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Distribution of ²²⁶Ra in the Arctic Ocean and the Bering Sea and its hydrologic implications

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Abstract Radium-226 (²²⁶Ra) activities were measured in the surface water samples collected from the Arctic Ocean and the Bering Sea during the First Chinese National Arctic Research Expedition. The results showed that ²²⁶Ra concentrations in the surface water ranged from 0.28 to 1.56 Bq/m³ with an average of 0.76 Bq/m³ in the Arctic Ocean, and from 0.25 to 1.26 Bq/m³ with an average of 0.71 Bq/m³ in the Bering Sea. The values were obviously lower than those from open oceans in middle and low latitudes, indicating that the study area may be partly influenced by sea ice meltwater. In the Bering Sea, ²²⁶Ra in the surface water decreased northward, probably as a result of the exchange between the ²²⁶Ra-deficient sea ice meltwater and the ²²⁶Ra-rich Pacific water. In the Arctic Ocean, ²²⁶Ra in the surface water increased northward and eastward. This spatial distribution of ²²⁶Ra reflected the variation of the ²²⁶Ra-enriched river component in the water mass of the Arctic Ocean. The vertical profiles of ²²⁶Ra in the Canadian Basin showed a concentration maximum at 200 m, which could be attributed to the inputs of the Pacific water or/and the bottom shelf water with high ²²⁶Ra concentration. This conclusion was consistent with the results from ²H, ¹⁸O tracers.

Keywords: Arctic Ocean, Bering Sea, ²²⁶Ra, distribution.

The Arctic Ocean, the northernmost parts of the earth, covers the total surface area of 14.79 million square kilometers and amounts to only about 4% of global ocean surface area. Although its surface area is the smallest in the four major oceans, the Arctic Ocean plays an important role in the regulation of global climate, material balance and energy circulation of global ecosystem. The Arctic Ocean has seasonal or long-term ice cover, extended periods of darkness during the Arctic winter and the wide continental shelves covering about 36% of the total surface area of the Arctic Ocean, a percentage unparalleled by other oceans. These unique features have led to the recent interest in the hydrology of the Arctic Ocean.

The major water inputs to the Arctic Ocean are the Atlantic water, river water, and the Pacific-originated water entering through the Bering Strait. The Atlantic water is characterized by its high salinity and temperature. The high freshwater component in Arctic surface water, to a large extent, is due to river runoff to the Arctic Basin, which amounts to 10% of global river discharge^[1]. Additional freshwater inputs come from sea ice meltwater and precipitation. Among the freshwater components of the Arctic surface water, sea ice meltwater and meteoric water can be distinThe upper halocline water between 50 and 200 m of the Arctic Basin, which is characterized by a constant temperature at most the freezing point and a salinity that increases with depth, has interested many oceanographers. Jones and Anderson^[4] showed that the upper halocline water is associated with a maximum in silicate, phosphate, nitrate, normalized total carbonate and a minimum in oxygen at the CESAR Ice Station (81°43.4′N, 93°25′W). Moore and Smith^[5] showed that this layer also had a maximum in ²²⁶Ra and a minimum in ²¹⁰Pb at this station. Temperature, salinity, NO/PO ratio, ²H and ¹⁸O tracers have been used to investigate the sources and mechanism of the upper halocline water.

²²⁶Ra (half-life = 1622 a) has been successfully used to study circulation and mixing of the ocean water masses. It was suggested that ²²⁶Ra, just like ¹⁴C, was an ideal naturally occurring radionuclide for water movement in the ocean. With the implementation of several international research plans, such as GEOSECS, WOCE, and JGOFS, ²²⁶Ra activities in the world oceans have been determined extensively. However, very few data about ²²⁶Ra in the Arctic Ocean and the Bering Sea have been reported. Tsunogai and Harada^[6] determined ²²⁶Ra and ²¹⁰Pb activities in the samples collected from four stations in the Bering Sea, and found that a maximum in ²²⁶Ra occurred in mid-depth waters. Moore and Smith^[5] studied the disequilibria among ²²⁶Ra, ²¹⁰Pb and ²¹⁰Po at the CESAR Ice Station (81°43.4′N, 93°25′W), and found that the ²²⁶Ra activities increased to a maximum at depths corresponding to the nutrients maximum and upper halocline. By measuring ²²²Rn and ²²⁶Ra in the southeastern Bering Sea shelf waters and sediments, Glover and Reeburgh^[7] pointed out that the ²²⁶Ra activities showed a heterogeneous distribution in the southeastern Bering Sea shelf and a positive correlation with dissolved silicate.

In this study, thirty-three surface water samples and twelve vertical samples were collected from the Arctic Ocean and the Bering Sea during the First Chinese National Arctic Research Expedition conducted from July to September in 1999. The spatial distribution of ²²⁶Ra in surface waters gave information about the influence of sea ice meltwater on ²²⁶Ra activities. The relationship between ²²⁶Ra activities and dissolved silicate concentrations was evaluated. In addition, the vertical profiles of ²²⁶Ra were used to provide clues to the possible sources of the upper halocline water in the Canadian Basin.

1 Methods

1.1 Sample location

During the First Chinese National Arctic Research Expedition (from July to September in 1999), 13 and 20 large-volume surface water samples were collected in the Arctic Ocean and the Bering Sea, respectively (fig. 1). Sampling stations covered the Chukchi Sea (9 stations), the Beaufort Sea (Station C30 and Station C33), the Canadian Basin (Station C34 and Station C39), the western Bering Sea (18 stations) and the Norton Sound (Station BJ10).

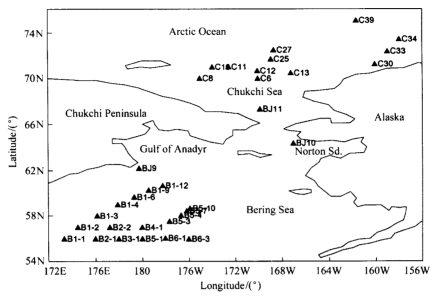


Fig. 1. Location of sampling stations in the Arctic Ocean and the Bering Sea.

1.2 Sample collection

120 dm³ surface water samples were collected and passed through a PVC pipe packed with the MnO₂-impregnated fibers. Flow rate was controlled at 200—250 cm³/min in order to absorb radium isotopes quantitatively. After the enrichment, MnO₂-fibers were taken out and enveloped using plastic bags. The fibers were taken back to the onshore laboratory for ²²⁶Ra measurement.

Vertical samples at Station C34 and Station C39 in the Canadian Basin were obtained by hanging MnO₂-fibers at different depths. After certain hours, MnO₂-fibers were taken out and enveloped using plastic bags. The fibers were taken back to the shore-based laboratory.

1.3 Methodology

1.3.1 ²²⁶Ra analysis. In the laboratory, MnO_2 -fibers were taken out from the plastic bags. Residual waters in the fibers were removed. The fibers were filled into a diffusion tube, which was subsequently sealed and vacuumized. After 5 –7 d, when ²²²Rn ingrown reached a significant level, a vacuumized scintillation counting cell was sequentially connected to the diffusion tube packed with MnO_2 -fibers and an airflow adjustor. ²²²Rn was emanated from MnO_2 -fibers into the counting cell for 15 min. After sealing in the cell for 3 h, ²²²Rn will be in equilibrium with his daughters. The activities were subsequently measured by using a Rn-Th analyzer (FD-125)^[8].

The specific activity of 226 Ra, A_{226} , was calculated from the following equation:

$$A_{226} = \frac{k_{226}(N_{\rm s} - N_{\rm b})}{\boldsymbol{a} \cdot \boldsymbol{V} \cdot \boldsymbol{h} \cdot \boldsymbol{t}},$$

where A_{226} is the specific activity of ²²⁶Ra in the water sample (Bq/m³), k_{226} is the counting coefficient for ²²⁶Ra (Bq/cpm), N_s and N_b are the total counts of the sample and the background respectively.

tively (with a consistent counting time *t*), **a** is the ingrown coefficient of ²²²Rn; *V* is the sample volume (m³), **h** is the emanation efficiency of ²²²Rn from MnO₂-fibers, *t* is the counting time of the sample and the background (min).

1.3.2 Influence of residual water in MnO₂-fibers upon ²²⁶Ra measurement. In our laboratory, the aforementioned method for ²²⁶Ra measurement has been set up for many years. However, the procedure for removing residual water in MnO₂-fibers was different for different samples. In order to check the influence of this procedure on ²²⁶Ra measurement, five vertical samples from Station C34 were selected for measuring ²²⁶Ra in MnO₂-fibers and residual water. The residual water in the fibers was extruded by pressure and collected completely. ²²⁶Ra in MnO₂-fibers and residual waters were determined by the radon emanation method aforementioned. These results (table 1) showed that ²²⁶Ra activities in residual water were considerably lower than those in MnO₂-fibers. Among the five samples, ²²⁶Ra activities in the residual water of three samples were below the detection limit. In the other samples, ²²⁶Ra activities provided the evidence showing that ²²⁶Ra contents in the extruded waters were negligible and the method used for ²²⁶Ra determination was reliable.

Sample layer/m	$\frac{^{226}\text{Ra}}{V} \times 10^2/\text{Bq} \cdot \text{m}^{-3 \text{ a})}$	Residual water	$\frac{\frac{226}{\mathrm{Ra}}}{V} \times 10^{2}/\mathrm{Bq} \cdot \mathrm{m}^{-3 \mathrm{a}}$
25	3.04 ± 0.06	L-C34-25 m	$B.D^{b)}$
50	4.49 ± 0.07	L-C34-50 m	0.09 ± 0.07
100	3.99 ± 0.06	L-C34-100 m	B.D ^{b)}
200	4.65 ± 0.07	L-C34-200 m	0.07 ± 0.07
500	2.44 ± 0.06	L-C34-500 m	B.D ^{b)}

Table 1 ²²⁶Ra activities in MnO₂-fibers and residual water from Station C34

a) Assuming that the water volume flowing through MnO₂-fiber is V (in dm³); b) B.D, below the detection limit.

1.4 Analysis of other parameters

Temperature and salinity were determined *in-situ* via MARK III C/WOCE-CTD. Nutrients were analyzed according to the national standards in China "Marine Investigation Criteria · Measurement of Chemical Parameters in Seawater" (GB1276-4-01). NO_3^- data were calibrated for zinc reduction efficiency^[9].

2 Results and discussion

2.1 ²²⁶Ra activities in the surface seawater

 226 Ra activities in 33 surface seawater samples collected from the Arctic Ocean and the Bering Sea are listed in table 2. The results showed that 226 Ra activities in the surface waters ranged from 0.28 to 1.56 Bq/m³ with an average of 0.76 Bq/m³ in the Arctic Ocean, and from 0.25 to 1.26 Bq/m³ with an average of 0.71 Bq/m³ in the Bering Sea. Our results were consistent with previous

Sea area	Station	Longitude	Latitude	Depth/m	Salinity (%)	226 Ra/Bq · m ⁻³
	C6	170°0.6'W	69°59.9'N	50	25.633	0.28 ± 0.02
	C8	174°59.5 ′ W	70°0.7'N	60	31.147	0.34 ± 0.02
	C10	173°54.3′W	71°0.1′N	35	26.669	0.43 ± 0.03
	C11	172°29.6′W	71°1.3′N	38	30.500	0.28 ± 0.03
	C12	170°2.1'W	70°40.1'N	30	30.520	1.17 ± 0.04
A	C13	167°10.2 ′ W	70°28.8'N	50	31.021	1.31 ± 0.04
Arctic	C25	168°52.6′W	71°42.0'N	50	29.099	0.28 ± 0.03
Ocean	C27	168°38.2 ′ W	72°29.6'N	54	30.291	1.56 ± 0.04
(C30	160°0.6'W	71°15.3′N	44	29.850	0.52 ± 0.03
	C33	158°56.5 ′ W	72°22.5'N	50	28.395	0.87 ± 0.04
	C34	157°55.9 ′ W	73°25.2'N	2700	26.409	1.19 ± 0.04
	C39	161°55.3′W	75°16.1'N	2080	28.295	0.94 ± 0.03
	BJ11	169°49.5 ′ W	67°18.7'N	47	31.852	0.91 ± 0.04
	BJ9	179°45.9′E	62°11.2′N	-	31.708	0.60 ± 0.02
	BJ10	166°59.6′W	64°21.8'N	_	26.987	0.78 ± 0.03
	B1-12	178°14.2 ′ W	60°39.7'N	165	32.168	0.33 ± 0.03
	B1-9	179°25.8′W	60°15.2'N	840	32.665	0.29 ± 0.03
	B1-6	179°19.7′E	59°39.6′N	3200	32.730	0.42 ± 0.03
	B1-4	177°55.4′E	58°59.4'N	3720	32.848	0.26 ± 0.04
	B1-3	176°9.4′E	58°0.2'N	3780	32.968	0.46 ± 0.02
	B1-2	174°30.5 ′ E	56°59.8′N	3650	33.011	1.22 ± 0.04
	B1-1	173°21.1 ′ E	55°59.8'N	3850	32.970	0.60 ± 0.03
Bering	B2-1	176°0.6′E	56°0.1′N	3860	32.821	1.26 ± 0.04
Sea	B3-1	177°59.8′E	56°0.0'N	3860	32.972	0.76 ± 0.03
	B5-1	179°59.1 ′ W	55°59.9'N	3830	32.768	0.76 ± 0.04
	B6-1	178°1.2'W	56°2.8'N	3780	32.932	0.25 ± 0.02
	B6-3	175°59.9 ′ W	55°59.5'N	3720	32.866	0.72 ± 0.04
	B5-3	177°39.3 ′ W	57°31.3'N	3680	32.906	1.09 ± 0.05
	B4-1	180°0.0 ′ W	56°59.9'N	3810	32.832	0.62 ± 0.03
	B2-2	177°14.0 ′ E	56°59.7′N	3830	32.996	0.85 ± 0.04
	B5-4	176°39.5 ′ W	58°3.6′N	3370	32.688	1.19 ± 0.05
	B5-7	176°10.8 ′ W	58°26.0'N	2440	32.628	0.95 ± 0.05
	B5-10	175°53.9 ′ W	58°39.9'N	139	32.409	0.83 ± 0.04

data from the Arctic Ocean and the Bering Sea, but obviously lower than those from open oceans in middle and lower latitudes (table 3).

The low ²²⁶Ra activities in the study area may be caused by sea ice meltwater. Sea ice is composed of pure-water ice, seawater and air packed in ice when the temperature is higher than -8.2 . When the temperature is equal to -8.2 , $Na_2SO_4 \cdot 10H_2O$ can be crystallized and separated out. NaCl \cdot 2H₂O can be separated out after the temperature decreases to -22.9 . Other salts will be congealed entirely at -55 . During the formation of sea ice, the fractionation of ²²⁶Ra may occur and result in the low ²²⁶Ra activities in sea ice and the high activities in saline waters. Besides, ²²⁶Ra activities in the snow covering on the sea ice could be low. Both effects will

Sea area	226 Ra/Bq $\cdot m^{-3 a)}$	Reference	
Pacific Ocean	Average ≈1.07	[10]	
Atlantic Ocean	Average ≈1.23	[11]	
Indian Ocean	1.30-2.28(16)		
Western Pacific Ocean	1.30-1.40(2)	[12]	
Indonesian Sea	1.38-2.77(6)		
Nansha Sea Area	1.57(1)		
Amazon Estuary	0.83-3.62(20)	[13]	
Dead Sea (100—300 m)	1153—1927(14)	[14]	
East China Sea, Yellow Sea	0.95-3.33(31)	[15]	
Seto Inland Sea	0.22-3.07(82)	[16]	
Orinoco Estuary	0.48-2.99(22)	[17]	
Jiulongjiang Estuary	2.10-5.92(11)	[18]	
Bering Sea(KH-75-4)	²²⁶ Ra (salinity(%))		
Station 8(57°00'N,171 °00'E)	$1.50 \pm 0.13(32.83)$		
Station 9(57°00'N,174 °54'E)	$1.73 \pm 0.13(33.17)$	[6]	
Station 10(57°01'N, 179°03'E)	$1.00 \pm 0.07(32.95)$		
Station 11(57°05'N, 177°05'W)	$1.03 \pm 0.10(32.53)$		
Arctic Ocean(CESAR Station)	1.78 (depth: 25 m)	[5]	
Eastern Bering Sea	~0.33—~1.67	[7]	
Arctic Ocean	0.28-1.56	this study	
Bering Sea	0.25-1.26		

result in low ²²⁶Ra contents in sea ice meltwater. Unfortunately, there has not been any data reported about ²²⁶Ra activities in the sea ice at present.

Table 2 ²²⁶Do activition in the surface water from different and and

a) Except for particular annotation, the values in the brackets are the sample numbers.

2.2 Spatial distribution of ²²⁶Ra and other parameters in the surface seawater——the composition of the water masses in the Arctic Ocean

Spatial distributions of temperature, salinity and nutrients (NO_3^- , PO_4^{3-} , SiO_3^{2-}) in the surface water are shown in fig. 2(a)—(e). Surface temperature in the study area decreased northward. The distribution of surface salinity was relatively complicated. In the Bering Sea, surface salinity increased westward. In the Arctic Ocean, salinity in the surface water decreased in the offshore direction and reached the minimum (25.633‰) at Station C6 in the middle region. This distribution pattern indicated that the freshwater composition in the Arctic Ocean increased in the offshore direction and converged at the internal region. The spatial distribution of nutrients (nitrate, phosphate and silicate) in the surface water had a similar feature: nutrient concentrations decreased northward in the Bering Sea and increased northward and formed a high concentration area in the middle region of Arctic Ocean.

Spatial distribution of ²²⁶Ra activities in the surface water of the Arctic Ocean and the Bering Sea is shown in fig. 3. In the Bering Sea, ²²⁶Ra in the surface water decreased northward, probably as a result of the exchange between the ²²⁶Ra-deficient sea ice meltwater and the ²²⁶Ra-rich Pacific water. In the Arctic Ocean, ²²⁶Ra in the surface water increased northward and eastward,

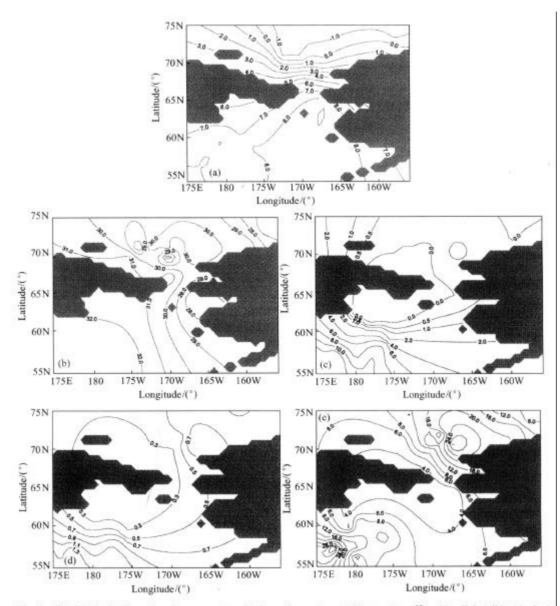


Fig. 2. Spatial distributions of various parameters in the surface water. (a) Temperature (°C); (b) salinity (%c); (c) nitrate $(\mu mol/m^3)$; (d) phosphate $(\mu mol/m^3)$; (e) silicate $(\mu mol/m^3)$.

and formed a high concentration area in the internal region. Since river water carried not only "original" dissolved ²²⁶Ra, but also the ²²⁶Ra released from suspended particles by desorption, it had a high ²²⁶Ra characteristic. The distribution pattern of ²²⁶Ra thus indicated that the river component increased in the offshore direction and converged at the internal ocean. Additionally, since continental shelves covered about 1/3 of the total surface area of the Arctic Ocean and most water inputs had a long residence time on the shelves before entering the internal ocean, the hydrogra

phy of the Arctic Ocean was strongly influenced by these extensive shelf areas^[19]. Thus, river water and shelf water with high ²²⁶Ra converged at the internal Arctic Ocean. The distribution of ²²⁶Ra in the study area showed a unique pattern, contrary to the general pattern in which ²²⁶Ra decreased in the offshore direction. The information about water mass composition obtained from the spatial distribution of ²²⁶Ra was coincident with that of the results from salinity and ²H, ¹⁸O tracers^[20].

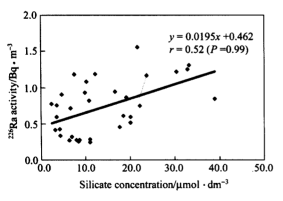
70N ()) 65N 66N 60N 55N 175E 180 175W 170W 165W 160W 1

Although the spatial distributions of

²²⁶Ra activities and salinities indicated the similar information on the water mass composition in the Arctic Ocean, there was no statistically significant linear relationship between ²²⁶Ra activity and salinity. This was mainly due to the interstitial water input from bottom sediment. In addition, sea ice meltwater with low salinity may also contribute to the insignificant linear relationship between ²²⁶Ra and salinity.

2.3 Correlation between ²²⁶Ra activity and dissolved silicate concentration

Both ²²⁶Ra activities and dissolved silicate concentrations decreased northward in the Bering



Sea and increased northward in the Arctic Ocean and formed a high concentration region in the internal Arctic Ocean (fig. 3 and fig. 2(e)). Linear regression showed that ²²⁶Ra activities had a statistically significant positive correlation with dissolved silicate concentrations (fig. 4, correlation coefficient was 0.52 at the 99% confidence level). Glover and Reeburgh^[7] also explored the relationship between ²²⁶Ra activities and dissolved silicate for all samples collected from the southeastern Bering Sea, and found a

significant positive linear relationship between 226 Ra and silica at the 95% confidence level (correlation coefficient is 0.66).

2.4 Profiles of ²²⁶Ra in the Canadian Basin——sources and mechanism for the upper halocline water

Vertical samples in the upper water between 0 and 500 m were collected by hanging MnO_2 -fibers at certain depths for several hours at Station C34 and Station C39 in the Canadian

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Basin. Because of no data about the water volume flowing through MnO₂-fibers, ²²⁶Ra activities cannot be calculated quantitatively. However, taking into account the fact that the MnO2-fibers drifted continually with the ship during sampling and the drift velocity far exceeded the horizontal flow rate of waters, it is reasonable to assume that the flow rate at each depth was similar and the water volume through MnO₂-fibers is consistent. Thus, ²²⁶Ra relative activities can be calculated. The results are listed in table 4.

Station	Depth/m	Temperature/	Salinity (%)	$\frac{^{226}\text{Ra}}{V} \times 10^2/\text{Bq} \cdot \text{m}^{-3 \text{ a}}$
C34 (157°55.9'W, 73°25.2'N)	25	-1.100	29.616	3.04 ± 0.06
	50	-1.016	31.769	4.49 ± 0.07
	100	-1.510	32.522	3.99 ± 0.06
	150	-1.509	33.040	4.08 ± 0.07
	200	-0.974	33.975	4.65 ± 0.07
	500	0.470	34.823	2.44 ± 0.06
C39 (161°55.3'W, 75°16.1'N)	25	-0.944	29.382	0.57 ± 0.03
	50	-0.732	31.642	1.14 ± 0.04
	100	-1.396	32.590	1.56 ± 0.04
	200	b)	b)	1.93 ± 0.05
	300	0.410	34.694	1.43 ± 0.05
	500	0.569	34.823	2.02 ± 0.05

a) Assuming that the water volume flowing through MnO₂-fiber is V/dm³; b) no data.

The profiles of ²²⁶Ra and other parameters at the two stations (C34 and C39) in the Canadian Basin are shown in figs. 5 and 6. The results showed that the water columns had the characteristic of "the upper halocline water" with a constant temperature of at most the freezing point and a salinity that increases with depth (figs. 5(a) and 6(a)). Nitrate, phosphate and silicate had maximal concentrations at depths of 150–200 m (figs. 5(b) and 6(b)). Conservative parameters NO and PO had maximums at 200 m and minimums at 300 m, while NO/PO ratio had a minimum at 200 m. The ratio increased distinctly from this depth and kept almost constant below 300 m (figs. 5(c) and 6(c)). Corresponding with these hydrochemical characteristics, ²²⁶Ra showed a concentration maximum at 200 m (figs. 5(d) and 6(d)). The maximum could be attributed to the inputs of the Pacific water or/and the bottom shelf water with a high ²²⁶Ra concentration. The results based on 226 Ra were supported by those from 2 H, 18 O tracers $^{[21]}$.

Aagaard et al.^[21] proposed two possibilities for the maintenance of the upper halocline water: the first is that it was maintained by the lateral advection of cold, saline water formed by sea ice formation on the continental shelves; the second is that it was formed by the upwelling and cooling of the Atlantic layer during winter. Moore and Smith^[5] argued that this water mass had its origin in the Bering Sea, but was strongly modified during its residence on the shelf, especially by

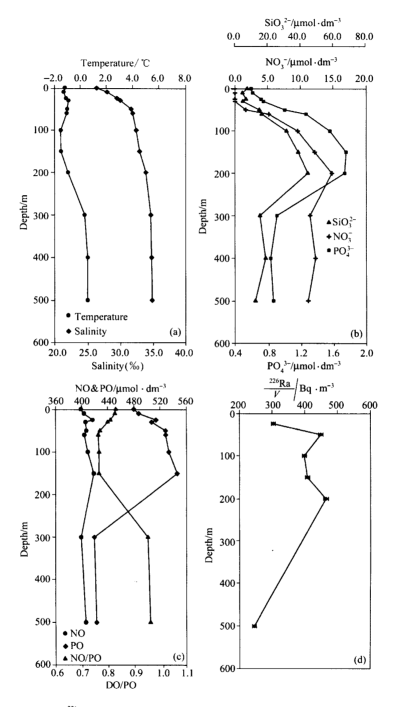


Fig. 5. The vertical profiles of ²²⁶Ra and other parameters at Station C34. (a) Temperature (\bullet) and salinity (\bullet); (b) nitrate (+), phosphate (\blacksquare) and silicate (\blacktriangle) (µmol/m³); (c) NO (\bullet), PO (\bullet) (µmol/m³) and NO/PO (\blacktriangle); (d) ²²⁶Ra specific activity (Bq/m³).

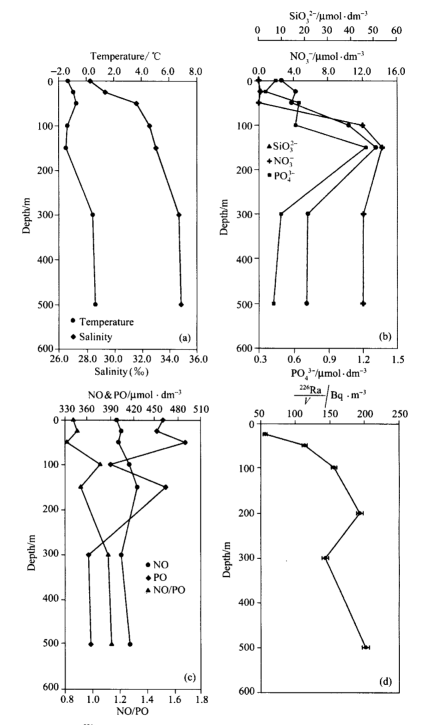


Fig. 6. The vertical profiles of 226 Ra and other parameters at Station C39. (a) Temperature (°C) and salinity (‰); (b) nitrat phosphate and silicate (µmol/m³); (c) NO, PO (µmol/m³) and NO/PO; (d) 226 Ra specific activity (Bq/m³). Designators are the same as in fig. 5.

the halocline water. They analyzed the distribution of NO/PO data in the Arctic Ocean and over the shelves and concluded that the upper halocline water must be derived from the Pacific Ocean surface water, which was modified on the Chukchi/Bering shelf. Bauch et al.^[23] developed a 4-component model to determine the contribution of river water, meltwater, Atlantic Ocean and Bering Strait inflows, using d^{18} O, salinity and silicate as tracers. Applied to the data from CESAR Ice Station, their model reproduces the high proportion of Bering Strait Inflow in the upper halocline water. The results presented above were in agreement with our present understanding of the sources and mechanism of the upper halocline water in the Canadian Basin, primarily based on ²²⁶Ra profiles.

Note that ²²⁶Ra also had a maximum at 500 m at Station C39 (fig. 6(d)). Similar phenomenon was also observed in the profile of ²²⁸Ra. This phenomenon has not been explained reasonably and is worthy of further research.

3 Conclusions

(1) 226 Ra activities in the surface water ranged from 0.28 Bq/m³ to 1.56 Bq/m³ with an average of 0.76 Bq/m³ in the Arctic Ocean, and from 0.25 Bq/m³ to 1.26 Bq/m³ with an average of 0.71 Bq/m³ in the Bering Sea. These values were obviously lower than those from open oceans in middle and lower latitudes, indicating that the study area may be partly influenced by sea ice meltwater.

(2) In the Bering Sea, ²²⁶Ra in the surface water decreased northward, probably as a result of the exchange between the ²²⁶Ra-deficient sea ice meltwater and the ²²⁶Ra-rich Pacific water. In the Arctic Ocean, surface ²²⁶Ra increased northward and eastward. This spatial distribution of ²²⁶Ra reflected the variation of the ²²⁶Ra-enriched river component in the water mass of the Arctic Ocean.

(3) Profiles of ²²⁶Ra in the Canadian Basin showed a concentration maximum at depth of 200 m. The maximum could be attributed to the inputs of the Pacific water or/and the bottom shelf water with high ²²⁶Ra concentration.

In brief, ²²⁶Ra distribution in the Arctic Ocean and Bering Sea has a unique pattern compared to that of other sea areas. These features can be used to study the water inputs in the Arctic Ocean, to evaluate the sources and mechanism of the upper halocline water in the internal Arctic Basin, and to quantify the exchange between the Arctic Ocean and the northern Pacific Ocean. Furthermore, it will be helpful to understanding the regulation of Chinese and global climates.

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