800	39.1 ± 3.1	93.6
3900	27.0 + 3.0	93.6
3930	27.4 ± 2.9	93.6

^{*}Dissolved plus particulate.

 $^{^+}$ Calculated from correlation between dissolved silicate and 226 Ra (see text).

	samples	
Depth (m)	²⁴¹ Am (dissolved) (dpm/1000 l)	^{239,240} Pu (total) (dpm/1000 l)
Sta. 287		
6	$0.116 \pm 0.014*$	$1.07 \pm 0.12*$
350	$0.113 \pm 0.013*$	$1.75 \pm 0.16*$
500	0.097 ± 0.015	1.61 ± 0.13
800		1.60 ± 0.13
1050	0.138 ± 0.011	-
1300	No. of Assets	1.62 ± 0.14
1450	0.129 ± 0.018	
1900		1.20 ± 0.10
2050	0.043 ± 0.020	
2150	0.036 ± 0.027	
Sta. 358		
20		1.00 ± 0.14
270	0.137 ± 0.009	1.14 ± 0.16
5''20	0.248 ± 0.044	1.44 ± 0.17
1120		1.16 ± 0.14
1500	0.148 ± 0.010	
2500	0.062 ± 0.009	0.54 ± 0.06
3400	0.027 ± 0.008	0.11 ± 0.08
39.50		0.57 ± 0.06
4000	0.018 ± 0.006	0.76 ± 0.09

Taole 4. Americium and plutonium analyses of Nansen Basin water samples

the cartridge extraction efficiency calculated from the Th data. Previous work has shown that Am and Th behave similarly with respect to their uptake on the Mn-impregnated cartridges (Cochran ϵt al., 1987).

DISCUSSION

Scavenging of reactive radionuclides—thorium isotopes and ²¹⁰Pb

In the Arctic Ocean, low particle concentrations and fluxes often have been cited as causing low scavenging efficiency of reactive radionuclides. Ku and Broecker (1967) advanced this hypothesis to explain the anomalously low ²³⁰Th and ²³¹Pa inventories in a deep-sea sediment core collected under ice island T-3 and Finkel *et al.* (1977) similarly used the argument to explain low ¹⁰Be inventories in Arctic Ocean cores. More recently Moore and Smith (1986) and Bacon *et al.* (1989) used water column data to show that scavenging of ²³⁰Th, ²¹⁰Po and ²¹⁰Pb is slower in the Canada Basin than in the Atlantic Ocean.

In contrast, the central Nansen Basin 230 Th profiles (Stas 358, 370; Fig. 2) are similar to those observed in the open North Atlantic Ocean (Cochran *et al.*, 1987). At Station 358, dissolved 230 Th increases to relatively constant values of $\sim 0.5-0.6$ dpm/1000 l at depths greater than 2500 m (Fig. 2). The ratio of particulate to dissolved 230 Th ranges from 0.13 to 0.25, also similar to the north-west Atlantic. The profile at Sta. 358 is comparable to one

^{*}Particulate 241 Am: 6 m, 0.006 \pm 0.001 dpm/1000 l; 350 m, 0.011 \pm 0.002 dpm/1000 l.

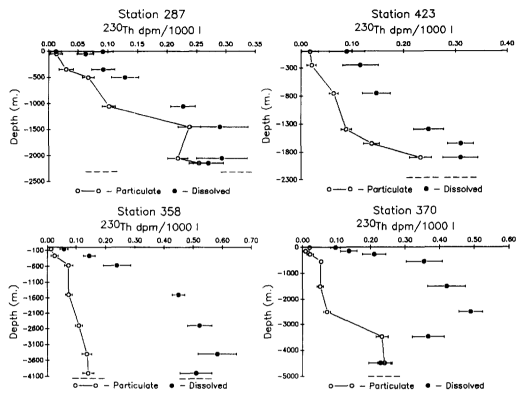
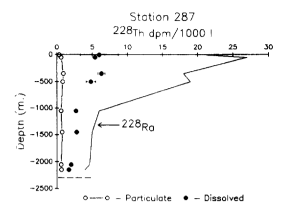


Fig. 2. Thorium-230 profiles obtained from in-situ pump casts at four stations in the Nansen Basin: Sta. 287, 423—Barents Sea slope, Sta. 358—Central Basin, Sta. 370—Gakkel Ridge. Dashed lines indicate bottom depths; open circles indicate particulates ($> 0.5 \mu m$); filled circles indicate dissolved material.

obtained from the central Nansen Basin in 1991 (Scholten et al., 1994), indicating good intercalibration. At Sta. 370 in the vicinity of the Gakkel Ridge, dissolved 230 Th increases to $\sim 0.5 \text{ dpm/1000 l}$ at a depth of about 2500 m then decreases toward the bottom. This decrease in dissolved 230 Th is complemented by a marked increase in particulate 230 Th, suggesting enhanced scavenging associated with the ridge (Kadko et al., 1986; Shimmield and Price, 1988). Indeed Schlosser et al. (1990) have identified the Gakkel Ridge as a possible source of primordial 3 He to the deep Nansen Basin, and hydrothermal activity or nepheloid layers associated with the ridge may explain the enhanced removal of 230 Th.

In the marginal areas of the Nansen Basin, scavenging of ²³⁰Th is enhanced relative to the basin interior. This is seen at Sta. 287, taken close to the Barents Sea continental slope (Fig. 2), where dissolved ²³⁰Th activities do not exceed 0.3 dpm/1000 l. Particulate activities also show high values, producing particulate/dissolved ²³⁰Th ratios ranging from 0.1 to 0.6, a factor of about 2 greater than in the central basin. Similarly, dissolved ²³⁰Th is low and the proportion of particulate to dissolved ²³⁰Th is high at Sta. 423, taken near the slope off Svalbard.

These trends in ²³⁰Th are reflected in the ²²⁸Th data (Fig. 3). The two stations for which dissolved ²²⁸Th could be calculated reliably, 287 and 358, provide examples of both basin



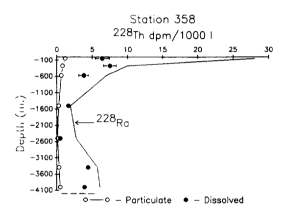


Fig. 3. Thorium-2:28 profiles at the Barents Sea slope (Sta. 287) and Central Nansen Basin (Sta. 358). Open circles ir dicate particulates (>0.5 μm); filled circles indicate dissolved ²²⁸Ra activities (solid line) determined from measured ²²⁸Ra/²²⁶Ra activity ratios (Rutgers van der Loeff *et al.*, 1995) and calculated ²²⁶Ra activities (see text). Dashed lines indicate bottom depths.

margin (Sta. 287) and central basin (Sta. 358) profiles. Particulate 228 Th is greater and dissolved 228 Th is somewhat lower at Sta. 287 than at 358. The total 228 Th is lower in the deep water column at Sta. 287 than at 358, indicating enhanced removal at Sta. 287. Both stations show low 228 Th/ 228 Ra activity ratios in the upper 50 m. At Sta. 358 the ratios approach equilibrium at depth, while at Sta. 287 they remain < 0.6.

Assuming that the Th isotope profiles are steady state, residence times for both dissolved and particulate Th car. be calculated from the data (Coale and Bruland, 1987). The residence time of dissolved Th with respect to scavenging onto particles, τ_d , is

$$\tau_{\rm d} = \frac{1}{\lambda} \frac{A_{\rm d}}{(P - A_{\rm d})} \tag{3}$$

where λ is the Th decay constant (year⁻¹), A_d is the dissolved ²²⁸Th or ²³⁰Th activity (dpm/l) and P is the parent activity (²²⁸Ra or ²³⁴U; dpm/l)

The residence time of particulate Th is

$$\tau_{\rm p} = \frac{A_{\rm p}}{\lambda [P - (A_{\rm d} + A_{\rm p})]} \tag{4}$$

where $A_{\rm p}$ is the particulate ²²⁸Th or ²³⁰Th activity (dpm/l).

Equations (3) and (4) both depend on the extent of disequilibrium between parent and daughter Th isotope. In the Nansen Basin deep water, 228 Th is close to equilibrium with 228 Ra (228 Th/ 228 Ra \sim 0.8) and is consequently not a useful tracer for calculating τ s. In the surface water, 228 Th/ 228 Ra disequilibrium is greater and this isotope can be used. Thorium-230 displays disequilibrium relative to 234 U throughout the water column, but in the surface water "dissolved" 230 Th activities are low and often are not significantly greater than "dissolved" 232 Th (Figs 3 and 4). Both 230 Th and 232 Th can be present in the "dissolved" fraction in surface waters by leaching of atmospherically supplied dust particles (Cochran et al., 1987) or through particles finer than 0.5 μ m that pass the prefilter but may be retained on the manganese cartridges. The correction for 234 U-supported 230 Th is large in these samples and 230 Th is consequently not a sensitive tracer for calculating scavenging rates in surface water.

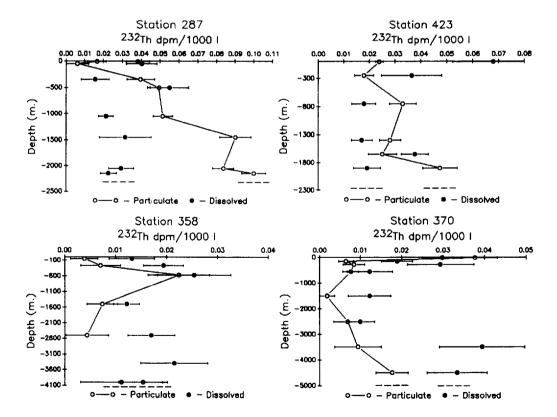


Fig. 4. Particulate and dissolved 232 Th profiles in the Nansen Basin. Open circles indicate particulates (> 0.5 μ m); filled circles indicate dissolved material; dashed lines indicate bottom depths.

Table 5 gives residence times for dissolved and particulate Th in surface water (calculated from 228 Th/ 228 Ra disequilibrium) and deep water (calculated from 230 Th/ 234 U disequilibrium) for the stations sampled. Corrections for supported 230 Th were made to the particulate activities by assuming that the activity ratio of supported 230 Th to 232 Th is 0.6 (R. Anderson, personal communication). This correction is generally $\leq 10\%$ of the particulate 230 Th activity in the central basin and ranges up to 25% at Sta. 287. The Eurasian Basin Deep Water (Stas 358 and 370) has a dissolved Th residence time with respect to scavenging of 18–19 years. Within the Gakkel Ridge complex, the value drops to ~ 12 years near the bottom. Dissolved 230 Th in deep water of the Barents Sea slope has mean residence times of ~ 10.5 –12 years, indicating more rapid removal onto particles in this ocean margin environment. Particulate Th residence times are longer (~ 6 years) in the slope deep water relative to the basin interior (~ 2 –4 years), probably indicating greater particle resuspension rates caused by slope boundary currents. In the surface water, dissolved Th has a mean residence time of about 1 year in both central basin and slope environments, and particulate Th is removed rapidly (0.06–0.2 years).

The mean residence time of total ²¹⁰Pb with respect to scavenging and particle removal also may be calculated using a variation of equation (4):

$$\tau_{Pb} = 32.3 \frac{A_{Pb}}{(A_{Ra} - A_{Pb})} \tag{5}$$

Data are available for two central basin stations, 340 and 371 (Fig. 5). Total ²¹⁰Pb activities in the deep water (1500–3000 m) average 35 \pm 10 dpm/1000 l. Relative to average ²²⁶Ra activities of ~9.3 dpm/1000 l (Table 3), τ_{Pb} is about 19 years. This value is similar to that for the North Atlantic (Cochran *et al.*, 1990).

The general increases in both dissolved and particulate ²³⁰Th with depth support the hypothesis that a sorption equilibrium is occurring between dissolved and particulate Th (Nozaki *et al.*, 1981; Bacon and Anderson, 1982; Bacon *et al.*, 1985; Nozaki *et al.*, 1987; Bacon *et al.*, 1989). Using box models of the dissolved and particulate forms of ²³⁰Th and ²²⁸Th gives

$$\frac{A_{\rm p}}{A_{\rm d228}} = \frac{k_1}{k_{-1} + \lambda_{228}} \tag{6}$$

$$\frac{A_{\rm p}}{A_{\rm d230}} \ge \frac{k_1}{k_{-1}} \tag{7}$$

where A_p is the particulate ²³⁰Th or ²²⁸Th (dpm/1000 l), A_d is the dissolved ²³⁰Th or ²²⁸Th (dpm/1000 l), k_1 is the adsorption rate constant (year⁻¹), k_{-1} is the desorption rate constant (year⁻¹), λ_{228} is the decay constant for ²²⁸Th (0.3647 year⁻¹).

Equation (7) assumes that $\lambda_{230} \Theta k_{-1}$, and if λ_{228} is comparable to k_{-1} , individual values for k_1 and k_{-1} can be calculated from the data. If $\lambda_{228} \Theta k_{-1}$, only the quotient k_1/k_{-1} can be calculated.

The mean $A_{\rm p}/A_{\rm d}$ values below 500 m (Table 1), corrected for supported particulate Th using 232 Th, are: $A_{\rm p}/A_{\rm d}\Theta^{228}=0.23\pm0.06$ and 0.13 ± 0.04 , and $A_{\rm p}/A_{\rm d}\Theta_{230}=0.51\pm0.19$ and 0.22 ± 0.04 for Stas 287 and 358, respectively. From equations (6) and (7), we obtain $k_1=0.15\pm0.07$ and 0.12 ± 0.07 year $^{-1}$ and $k_{-1}=0.29\pm0.17$ and 0.55 ± 0.34 year $^{-1}$ at Stas 287 and 358, respectively. The large errors associated with the estimates of k_1 and k_{-1}

Th
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Residence
Table 5.

Location (10								
	A _{part} 10 ³ dpm/l)	Tpart (year)	Adiss (10 3 dpm/l)	t _{diss} (year)	Apart* (10 3dpm/l)	Tpart (year)	$A_{\rm diss}$ (10 $^3{ m dpm/l}$)	τ _{diss} (year)
	1.2 ± 0.06	0.20 ± 0.01	7.2 ± 1.2	1.0 ± 0.13			Manage of the Control	1
EBDW (1500-2500 m)†	1		ŀ	İ	0.088	3.4	0.486	18.9
EBBW (3400-4000m)†	-		i		0.135	5.3	0.546	21.2
Sta. 370 (Gakkel Ridge) EBDW (1500–2500 m)†				I	0.060	2.3	0.455	17.7
EBBW (3500-4500 m)†	I	1	1	1	0.228	6.8	0.298	11.6
lope								
Surface water 0.3 Deepwater (1450–2150 m)	$0.341 \pm .041$	0.06 ± 0.01	7.3 ± 3.3 —	1.1 ± 0.5	$\frac{-}{0.182 \pm 013}$	7.1 + 0.5	$\frac{-}{0.270 + .030}$	$\frac{-}{10.5 + 1.2}$
Sta. 423 Deepwater (1650-1900 m)	1	I	l	I	0.163	5.8	0.310	12.1

*Corrected for supported ²³⁰Th using measured particulate ²³²Th activities and assuming (U-supported ²³⁰Th/²³²Th) activity ratio in particles = 0.6. †EBDW = Eurasian Basin Deep Water, EBBW = Eurasian Basin Bottom Water. Activities are average activities for the depths indicated.

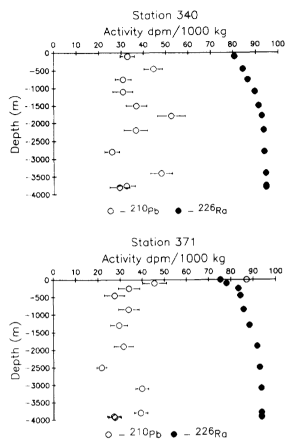


Fig. 5. Total ²¹⁰Pb (particulate + dissolved; open circles) at two stations in the Nansen Basin. Sta. 340—Central Basin, Sta. 371—Gakkel Ridge. Filled circles ²²⁶Ra activity calculated from Ra vs Si correlation (see text).

make it difficult to discern differences at the two stations. However, the calculated values of k_1 and k_{-1} are within the range of values from open ocean sites in the North Atlantic and North Pacific (Murnane *et al.*, 1994) and reinforce the notion that scavenging of Th is not markedly different in the Nansen Basin from the Atlantic or Pacific.

Although the residence times calculated for Th in the Nansen Basin surface water are not significantly different from values obtained for the open North Atlantic Ocean (Kaufman et al., 1973; Broecker et al., 1973; Li et al., 1980), they do stand in contrast with the results of Bacon et al. (1989), who found Th residence times of 6 years in the surface waters of the Alpha Ridge, separating the Canada and Makarov Basins. In the deep water, Th residence times of about 20 years in the central Nansen Basin are significantly shorter than values of 55 years estimated by Scholten et al. (1994) for the Makarov Basin. Several factors may explain these differences: (i) particle fluxes are greater in the Nansen Basin than in the Makarov Basin; (ii) ventilation time scales are more rapid for the Nansen Basin than the Makarov Basin; (iii) the Nansen Basin margin, specifically the Barents Sea slope, is a site of enhanced removal of Th and this sink influences the basin's interior.

There are few data on particle fluxes in the Arctic Ocean, but given the ice coverage of both the Amerasian (Canada and Makarov) and Eurasian (Amundsen and Nansen) Basins there seems no reason to expect particle fluxes to be different a priori. There are, however, well documented differences in ventilation times of the basins. In the Nansen Basin, the chlorofluoromethane (freon) data of Wallace et al. (1992) show detectable F-11 data throughout the water column at stations along the Barents slope (Sta. 287) and Gakkel Ridge (Sta. 370), with deep water (1000-3000 m) ages of 15-30 years. At Sta. 358 in the basin interior, deep water ages are about 5 years older than at the basin boundaries. FREON is undetectable in the Eurasian Basin bottom water (EBBW: > 3000 m), but this water mass has measurable CCl₄ and tritium, with a $T/^{3}$ He age of 40–50 years (Schlosser et al., 1990). However, the radiocarbon age of this water is 200-250 years (Schlosser et al., 1990). Even in the central Nansen Basin, water column FREON inventories are relatively high compared with the central Canada Basin, indicating more rapid ventilation of the former (Wallace and Moore, 1985; Wallace et al., 1992). Indeed, ventilation of Nansen Basin Deep Water on a timescale not very different from the scavenging residence time of Th will tend to cause the ²³⁰Th profiles to reach a constant value, as observed (Fig. 2), rather than the linear increase seen in the deep Pacific for example (Nozaki et al., 1981). The ²³⁰Th activity of the basin deep water thus represents a balance between in-situ production, scavenging and mixing with water with low ²³⁰Th.

Our data also show evidence that removal of ²³⁰Th is enhanced at the Barents Sea slope stations (287 and 423). Total ²³⁰Th is slightly lower at these stations than in the basin interior (Sta. 358), but greater differences are apparent between dissolved and particulate activities. Enhanced scavenging of Th near the margin of the Nansen Basin may be due in part to higher particle concentrations and fluxes there. Although particle concentration data are not available for the ARK IV/3 samples, evidence for high values near the margin is seen in the particulate ²³²Th distributions (Fig. 4). Unlike ²³⁰Th and ²²⁸Th, ²³²Th is not produced *in situ* in the oceans, but rather is added in association with continentally-derived detrital particles. The concentrations of particulate ²³²Th at the margin Stas 287 and 423 are factors of 3 to 8 greater than those at the interior Stas 358 and 370 (Table 1). Elevated particle concentrations near the basin margin, if coupled with greater particle fluxes, could produce enhanced removal of reactive radionuclides from the water column.

Americium-241 and ^{239,240}Pu distributions

The distributions of the transient tracers 241 Am and 239,240 Pu provide additional information on the pattern of scavenging suggested from Th isotope distributions in the Nansen Basin. At the Barents Sea slope station (287), Pu shows a broad maximum over the depth interval 350–1300 m. Americium-241 activities display more variation within this depth range, with a maximum at ~ 1000 m (Fig. 6). In contrast, both Pu and Am show distinct maxima at ~ 600 m in the central Nansen Basin (Sta. 358; Fig. 6). Moreover, 241 Am/ 239,240 Pu ratios are lower at the Barents slope than in the central Nansen Basin.

These patterns must be viewed in the context of the sources of these anthropogenic radionuclides to the Arctic. Livingston et al. (1984) have identified two important sources as direct atmospheric fallout from the testing of atomic weapons and advective supply from the North Atlantic. The former arrived in the period 1956–1966 with a peak in 1963–1964, and the latter is a continuous input that has varied over time due to the contribution of the discharge of these radionuclides from the nuclear fuel reprocessing facilities at Sellafield,

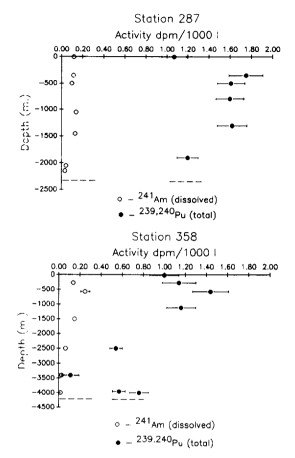


Fig. 6. Americium-241 (dissolved) and ^{239,240}Pu (total) profiles in the Nansen Basin. Sta. 287—Barents Sea slope, Sta. 358—Central basin Samples for ²⁴¹Am were collected using *in situ* pumps and samples for Pu were taken from large volume water casts at the same stations.

U.K. (Livingston et al., 1982). Sellafield inputs of 137 Cs, 239 240 Pu and 241 Am were particularly high from 1974 to 1978, and during the period 1969–1978 greatly exceeded the fallout delivery (Livingston et al., 1984). An additional source of 137 Cs was the Chernobyl reactor accident in April 1986 (see discussion below). The 239,240 Pu inventories at Stas 287 and 358 are ~ 50 Bq/m² (Table 6), about a factor of 10 greater than the direct fallout input at these latitudes (Livingston et al., 1984).

Livingston et al. (1984) documented the presence of Sellafield ¹³⁷Cs in the Amundsen Basin and determined a transit time of about 5 years from the Irish Sea to the Barents Sea and an additional 3 years to the Amundsen Basin, the northernmost basin of the Eurasian Basin (Fig. 1). Tritium/helium (Schlosser et al., 1990) and chlorofluoromethane data (Anderson et al., 1989; Wallace et al., 1992) also indicate fairly rapid ventilation (< 50 years) of the Nansen Basin for depths less than 3500 m (see below). Thus it is likely that the ^{239,240}Pu distributions in the Nansen Basin are continuing to evolve, but have been characterized by high inventories since the 1970s.

	Reference	Water Depth (m)	Date	$\Sigma^{239,240} Pu$ (Bq/m ²)	Σ^{241} Am (Bq/m ²)	Am/Pu (Bq/m ²)	Σ ¹³⁷ Cs pre- Chernobyl	Σ ¹³⁷ Cs post- Chernobyl
Arctic Ocean								
Nansen Basin (Sta. 358)	This study	4050	1987	52	7.3	0.14	7811	8749
Barents Sea Slope (Sta. 287)	This study	2250	1987	55	3.9	0.07	8367	13,740
Fram Basin	Livingston et al. (1984)	4000	1979	35	6.4	0.18	6790	
Makarov Basin	Livingston et al. (1984)	4000	1979	24	6.2	0.26	3670	
Northwest Atlantic	Ocean							
Nares Abyssal Plain	Cochran <i>et al.</i> (1987)	5840	1984	44	12	0.27	2920	
Hatteras Abyssal	Cochran et al.	5400	1980-81	93	13	0.14	4770	

Table 6. Anthropogenic radionuclide inventories in the Arctic and north-west Atlantic oceans

Americium-241 shows the opposite trend from Pu or ¹³⁷Cs, namely higher activities and water column inventories in the central basin (Sta. 358) than at the margin (Sta. 287, Table 6). Because ²⁴¹Am is produced from ²⁴¹Pu decay, this pattern can be explained in the context of rapid supply of Pu to the Nansen Basin via the Barents Sea followed by production of ²⁴¹Am in the lower scavenging environment of the central basin. Americium-241 that accompanied the Pu or was produced *en route* was likely scavenged in the rapid scavenging regimes of the Barents shelf or slope, consistent with the observation of lower ²⁴¹Am inventories at slope Sta. 287.

Mixing timescales in the Nansen Basin: Evidence from Cs isotope distributions

We have argued that the particle-reactive tracer distributions in the Nansen Basin are affected by the relatively rapid mixing or ventilation timescales for the Eurasian Basin Deep Water (EBDW). These timescales are estimated to be less than about 50 years, based on FREON, CCl_4 and $T/^3$ He distributions (Wallace *et al.*, 1992). The cesium isotope data provide additional constraints on ventilation timescales of the Nansen Basin.

Two important sources of 137 Cs to the Arctic Ocean are direct fallout from nuclear weapons testing and Cs advected from the Sellafield reprocessing facility (Livingston *et al.*, 1984, 1985). The ARK IV/3 cruise was about one year after the accident at the Chernobyl nuclear power generating station, and the Chernobyl accident constitutes another potential source of 137 Cs to the Nansen Basin. The presence of Chernobyl-derived Cs is evident in the high 134 Cs/ 137 Cs activity ratios in the upper water column (0.1–0.2, Table 7). In comparison, corresponding pre-Chernobyl 134 Cs/ 137 Cs ratios measured in the same area in 1981 did not exceed 0.03 (Casso and Livingston, 1984). The Sellafield source term has been declining, and the 134 Cs/ 137 Cs would have decreased to \leq 0.005 in 1987 in the absence of Chernobyl input. The short half-life of 134 Cs (2.1 years) places constraints on the rate of mixing, and we note that 134 Cs is present to 1000 m at the Barents slope stations (287, 423, Fig. 7) but is mixed to

Table 7. Cesium isotope data for Nansen Basin water samples

	¹³⁷ Cs	¹³⁴ Cs	Pre-Chernobyl ¹³⁷ Cs
Depth	(dpm/10001)		(dpm/1000 l)
Sta. 287			
6	702 ± 24	82 ± 7	477 ± 27
50	708 ± 9	78 ± 2	495 ± 11
100	631 ± 22	65 ± 6	453 ± 25
350	702 ± 16	78 ± 4	489 ± 21
400	583 ± 14	62 ± 3	412 ± 16
500	599 ± 10	70 ± 3	407 ± 12
800	149 ± 7	$\frac{-}{2.2 \pm 1.1}$	143 ± 8
1050	150 ± 7	0.8 ± 1.3	1 4 8 ± 8
1450	95 ± 7	-0.1 ± 1.6	95 ± 8
1850	64 ± 1	-0.6 ± 2.0	64 ± 1
2050	61 ± 4	-0.9 ± 1.9	61 ± 4
2150	61 ± 8	-1.1 ± 3.0	61 ± 8
Sta. 358			
6	762 ± 8	9.4 ± 1.2	737 ± 8
50	1090 ± 10	105 ± 4	803 ± 13
270	440 ± 5	5.9 ± 0.6	423 ± 5
400	384 ± 10	-0.1 ± 1.3	384 ± 10
570	240 ± 8	1.3 ± 1.3	236 ± 8
870	193 ± 37	-15 ± 16	193 ± 37
1500	59 ± 4	0.1 ± 1	59 ± 4
2482	17 ± 2	0.1 <u>1</u> 1	16 ± 4
2500	11 ± 1	0 ± 0.8	10 ± 4 10 ± 2
3100	6 ± 2	$\frac{0 \pm 0.8}{1.8 \pm 1.7}$	0.5 ± 1.0
3372		1.0 ± 1.7	9 ± 2
3400	11 ± 2 8 ± 4	0.6 ± 0.8	6 ± 2
3916		0.0 ± 0.0	
	9 ± 2	20 1 21	$\begin{array}{c} 6 \pm 2 \\ 101 \pm 3 \end{array}$
3950	15 ± 4	2.0 ± 2.1	
3966 4000	14 ± 4 150 ± 10	48 ± 4	9 ± 4 181 17
Sta. 370			
6	557 ± 7	4.4 ± 1	545 ± 7
275	401 ± 13	-0.6 ± 2.2	401 ± 13
550	204 ± 11	-1.2 ± 1.9	204 ± 11
1500	111 ± 5	1.5 ± 1.3	107 ± 5
2500	45 ± 3	-1.6 ± 10	45 ± 3
3500	$\frac{45 \pm 3}{31 \pm 9}$	0.6 ± 3.4	29 ± 9
4500	$\frac{31 \pm 9}{27 \pm 3}$	-1.2 ± 1	27 ± 3
Sta. 423			
6	859 ± 136	55.9 ± 9	706 ± 138
250	596 ± 95	46.3 ± 8	469 ± 197
750	353 ± 56	9.1 ± 1.9	328 ± 56
1400	69 ± 11	0.2 ± 0.8	69 ± 11
1650	_	-0.6 ± 0.7	51 ± 8
1900	51 ± 8		
2150	66 ± 11	0.4 ± 1.2	65 ± 11
413U	4 ± 8		4 ± 8

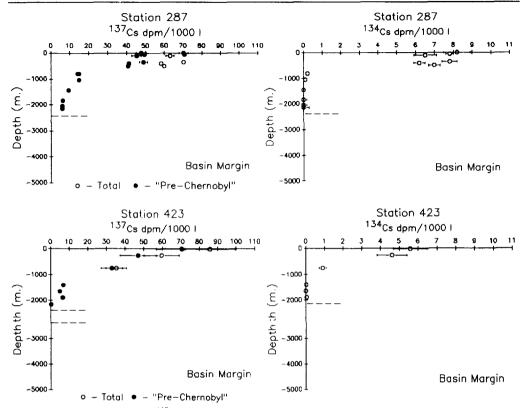


Fig. 7. Cesium-134 and ¹³⁷Cs profiles from the Nansen Basin. Open circles correspond to activities measured in samples collected in 1987. Filled circles indicate pre-Chernobyl ¹³⁷Cs activities calculated from ¹³⁴Cs data (see text).

depths of < 300 in the basin interior (Stas 358, 370, Fig. 7). Because about 1 year had elapsed since the introduction of this 134 Cs to the Nansen Basin, mixing of 134 Cs to these depths was accomplished within 1 year.

The EBDW at Stas 358 and 370 is generally unlabelled by Chernobyl Cs at the time of sampling, but an interesting exception to this pattern is the near-bottom sample at Sta. 358, which shows the highest ¹³⁴Cs/¹³⁷Cs measured at any of the stations (Fig. 7). Cesium is only weakly reactive in seawater (Bowen *et al.*, 1980), but we attribute this high ¹³⁴Cs to Cs associated with particulate material from the Chernobyl accident that sank rapidly to the bottom. Although sea ice coverage of the Nansen Basin is heavy in the winter, breakup is common in the spring and summer, allowing particles trapped by the ice to enter the water column and sink. Moreover, Sta. 358 is located in one of the branches of the Transpolar Ice Drift, with most of the ice in this area originating in Siberian shelves in the Laptev Sea (Thiede, 1988; Anderson and Jones, 1992). It is possible that particulate Cs from Chernobyl could have labeled the ice in this area and been rapidly transported to Sta. 358.

The ¹³⁴Cs/¹³⁷Cs ratios, corrected for decay to the date of the Chernobyl accident, can be coupled with the ¹³⁴Cs/¹³⁷Cs ratio of Cs released by the accident to determine the "pre-Chernobyl" profile of ¹³⁷Cs at the four stations (Fig. 7; Table 7). The profiles show that pre-Chernobyl ¹³⁷Cs is present throughout the water column at the slope Stas 287 and 423, but is

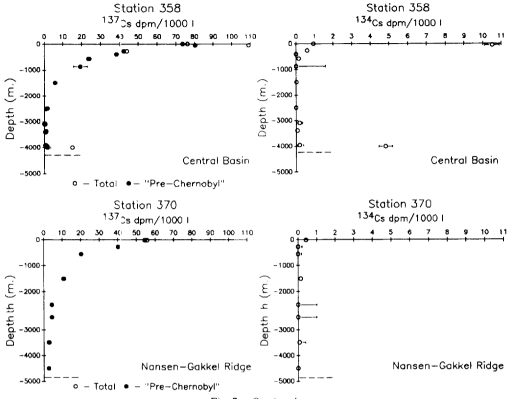


Fig. 7. Continued.

present at very low activities ($\leq 10 \text{ dpm}/1000 \text{ l}$) in the deep central basin. Station 370, the Gakkel Ridge station, shows higher ¹³⁷Cs at depth relative to Sta. 358, and this pattern is consistent with ²²⁸Ra (Rutgers van der Loeff *et al.*, 1995) and freon data (Wallace *et al.*, 1992), which show deeper mixing in the vicinity of the Nansen Ridge. The presence of very low ¹³⁷Cs activities in the EBDW are consistent with the ages of about 50 years for this water estimated from T/³He and CCl₄ distributions (Schlosser *et al.*, 1990; Wallace *et al.*, 1992).

Inventories of pre-Chernobyl ¹³⁷Cs in the Nansen Basin are greater than those measured in 1979 in the Fram and Makharov Basins (Table 6). The latter in turn are greater than those measured in the Northwest Atlantic in the early 1980s (Table 6). This pattern emphasizes the importance of the Sellafield source of ¹³⁷Cs to the Nansen Basin. The Barents Sea slope and Nansen Basin have been strongly labeled with this source to depths of at least 1500 m in the central basin and throughout the water column at the Barents slope. This implies ventilation timescales of 9–18 years for these waters, given the release pattern of ¹³⁷Cs from Sellafield (Livingston *et al.*, 1982).

CONCLUSIONS

The distributions in the Nansen Basin of the naturally occurring thorium isotopes ²³⁰Th and ²²⁸Th and the anthropogenic radionuclide ²⁴¹Am suggest that scavenging is less intense in the Eurasian Basin Deep Water in the central basin than at the basin margins (e.g. Barents

Sea slope). Residence times of dissolved Th in the basin deep water are 18–19 years in the central Nansen basin and 10–12 years on the Barents Sea slope. Mixing timescales are estimated from ¹³⁴Cs and ¹³⁷Cs distributions. Cesium-134 derived from the accident at the Chernobyl nuclear power station 1 year prior to sampling has been mixed to depths of 1000 m at the Barents Slope and to ~300 m in the central Nansen Basin. Pre-Chernobyl ¹³⁷Cs, derived from fallout from atmospheric testing of atomic weapons and the nuclear fueled reprocessing facility at Sellafield, U.K., has been mixed throughout the water column at all stations, but is present in very low activities in the Eurasian Basin Deep Water in the central Nansen Basin. This pattern is consistent with ages of about 50 years for this water mass, based on T/³He and FREON distributions (Schlosser *et al.*, 1990; Wallace *et al.*, 1992). The relatively rapid mixing time of Eurasian Basin Deep Water in the Nansen Basin, coupled with the texture and rate of Th scavenging within the basin, maintains dissolved ²³⁰Th activities at a lower value in the Nansen Basin compared with previously reported values in the deep Canadian Basin.

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