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Insight into the measurement of dissolved ²²⁷Ac in seawater using radium delayed coincidence counter

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ABSTRACT

Due to the low abundance of 227 Ac in seawater, the analysis of this radionuclide requires the use of high-sensitivity, low-background instruments and the collection of large volume samples. A promising technique relies on the pre-concentration of 227 Ac in seawater using cartridges impregnated with manganese oxide (Mn-cartridges) that are mounted on in situ pumps, and its measurement on a Radium Delayed Coincidence Counter (RaDeCC), usually used to analyze short-lived radium isotopes. In this work, we present an evaluation of this technique, including 1) the study of the performance of the RaDeCC measurements for 227 Ac fixed on Mn-cartridges (backgrounds, detector efficiency, repeatability), and 2) the determination of the efficiency of seawater 227 Ac extraction of the Mn-cartridges and its reproducibility for the first time, by using Mn-cartridges placed in series. Overall, we found a Mn-cartridge extraction efficiency of 47 \pm 129 % (1 SD). Repeatability experiments allowed us to estimate the uncertainties of the entire measurement of 19 % (1 SD). Finally, in the aim to validate the method, the 227 Ac activities duetermined in several samples using Mn-fibers (assuming 100 % yield of 227 Ac extraction) and 2) to the 231 Pa activities determined at the same stations during the GEOVIDE cruise (GEOTRACES GA01), 231 Pa being the parent nuclide of 227 Ac (Deng et al., 201 B). Only few studies 227 Ac and 231 Pa have been published so far due to the difficulty to analyze these two nuclides. First, the 227 Ac activities determined using Mn-cartridges agree well with the 227 Ac activities determined using Mn-cartridges agree well with the 227 Ac activities determined using Mn-cartridge extraction efficiency.

1. Introduction

Actinium-227 (227 Ac; $T_{1/2} = 21.8$ y) is produced by the radioactive decay of its parent isotope protactinium-231 (231 Pa; $T_{1/2} = 32,760$ y), itself produced by the decay of uranium-235 (235 U; $T_{1/2} = 7.04 \times 10^8 \text{ y}$). Oceanic 235U concentrations are mostly constant through space and time due to a long residence time (~0.5 Ma; Ku et al., 1977), which leads to a uniform production rate of ²³¹Pa. Once produced, ²³¹Pa is adsorbed onto particles, and is transported to the seafloor, where it accumulates in deep-sea sediments (Anderson et al., 1983). The decay of ²³¹Pa to ²²⁷Ac in the sediments is followed by partial release of ²²⁷Ac due to its higher solubility (Anderson et al., 1983; Nozaki, 1993, 1984) thus leading to an ²²⁷Ac flux to the overlying water column (Nozaki, 1993, 1984; Nozaki et al., 1990). Being mostly soluble, ²²⁷Ac is redistributed in the ocean through water mass transport and mixing. Assuming steady state, ²²⁷Ac produced in the water column is expected to be in secular equilibrium with ²³¹Pa. ²²⁷Ac that diffuses out of deep-sea sediments leads to the observation of excess ²²⁷Ac activities (²²⁷Ac_{ex}) in deep waters, usually below 2000 m (Geibert et al., 2002; Nozaki, 1984; Nozaki et al., 1990). Hydrothermal vents have also been identified as an additional source of ²²⁷Ac to the deep ocean (Kipp et al., 2015). Due to its deep-sea source, ²²⁷Ac has been used to quantify vertical mixing and as a tracer of deep ocean circulation at basin scales and/or timescales of ca. 100 y (Geibert et al., 2002; Koch-Larrouy et al., 2015; Nozaki, 1984). Because vertical advection can also bring ²²⁷Ac towards the sea surface, ²²⁷Ac has also been used to estimate upwelling rates (Geibert et al., 2002)

²²⁷Ac concentrations are especially low within the ocean, the total oceanic inventory being only 37 mol or 8.4 kg (Geibert et al., 2008). As a result, relatively few studies have been reported to date (Dulaiova et al., 2012; Geibert et al., 2008, 2002; Geibert and Vöge, 2008; Kipp et al., 2015; Koch-Larrouy et al., 2015; Nozaki, 1993, 1984; Nozaki et al., 1998, 1990; Shaw and Moore, 2002). The analysis of ²²⁷Ac thus requires handling large volumes (typically > 200 L), from which ²²⁷Ac is extracted, either through Fe or Mn hydroxide coprecipitation (Geibert and Vöge, 2008) or through sorption onto media impregnated with

MnO₂ (Shaw and Moore, 2002). Shaw and Moore (2002) reported a method for the analysis of ²²⁷Ac in seawater based on the extraction of ²²⁷Ac from seawater on acrylic fibers impregnated with manganese oxide (Mn-fibers), followed by its analysis using a Radium Delayed Coincidence Counter (RaDeCC; Moore and Arnold, 1996). However, passing hundreds of liters of seawater through Mn-fibers is not easy is time consuming and prevents from building vertical profiles of ²²⁷Ac with a high resolution. To reduce the sample volume requirement, ²²⁷Ac can be determined using alpha spectrometry that requires only 20-80 L of seawater at typical ocean activities (Geibert and Vöge, 2008). However, this approach requires lengthy steps of chemical purification and separation of ²²⁷Ac prior to analysis.

Alternatively. ²²⁷Ac can be extracted from large volumes of seawater using Mn-cartridges mounted on in situ pumping systems (Henderson et al., 2013). This allows for extraction of ²²⁷Ac from seawater at depth without having to bring the water onboard for extraction onto Mn fibers. In this work, we evaluate the performance of this extraction method that is followed by the use of RaDeCC to quantify ²²⁷Ac activities. Samples used herein were collected along the GEOVIDE section in the North Atlantic (GEOTRACES GA01). We discuss the limit of quantification and the uncertainties associated with the 227Ac activities determined using RaDeCC. The repeatability (see Joint Committee for Guides in Metrology (2008) for metrology vocabulary definitions) is assessed by repeated analysis. We quantify the yield of ²²⁷Ac extraction onto the Mn-cartridges using two Mn-cartridges mounted in series. The ²²⁷Ac activities thus obtained can be compared with the ²²⁷Ac activities determined in several samples using Mn-fibers (following Shaw and Moore, 2002). Finally, in the aim to further validate the method, we report two seawater ²²⁷Ac vertical profiles in the North Atlantic Ocean and compare them to the vertical profiles of ²³¹Pa activity from the same stations and with historical data. Above 2000 m, where ²²⁷Ac is expected to be in secular equilibrium with ²³¹Pa, such a comparison provides independent validation and assesses the trueness (see Joint Committee for Guides in Metrology, 2008 for metrology vocabulary definitions) of the procedure as a whole.

2. Materials and methods

2.1. Sample collection

The samples were collected in the framework of the GEOVIDE section (GEOTRACES GA01; PIs: Géraldine Sarthou, LEMAR, France and Pascale Lherminier, LOPS, France) that was conducted in the North Atlantic Ocean between Lisbon, Portugal and St John's, Canada (15 May - 30 June 2014). Along the section, nine stations were sampled for ²²⁷Ac. ²²⁷Ac was extracted from seawater using two Mn-cartridges placed in series on in situ pumps. As a comparison, several discrete, large volume samples were also collected using Niskin bottles, and passed by gravity through Mn-fibers. Here, we report on the validation of the method used to determine dissolved ²²⁷Ac activities. The ²²⁷Ac activities determined for the entire GEOVIDE section will be shown in a separate paper.

2.1.1. Sample preparation and extraction of ²²⁷Ac using Mn-cartridges

Several cartridges designed to be mounted on in situ pumps were prepared using a protocol slightly modified from Henderson et al. (2013). Acrylic fiber cartridges of 25.4 cm (10 in.) long with 5 μm porosity were cut to obtain a final size of 77 mm \pm 0.4 mm, which is slightly smaller than the cartridges prepared by Henderson et al. (2013). The cartridges were soaked in a milli-Q water bath for 48 h. The cartridges were then impregnated with MnO2 by soaking in a saturated KMnO4 bath at room temperature for 48 h. In addition, circulation of a saturated KMnO4 solution through the cartridges served to improve the MnO2 impregnation within the cartridge itself with the aim of increasing the cartridge extraction efficiencies for $^{227} Ac$ (and other radionuclides). Mn-cartridges were rinsed after the final step with

radium-free milli-Q water and then vacuum packed in a sachet.

At each station, large volume in situ pumps (ISP) (Challenger and McLane) were deployed at 6 to 13 depths for 3 to 4 h, which resulted in filtration volumes of 418 to 1565 L at flow rates of 3-6 L min⁻¹. Seawater first passed through Supor (Pall, 0.8 µm pore size) or QMA (Sartorius, 1 um pore size) membranes (to collect suspended particles). then through the Mn-cartridges to preconcentrate ²²⁷Ac. For the deep samples, two Mn-cartridges (A Mn-cartridge and B Mn-cartridge) were placed in series in order to provide information on the yield of ²²⁷Ac fixation (see section 3.2.). For shallow samples, single cartridges were mounted on the in-situ pumps. Following collection, each Mn-cartridge was rinsed with Ra-free milli-O water and slightly dried using compressed air. Similar to Mn-fibers (Sun and Torgersen, 1998), the moisture of the Mn-cartridges may affect the emanation efficiency of radon that is actually the radionuclide quantified using the RaDeCC systems (see 2.2). According to Sun and Torgersen (1998), the optimal range for the 220 Rn emanation from Mn-fibers is 0.3–1.0 g_{water}/g_{fiber} . Therefore, the Mn-cartridges were weighted to control their water content and were kept to a moisture range similar to that of the Mnfibers. The water content of the Mn-cartridges mostly ranged from 0.3 to 0.7 gwater/gcartridge.

2.1.2. Extraction of ²²⁷Ac using Mn-fibers

At two stations (stations 32 and 38), six discrete seawater samples were collected using Niskin bottles at the same depth as the ISPs in order to validate the $^{227}\mathrm{Ac}$ activities determined using Mn-cartridges. Seawater samples, which ranged from 120 to 541 L, were weighted using a Chatillon KPB-052-T Portable Bench Platform Scale. The samples were then stored in large plastic barrels and were passed by gravity through 20 g of acrylic fibers impregnated with MnO₂ (Mn-fiber) that quantitatively adsorb radium isotopes but also $^{227}\mathrm{Ac}$ when the flow rate is below 1 L min $^{-1}$; in such conditions, Mn-fibers have been shown to extract 99 \pm 1% of both Ra and Ac (Moore and Reid, 1973). $^{227}\mathrm{Ac}$ analysis using RaDeCC.

Following Shaw and Moore (2002), we used Radium Delayed Coincidence Counters (RaDeCC, Scientific Computer Instruments, USA), which were primarily designed to determine ²²³Ra and ²²⁴Ra in seawater samples (Giffin et al., 1963; Moore, 2008; Moore and Arnold, 1996). Dissolved ²²⁷Ac activities were determined by measuring the ²¹⁹Rn activities that are in secular equilibrium with ²²³Ra and ²²⁷Ac. Mn-cartridges (and Mn-fibers) were analyzed at least three months after sampling to allow ²¹⁹Rn, ²²³Ra, and ²²⁷Ac to equilibrate. The partially dried Mn-cartridges (and Mn-fibers) were placed in plastic cartridge (or fiber) holders in a closed helium circulation loop. Cartridges holders used for the analysis are smaller than the ones used for sampling. These small cartridges holders are designed for 4" cartridges but we use 3" cartridges raise with PVC pieces and are 169 mm high with a diameter of 133 mm. Helium was allowed to circulate over the Mn-cartridges (and Mn-fibers) and carried the ²¹⁹Rn to the scintillation cell coated with ZnS, where alpha particles produced by the ²¹⁹Rn decay into ²¹⁵Po are detected. A delayed coincidence system developed by Giffin et al. (1963) and then adapted by Moore and Arnold (1996) allowed us to discriminate the signal associated with 219Rn from that of other Rn isotopes (220Rn and 222Rn) that are not associated with 227Ac. Correction for chance coincidence counts was performed (Garcia-Solsona et al., 2008; Moore and Arnold, 1996).

The measurement of the Mn-cartridges using RaDeCC was carried out for at least 13 h and up to 24 h with the aim of increasing counting statistics. This could be achieved because of the very low background levels of the RaDeCC system (see below). Note that the background may increase due to decay products (accumulation of Rn and its daughter) remaining in the counting cells. In addition, Mn–cartridges lost up to 4.36 g of water during long counting sessions with a mean of 34 mg of water per hour (this study), which may also result in the accumulation of moisture within the detector. No effect of the moisture (between 30% and 70%) has been observed along counting or between repeated

measurements with different moisture content. In order to avert these latter issues, the scintillation cells were flushed for at least 4 h between samples to flush the system of these residual isotopes and to eventually dry the cell. We chose to conduct the analyses on only two RaDeCC systems in order to reduce the variability that may result from the use of multiple counting systems. Mn cartridges are known to extract radium isotopes, thorium isotopes but also ²³¹Pa. Therefore, a simple decay correction for ²²⁷Ac activities (as usually done) is not appropriate since ²²⁷Ac will also grow into equilibrium with its parent ²³¹Pa (due to the 3 to 4 year lag between sampling and analysis). Despite the fact that ²³¹Pa activities have been measured on separate samples along the section (Deng et al., 2018), the extraction efficiency of ²³¹Pa on Mn-cartridges is not known. Therefore, the amount of ²³¹Pa on Mn-cartridges are not known, which prevent from precisely quantifying the ingrowth of ²²⁷Ac. Assuming a ²³¹Pa extraction efficiency of 80% on Mn-cartridges, the 227 Ac activities would need to be lowered by 10% (\pm 13%, 1 SD). If no 231Pa was extracted by the Mn-cartridges, 227Ac activities would need to be increased by up to 7%. Therefore, in the absence of precise information on the 231 Pa activity on the Mn cartridges and because the correction is small, the ²²⁷Ac activities were not corrected for decay/ ingrowth that occurred between sampling and analysis. ²²⁷Ac activities were then corrected for Mn-cartridge extraction efficiency and normalized to the seawater volume. ²²⁷Ac activities are reported in dpm m ³. Despite the 100% extraction efficiency of the Mn-fibers, the ²²⁷Ac activities on the Mn-fiber samples were very low, due to the reduce amount of water that was collected using Niskin bottles. Each sample was counting several times, and the counts were summed up to decrease the uncertainties. The system was flushed for 5 min in between each counting and water was added if necessary to the Mn-fiber to maintain constant moisture.

3. Results and discussion

3.1. RaDeCC performances

3.1.1. Background measurements

The RaDeCC system displays very low backgrounds. We conducted several long background counts (up to 73 h; Table 1) using two sets of Mn-coated acrylic cartridges similar to those used during the GEOVIDE cruise. These long background analyses provided a precise estimation of the background for the two detectors used in this study (Table 1). For both Mn-cartridges and Mn-fibers, the chance coincidence corrected backgrounds on channel 219 (219Rn) were 0.004 and 0.002 cpm (counts per minute) for detectors #1 and #2 respectively. The backgrounds reported here were of the same order of magnitude as those previously reported in the literature for Mn-fibers (Garcia-Solsona et al., 2008; Giffin et al., 1963; Moore and Arnold, 1996). The limit of detection (LOD) and limit of quantification (LOQ) were calculated as 3 and 10 times the standard deviation of these backgrounds, respectively. As far as Mn-cartridges are concerned, LOD were 0.007 and 0.006 cpm for detectors #1 and #2, respectively, while LOQ were 0.014 cpm for both detectors (Table 1).

3.1.2. Detector efficiency

The two RaDeCC systems used in this study were calibrated using Mn-cartridge standards, i.e. Mn-cartridges containing a known amount of ²²⁷Ac. Mn-cartridges were prepared using radium free seawater spiked with an ²²⁷Ac solution and passed through the Mn-cartridges via gravity flow. When preparing the standard Mn-cartridge, the effluent was passed several times onto the Mn-cartridge to ensure quantitative adsorption of the ²²⁷Ac spike. Mn-cartridge standards were measured using small cartridge holders in a similar manner as for samples. This standard was measured by multiple laboratories over 13 months and it has proven to be stable over time (no decrease in activity as observed by Scholten et al. (2010)). The RaDeCC systems were also calibrated for Mn-fibers using a fiber standard. A decline with time has been observed

Table 1Backgrounds for ²¹⁹Rn of detectors #1 and #2 for both Mn-cartridges and Mn-fibers. The limit of detection (LOD) and limit of quantification (LOQ) were calculated as 3 and 10 times the standard deviation of these backgrounds, respectively.

		Detector #1	Detector #2
	Run time (min)	²¹⁹ Rn (cpm)	²¹⁹ Rn (cpm)
Mn-cartridges	959	0.004	0.003
_	1027	0.002	0.002
	1220	0.005	0.007
	1346	0.004	0.001
	2743	0.002	0.001
	3852	0.003	0.002
	3864	0.005	0.001
	3938	0.003	0.000
	4061	0.004	0.001
	4393	0.006	0.001
	Mean	0.004	0.002
	SD (1σ)	0.001	0.001
	LD	0.007	0.006
	LQ	0.014	0.014
Mn-fibers	377	0.003	0.003
	422	0.002	0.002
	464	0.000	0.000
	876	0.000	0.001
	924	0.001	0.000
	Mean	0.001	0.001
	SD (1σ)	0.001	0.001
	LD	0.004	0.004
	LQ	0.011	0.011

on the old Mn-fiber standard previously used to calibrated for ²¹⁹Rn (i.e., Mn fiber impregnated with a standard solution of ²²⁷Ac and prepared at Alfred Wegener Institute (AWI, Germany) and described in van Beek et al. (2010)). Therefore, the efficiency of the systems for fibers was determined from the efficiency of the ²²⁰Rn channel using a Mn-fiber impregnated with a known amount of ²²⁸Th following Scholten et al. (2010). The efficiency for ²²⁷Ac was then determined following Moore and Cai (2013). The standards were measured under controlled conditions (room temperature and humidity) over a 9-month period.

The detector efficiencies are reported in Table 2 for both Mn-cartridges and Mn-fibers. For Mn-cartridges, the detector efficiencies for the 219 channel were 35% \pm 2% and 34% \pm 2%, for detectors #1 and #2 respectively. For Mn-fiber, the efficiencies show good agreement with those for Mn-cartridges (i.e., 36% \pm 3% and 34% \pm 4% for

Table 2
Detector efficiencies for ²²⁷Ac (for both Mn-cartridges and Mn-fibers).

	Detection effici	ency	Detection effici	ency
	(Mn-cartridge)		(Mn-fiber)	
	Detector #1	Detector #2	Detector #1	Detector #2
	38%	37%	37%	28%
	36%	34%	37%	37%
	33%	34%	36%	35%
	34%	34%	37%	32%
	38%	35%	35%	29%
	34%	33%	40%	32%
	35%	32%	37%	37%
	35%	31%	37%	38%
	34%	33%	29%	38%
	35%	33%	37%	34%
Mean	35%	34%	36%	34%
SD (1σ)	2%	2%	3%	4%
n	10	10	10	10

Table 3

227 Ac activities determined on Mn-cartridges A for each analysis of several repeated samples are reported together with the average of ²²⁷Ac activities from these repeated analyses. The internal precision corresponds to the propagated standard error (SE) and relative standard error (RSE) determined following Garcia-Solsona et al. (2008). The external precision represents the calculated standard deviation (SD) and relative standard deviation (RSD) determined from repeated measurement of the same samples are reported.

										Internal p	recision	External 1	precision
Station	Longitude (°E)	Latitude (°N)	Depth (m)	Volume (L)	Runtime (min)	²¹⁹ Rn (counts)	²¹⁹ Rn (cpm)	²²⁷ Ac on Mn- cartridges (dpm)	Mean (dpm)	SE* (1σ) (dpm)	RSE* (1σ) %	SD (1σ) (dpm)	RSD (1σ) %
1	-10.036	40.333	2500	1565	1432	115	0.080	0.09	0.15	0.02	23	0.07	48
					1025	147	0.143	0.22		0.04	15		
					1002	80	0.080	0.12		0.02	20		
13	-13.888	41.383	1500	1341.5	1013	38	0.038	0.05	0.04	0.02	34	0.01	18
					1055	40	0.038	0.04		0.02	44		
			4000	1482	998	71	0.071	0.12	0.10	0.02	20	0.03	31
					1015	42	0.041	0.06		0.02	31		
					916	43	0.047	0.10		0.05	52		
			5280	647	1216	125	0.103	0.16	0.19	0.03	18	0.04	19
					949	114	0.120	0.23		0.03	15		
					1017	99	0.097	0.18		0.09	48		
21	-19.672	46.544	2700	1417	1222	66	0.054	0.06	0.05	0.02	37	0.00	8
					903	37	0.041	0.06		0.02	34		
					1228	49	0.040	0.05		0.01	31		
			3500	1482	911	74	0.081	0.14	0.14	0.03	20	0.00	1
					1007	84	0.083	0.14		0.03	19		
26	-22.603	50.278	2023	1423	962	37	0.038	0.06	0.06	0.02	26	0.00	3
					1070	50	0.047	0.06		0.02	29		
			2843	680	1281	40	0.031	0.06	0.06	0.02	25	0.00	8
					886	31	0.035	0.05		0.02	34		-
					1143	43	0.038				25		
			3563	1351	1053	58	0.055		0.09		24	0.01	7
			3303	1551	955	51	0.053	88 0.06 0.02 55 0.08 0.09 0.02 63 0.09 0.02 64 0.09 0.05 55 0.09 0.12 0.02 01 0.14 0.03	25	0.01	,		
	200			989	53	0.054				54			
			3868	581	1130	62	0.055		0.12		23	0.04	32
		3868	3606	361	1027	93	0.033		0.12		18	0.04	32
32	- 26 710	55 506	800	1378	1027	43	0.040	0.02	0.03	0.03	68	0.00	14
)2	-26.710 55.506 800 2267 2854	300	1376	1035	29	0.040	0.02	0.03	0.02	49	0.00	14	
		2267	1334	976	27	0.028	0.03	0.03	0.01	56	0.01	28	
		2207	1334	967	24	0.025	0.03	0.03	0.02	36	0.01	20	
			2054	1965	1041	40			0.05			0.01	20
			2854	1265	1041	35	0.038 0.07 0.05 0.02 27 0.01 0.034 0.05 0.02 31 0.019 0.04 0.01 38 0.031 0.04 0.02 42 0.047 0.03 0.04 0.02 61 0.00	0.01	28				
					967	35 18							
	01.067	E0.040	1016	1166	1089	34			0.04			0.00	12
38	-31.267	58.843	1316	1166	1002	47	0.047	0.03	0.04		61	0.00	12
					1003	51	0.051	0.04		0.02	46		
	00.054	F0.600	0560	1.40	1179	34	0.029	0.04	0.11	0.02	35	0.01	
14	-38.954	59.623	2560	149	1036	71	0.069	0.10	0.11	0.03	24	0.01	11
	46.000	50.000	400	1000	1292	101	0.078	0.12	0.06	0.03	20	0.00	0.6
54	-46.083	59.068	400	1068	1014	68	0.067	0.07	0.06	0.02	31	0.02	26
					962	55	0.057	0.05		0.02	44		_
			1716	1169	1027	34	0.033	0.05	0.05	0.02	33	0.00	3
					990	40	0.040	0.05		0.02	39		
59	-48.093	55.841	460	952	1032	44	0.043	0.06	0.05	0.02	53	0.02	40
					1074	49	0.046	0.03		0.02	54		
			2463	418	1147	58	0.051	0.07	0.08	0.02	24	0.01	8
					962	45	0.047	0.08		0.02	26		
			3616	738	1393	48	0.034	0.05	0.04	0.02	30	0.01	19
			1038	23	0.022	0.03		0.01	40				
					995	30	0.030	0.04		0.01	35		
								Weighted average			34%	0.02	19%
								Standard Deviatio	n		13%		13%

detectors #1 and #2, respectively). These efficiencies for the Mn-cartridges and Mn-fibers are in good agreement with those reported by Henderson et al., (2013) (30% to 38% and 28% to 47%, respectively).

3.1.3. Precision

The internal precision associated with the RaDeCC measurement is usually estimated through uncertainty propagation calculations as reported by (Garcia-Solsona et al., 2008). Here, the repeatability (i.e. external precision) was assessed by measuring a total of twenty (out of seventy-three) individual Mn-cartridge samples (A Mn-cartridges only) two, three, or four times (eleven duplicates, eight triplicates and one quadruplicate, yielding to a total of fifty measurements; Table 3).

Standard deviations of these replicates are < 0.09 dpm (1 SD) while the relative standard deviations (1 RSD) range between 1% and 48% (Table 3), with a weighted mean value of 19% (weight of 2 for duplicates, 3 for triplicates and 4 for quadruplicates). The Fig. 1 shows that there is no clear relationship between the RSD and the 227 Ac activities of the Mn-cartridges notably that the repeatability does not improve with increasing 227 Ac activity. As a comparison, the uncertainty calculated following Garcia-Solsona et al., (2008) is $34 \pm 13\%$ (1 SD; n = 50; Table 3, for A Mn-cartridges). Forty-one measurements out of the fifty ones constituting this repeatability study have an internal precision estimated by the counting statistic propagation method reported by Garcia-Solsona et al. (2008) larger than the mean

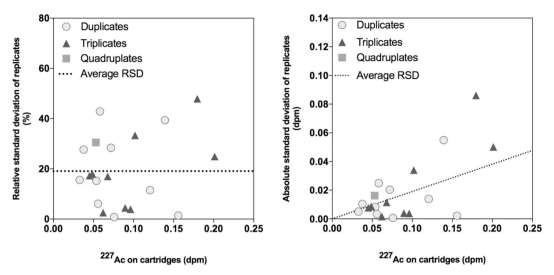


Fig. 1. Repeatability determined from replicate analyses of GEOVIDE Mn-cartridge samples expressed as the relative (left) and absolute (right) standard deviation of each replicate, versus the ²²⁷Ac activities on Mn-cartridges. The weighted average (weight of 2 for duplicates, 3 for triplicates and 4 for quadruplicates) value of relative standard deviations calculated for each individual replicate is shown as dotted lines on both plots.

repeatability of 19 \pm 13%. The RSD is lower (11%) for activities on Mn-cartridges from 0.06 to 0.08 cpm while for activities from 0.02 to 0.06 cpm and from 0.08 to 0.20 cpm, the RSD are 20% and 21%, respectively. The internal precision estimated by the counting statistic propagation decrease with increasing activities on the Mn-cartridges. This suggests that these uncertainty propagation calculations (Garcia-Solsona et al., 2008) may overestimate the real measurement uncertainties. This may be caused by the fact that these propagation calculations assume several hypotheses like the independence of the different variables that may not be verified. In any case, precision estimation from replicate analyses is always preferable to the error propagation calculations. Therefore, the average external precision reported above (19%, 1 SD, n=20) is considered as the best choice for the measurement uncertainty for all the 227 Ac activities reported in this study.

3.2. Mn-cartridge extraction efficiencies

3.2.1. Mn-cartridges in series

The extraction efficiency for dissolved ²²⁷Ac onto the Mn-cartridges was determined for each sample by measuring the ²²⁷Ac activity on two Mn-cartridges mounted in series on the in situ pumps as was done in the past for other radionuclides extracted from seawater samples in a similar manner (Baskaran et al., 1993; Livingston and Cochran, 1987; Mann and Casso, 1984; van der Loeff and Moore, 1999). The extraction efficiency is determined as follows:

$$E = 1 - \frac{^{227}Ac_{\text{Cart B}}}{^{227}Ac_{\text{Cart A}}}$$
 (1)

Where E is the dissolved 227 Ac extraction efficiency of the Mn-cartridges, 227 Ac_{cartA} and 227 Ac_{cartB} are the 227 Ac activities on the first and second Mn-cartridges placed in series, respectively. With this approach, E is assumed to be equal for both Mn-cartridges.

Among the thirty-six paired cartridges analyzed in this study, the ²²⁷Ac activities on B Mn-cartridges were found to be below the limit of quantification for fifteen samples, which prevented us from estimating the extraction efficiencies for these samples (Fig. 2). One could argue that this could be due to higher than average extraction efficiencies on A cartridges combined with average or below average extraction efficiencies on B cartridges. In such a case, excluding these samples would bias our mean extraction efficiency estimation. However, the activities measured on A Mn-cartridge of most of these fifteen samples (13 out of 15) were also low. Therefore, this rules out the above mention caveat,

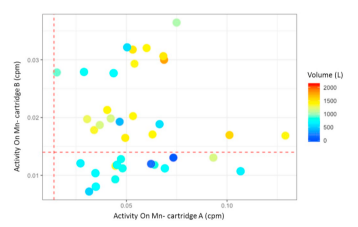


Fig. 2. ²²⁷Ac activities on Mn-cartridges A vs B in. The colour scale represents the volume (V) in L. The red dashed line represents the limit of quantification (LOQ). Below this line the samples were not use to estimate the extraction efficiency. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

and shows that these low activities on cartridges B rather reflect the combination of relatively small seawater volumes (shown in Fig. 2) and low seawater ²²⁷Ac activities. Only two samples display large ²²⁷Ac activities on A Mn-cartridges, suggesting that the ²²⁷Ac extraction efficiency onto the Mn-cartridges is higher on the A Mn-cartridges compared with the B Mn-cartridges. Excluding these data from the mean extraction efficiency estimation does introduce a negative bias. Unfortunately we have no choice but to exclude them as B cartridge measurement is unreliable because below detection limit. However, this involves only 2 data points compared to 21 used for our estimation, thus, the negative bias is likely negligible.

Among the remaining twenty-one samples, nine samples were chosen arbitrarily for repeatability estimation (Mn-cartridge B count rates above LOQ; Table 4). Repeated analyses were thus conducted in these samples, by analyzing both Mn-cartridges A and B, in order to better constrain the extraction efficiency. The obtained extraction efficiency for all Mn-cartridges ranged from 31% to 78%, with a mean value of 47 \pm 12% (1 SD, n=21). This mean value was obtained calculating a weighted average by multiplying by one samples analyzed once, by two samples analyzed twice, by three samples analyzed three times and by four samples analyzed four times. Other estimations done

 Table 4

 Extraction efficiencies calculated with Mn-cartridges A and B.

Column C	Column C	arion	oration ponferrate patricular volunte requires	Daning	, popu	T AOIM	day ar	reaces cantinge in				carriage p	9						
(4) (4) <th>(4) (4) (4) (4) (4) (40)</th> <th></th> <th></th> <th></th> <th></th> <th></th> <th></th> <th>Runtime</th> <th>²¹⁹Rn</th> <th></th> <th>²²⁷Ac on Mn- cartridge A</th> <th></th> <th></th> <th></th> <th>²²⁷Ac on Mn- cartridge B</th> <th>Extraction efficiency</th> <th>Replicates Mean</th> <th>Replicates SD (10)</th> <th>Replicates RSD (10)</th>	(4) (4) (4) (4) (4) (40)							Runtime	²¹⁹ Rn		²²⁷ Ac on Mn- cartridge A				²²⁷ Ac on Mn- cartridge B	Extraction efficiency	Replicates Mean	Replicates SD (10)	Replicates RSD (10)
40.00 150.00 </th <th>44.323 200 156 169 156 169<</th> <th></th> <th>(°E)</th> <th>(°N)</th> <th>(m)</th> <th>(T)</th> <th></th> <th>(min)</th> <th>(counts)</th> <th></th> <th>(mdp)</th> <th>(min)</th> <th></th> <th></th> <th>(mdp)</th> <th>%</th> <th>%</th> <th>%</th> <th>%</th>	44.323 200 156 169 156 169<		(°E)	(°N)	(m)	(T)		(min)	(counts)		(mdp)	(min)			(mdp)	%	%	%	%
40.333 55.00 156.6 a 1423 15.00 10.	40.333 35.00 156. 1 142 114 0.08 0.089 1142 1 1		-10.036	40.333	2000			800	55	0.07	0.114	800	24		0.069	40			
4. 5.24 1. 10.22 1.47 0.14 0.124 1.025 1.03	4. 381 4. 482 4. 47 0.44 0.224 1.025 1.0 0.018 0.023 0.038 0.033 0.038 0.033 0.034 0.033 0.034 0.032 0.034 0.032 0.034 0.033 0.034 0.033 0.034		-10.036	40.333	2500			1432	115	0.08	0.089	1432	22		0.024	73	78	7	6
44.383 3600 1382 4 1002 1002 1002 74 44.383 3600 1382 4 1002 1002 1002 74 74 44.383 3600 1382 4 999 70 1012 999 37 0000 1002 <t< td=""><td>4.5.3 3.6.0 1.86 0.103 1.00 1.0 <th< td=""><td></td><td></td><td></td><td></td><td></td><td>Р</td><td>1025</td><td></td><td></td><td>0.224</td><td>1025</td><td>18</td><td></td><td>0.031</td><td>98</td><td></td><td></td><td></td></th<></td></t<>	4.5.3 3.6.0 1.86 0.103 1.00 1.0 <th< td=""><td></td><td></td><td></td><td></td><td></td><td>Р</td><td>1025</td><td></td><td></td><td>0.224</td><td>1025</td><td>18</td><td></td><td>0.031</td><td>98</td><td></td><td></td><td></td></th<>						Р	1025			0.224	1025	18		0.031	98			
40.233 38.90 13.83 1008 13.0 0.13 0.03 10.0	44.381 300.0 183.5 100.8 11.0 10.0 10.1 0.02 40. 41.382 300.0 1842 3 100.8 11.0 0.01 0.02 40. 41.381 300.0 1842 3 100.8 11.0 0.07 0.07 40. 30. 40. 41.381 400.0 182 3 100.0 0.02 0.00 27. 40.						С	1002			0.123	1002	18		0.032	74			
41.383 400 1346 99 61 0.02 0.	41.383 300. 1356 999 70 0.052 0.073 0.073 0.073 0.073 0.073 0.073 0.074 0.073 0.074 0.073 0.074		-10.036					1008	_		980.0	1008	17		0.052	40			
41.383 46.39 47.9 40.3 91.8 40.0 41.3 89 77 0.07 0.123 99.8 34 0.03 0.04 41.3 89 77 0.07 0.123 99.8 34 0.03 0.04 27 41.3 89 77 0.04 0.03 0.04 0.03 0.04 27 41.3 8 9	41.383 46.9 47.9 49.8 34.0 40.0 43.9 35.9 41.383 40.0 41.383 40.0 41.383 40.0 41.383 40.0 41.383 48.0 40.0 41.383 40.0 41.383 48.0 40.0 41.0 40.0<		-13.888					666		90.0	0.122	666	32		0.071	42			
4.3.82 4.6.6 6.4 6.0 6.0 6.0 6.0 7. 4.3.82 4.6.6 6.4 6.0 6.0 6.0 6.0 6.0 7. 4.3.82 4.6.6 6.4 6.0 6.0 1.0 6.0 6.0 6.0 6.0 6.0 6.0 6.0 4.0 4.0 4.0 4.0 4.0 6.0 6.0 6.0 6.0 6.0 6.0 6.0 4.0 4.0 4.0 6.0 <t< td=""><td> 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1,</td><td></td><td>-13.888</td><td></td><td></td><td></td><td></td><td>866</td><td></td><td>0.07</td><td>0.123</td><td>866</td><td>34</td><td></td><td>0.070</td><td>43</td><td>35</td><td>8</td><td>23</td></t<>	1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1,		-13.888					866		0.07	0.123	866	34		0.070	43	35	8	23
41.383 48.50 48.70 49.90 49.0	41.383 4836 644 6 91 6 101 6 1						р	1015		0.04	0.063	1015	30		0.046	27			
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41.383 580 67 a 1216 124 1216	41.383 580 67 a 1216 1216 0.136 0.1146 1216 0.126 0.136 0.1146 0.12 0.134 0.114 0.12 0.233 0.00		-13.888		4850			626			0.094	626	26		0.048	49			
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52.27.8 38.6 5.4 0.02 98.9 2.2 0.022 0.046 5.0 52.27.8 3.86 5.81 a 1927 6.2 0.05 0.034 10.20 6.7 55.50.6 4.13.5 a 110.7 6.2 0.05 0.144 11.6 15 BDL BDL n.d. 55.50.6 2.267 133.5 a 9.76 2.7 0.03 0.02 0.04 0.03 0.03 0.03 0.03 0.03 0.04 0.03 0.04 0.03 0.04 0.03 0.04 0.03 0.04 0.03 0.04 0.03 0.04 0.03 0.04 0.03 0.04 0.03 0.04 0.03 0.04 0.04 0.03 0.04 0.04 0.03 0.04 0.04 0.03 0.04 0.04 0.03 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 </td <td>50.2.5.0 3.64 5.6 0.042 0.046 5.0 0.046 5.0 50.2.7.8 3.86 5.81 a 1130 6.0 0.041 0.042 0.046 5.0 50.2.7.8 4.12 b 1027 9.0 0.144 1146 15 0.04 0.05 0.04 0.04 0.05 0.04 0.04 0.05 0.04 0.04 0.05 0.04 0.04 0.05 0.04 0.04 0.05 0.04 0.04 0.05 0.04 0.04 0.05 0.04 0.04 0.05 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04</td> <td></td> <td></td> <td></td> <td></td> <td></td> <td>Р</td> <td>955</td> <td></td> <td>0.02</td> <td>0.088</td> <td>955</td> <td>31</td> <td></td> <td>0.059</td> <td>34</td> <td></td> <td></td> <td></td>	50.2.5.0 3.64 5.6 0.042 0.046 5.0 0.046 5.0 50.2.7.8 3.86 5.81 a 1130 6.0 0.041 0.042 0.046 5.0 50.2.7.8 4.12 b 1027 9.0 0.144 1146 15 0.04 0.05 0.04 0.04 0.05 0.04 0.04 0.05 0.04 0.04 0.05 0.04 0.04 0.05 0.04 0.04 0.05 0.04 0.04 0.05 0.04 0.04 0.05 0.04 0.04 0.05 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04						Р	955		0.02	0.088	955	31		0.059	34			
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5.5.06 1.87 1.89 1.027 9.9 0.144 1146 1.8 BDL BDL m.d. 55.506 2.267 1.813.5 a 961 7.2 0.078 0.078 4.3 9.0 9.0 1.0 0.0	5.5.02 1.02 0.03 0.144 1146 15 BDI BDI n.d. 55.506 1.815 9.61 7.2 0.07 0.134 961 1.05 0.016 0.017 0.017 0.016 0.017 0.017 0.017 0.017 0.017 0.017 0.017 0.017 0.017 0.018 0.017 0.018 0.017 0.018 0.017 0.018 0.017 0.018 0.017 0.018 0.017 0.018 0.018 0.019 0.018 0.019 0.011 0.019 0.011		-22.603	50.278	3868		В	1130		0.05	0.091	1130	16		0:030	29			
50,278 413 1031.5 961 72 0.13 961 35 0.036 0.078 43 55.506 1481 143.5 961 32 0.046 0.045 0.046 0.047 0.046 0.047 0.046 0.047 0.046 0.047 0.046 0.047 0.046 0.047 0.046 0.047 0.046 0.047 0.046 0.047 0.046 0.047 0.046 0.047 0.046 0.047 0.046 0.047 0.046 0.047 0.049 0.047 0.049 0.047 0.049 0.049 0.049 0.049 0.044 <td< td=""><td>50.278 41.26 1031.5 961 35 0.036 0.078 43 55.506 1481 142.5 961 35 0.036 0.078 43 55.506 1481 143.5 961 70 0.049 971 16 0.016 0.020 26 37 8 55.506 133.5 a 967 24 0.02 0.041 1279 29 0.023 0.02 37 8 55.506 126 a 1089 34 0.03 0.043 1041 17 0.016 0.02 0.041 1079 37 8 38 39 39 38 39 39 39 36 36 30 30 30 30 30 38 39</td><td></td><td></td><td></td><td></td><td></td><td></td><td>1027</td><td></td><td>0.09</td><td>0.144</td><td>1146</td><td>15</td><td></td><td>BDL</td><td>n.d.</td><td></td><td></td><td></td></td<>	50.278 41.26 1031.5 961 35 0.036 0.078 43 55.506 1481 142.5 961 35 0.036 0.078 43 55.506 1481 143.5 961 70 0.049 971 16 0.016 0.020 26 37 8 55.506 133.5 a 967 24 0.02 0.041 1279 29 0.023 0.02 37 8 55.506 126 a 1089 34 0.03 0.043 1041 17 0.016 0.02 0.041 1079 37 8 38 39 39 38 39 39 39 36 36 30 30 30 30 30 38 39							1027		0.09	0.144	1146	15		BDL	n.d.			
55.506 1481 1442.5 971 48 0.05 0.049 971 16 0.016 0.021 70 8 55.506 2854 133.5 a 976 27 0.027 0.047 17 16 0.026 37 36 37 8 55.506 2854 1265 a 1041 40 0.04 0.07 1041 17 0.016 0.023 37 38 39 55.506 3050 3050 0.03 0.041 1089 12 0.04	55.506 1481 1442.5 971 48 0.05 0.069 971 16 0.016 0.021 0.02 37 8 55.506 2854 1265 a 1041 40 0.04 0.027 1040 0.023 <td></td> <td>-22.603</td> <td></td> <td>4126</td> <td></td> <td>ı.</td> <td>961</td> <td></td> <td>0.02</td> <td>0.136</td> <td>961</td> <td>35</td> <td></td> <td>0.078</td> <td>43</td> <td></td> <td></td> <td></td>		-22.603		4126		ı.	961		0.02	0.136	961	35		0.078	43			
55.506 2854 133.5 a 976 27 0.03 0.027 976 14 0.014 0.020 37 8 55.506 2854 1265 a 1041 40 0.04 0.043 0.023 0.026 37 38 39 55.506 2854 1265 a 1048 35 0.03 0.041 108 24 0.023 0.025 38 39 55.506 370 370 37 0.04 1089 34 0.03 0.041 1089 12 BDL BDL DL n.d 39 39 39 39 39 39 39 39 30 30 30 30 30 39 39 39 39 30 30 30 39 39 39 39 39 39 39 39 39 39 39 30 30 30 30 30 30 30 30 <td>55.506 2854 1333.5 a 976 27 0.04 0.024 0.04 0.020 26 31 8 55.506 2854 1265 a 1041 40 0.04 0.057 17 0.016 0.023 0.026 37 39 39 55.506 2854 1265 a 1041 40 0.04 0.053 1038 24 0.023 0.047 111 89 34 0.03 0.053 1049 126 0.03 0.049 106 0.023 0.040 0.040 0.053 0.041 BDL n.d 99 99 99 90</td> <td></td> <td>-26.710</td> <td></td> <td>1481</td> <td>1442.</td> <td>τċ</td> <td>971</td> <td></td> <td>0.02</td> <td>690.0</td> <td>971</td> <td>16</td> <td></td> <td>0.021</td> <td>20</td> <td></td> <td></td> <td></td>	55.506 2854 1333.5 a 976 27 0.04 0.024 0.04 0.020 26 31 8 55.506 2854 1265 a 1041 40 0.04 0.057 17 0.016 0.023 0.026 37 39 39 55.506 2854 1265 a 1041 40 0.04 0.053 1038 24 0.023 0.047 111 89 34 0.03 0.053 1049 126 0.03 0.049 106 0.023 0.040 0.040 0.053 0.041 BDL n.d 99 99 99 90		-26.710		1481	1442.	τċ	971		0.02	690.0	971	16		0.021	20			
55.506 2854 1265 a 1041 40 0.04 0.073 0.023 0.026 37 89 39 55.506 2854 1265 a 1041 40 0.04 0.073 0.043 0.023 0.049 0.07 0.043 0.043 0.043 0.043 0.043 0.043 0.043 0.043 0.043 0.043 0.043 0.043 0.043 0.043 0.043 0.044 0.044 0.044 0.044 0.043 0.043 0.044	55.506 2854 1265 a 967 24 0.02 0.041 17 0.016 0.025		-26.710		2267	1333.		926		0.03	0.027	926	14		0.020	26	31	œ	24
55.506 2854 1265 a 1041 40 0.067 0.067 1041 17 0.016 0.023 0.047 11 38 39 55.506 305 306 30.63 0.053 0.043 1038 24 0.023 0.047 11 39 39 39 39 30	55.506 2854 1265 a 1041 40 0.047 1041 17 0.016 0.023 65 38 39 55.506 386 1088 35 0.03 0.041 1098 14 0.023 0.047 11 89 34 0.03 0.041 1098 13 8DL BDL BDL BDL n.d 89 13 8DL BDL BDL n.d 89 10 99 11 10 99 11 10 11 10							296		0.02	0.041	1279	29		0.026	37			
5.5.50 3.6 3.6 0.053 0.053 10.40 10.88 24 0.023 0.047 11 55.506 3.6 3.6 1.0 9.95 1.3 0.041 10.89 1.2 BDL n.d 55.506 3.10 7.1 8.94 4.3 0.05 0.042 10.15 1.8 BDL n.d 55.506 3.10 7.1 8.94 4.3 0.05 0.042 10.15 1.3 BDL n.d 58.843 6.45 7.0 1.005 0.050 0.042 10.15 1.0<	55.506 370 970<		-26.710		2854			1041		0.04	0.067	1041	17		0.023	65	38	39	101
S.5.56 30.50 89 34 0.03 0.041 1089 12 BDL BDL n.d. 5.5.50 30.50 89 73 0.07 0.009 995 13 BDL BDL n.d. 5.5.50 30.50 99 73 0.07 0.002 892 10 n.d. 5.8.843 8.0 101 48 0.05 0.042 101 BDL n.d. 5.8.843 8.0 108 0.05 0.042 1120 9 BDL n.d. 5.8.843 108 697 1120 39 0.05 0.045 93 12 8DL n.d. 58.843 1136 116 2 0.045 100 0.045 10 0.01 0.045 10 0.01 0.01 0.01 0.01 0.01 0.01 0.01 0.01 0.01 0.02 0.02 0.02 0.02 0.02 0.02 0.02 0.02 <t< td=""><td>55.50 395 34 0.03 0.041 1089 12 BDL BDL m.d. 55.506 3170 395 73 0.07 0.009 995 13 BDL BDL m.d. 55.506 3170 71 894 43 0.05 0.042 995 13 BDL BDL m.d. 58.843 640 1389 0.05 0.042 0.05 0.07 0.025 0.07</td><td></td><td></td><td></td><td></td><td></td><td>Р</td><td>1038</td><td></td><td>0.03</td><td>0.053</td><td>1038</td><td>24</td><td></td><td>0.047</td><td>11</td><td></td><td></td><td></td></t<>	55.50 395 34 0.03 0.041 1089 12 BDL BDL m.d. 55.506 3170 395 73 0.07 0.009 995 13 BDL BDL m.d. 55.506 3170 71 894 43 0.05 0.042 995 13 BDL BDL m.d. 58.843 640 1389 0.05 0.042 0.05 0.07 0.025 0.07						Р	1038		0.03	0.053	1038	24		0.047	11			
55.506 395 73 0.07 0.009 995 13 BDL BDL BDL BDL 55.506 3170 771 894 43 0.05 0.062 995 10 BDL BDL BDL n.d 58.434 54.3 7.0 1015 48 0.05 0.042 1015 10 BDL BDL BDL n.d 58.843 1389 103 0.05 0.05 0.05 1120 9. BDL BDL n.d 58.843 1386 106 0.03 0.045 9.93 12 BDL BDL n.d 58.843 1316 116 a 1002 47 0.03 0.045 1002 14 BDL BDL n.d 58.843 1316 116 a 1002 17 1004 21 0.01 n.d 0.02 18 58.843 1316 10 0.03 0.042 <t< td=""><td>55.506 3950 349 995 73 0.07 0.099 995 13 BDL BDL m.d. 55.506 3170 771 1015 43 0.05 0.062 892 10 BDL BDL m.d. 58.843 3470 1379 93 0.06 0.050 937 1120 99 m.d. 58.843 1087 697 1120 39 0.06 0.050 937 12 BDL BDL m.d. 58.843 1087 697 1120 39 0.025 1120 99 10 m.d. 58.843 1166 a 1002 1045 993 12 BDL BDL m.d. 58.843 136 106 0.045 1004 21 0.021 m.d. 0.04 10 10 m.d. 0.04 10 10 m.d. 0.02 0.02 0.021 0.021 0.021 0.021 0.02<</td><td></td><td></td><td></td><td></td><td></td><td>С</td><td>1089</td><td></td><td></td><td>0.041</td><td>1089</td><td>12</td><td>BDL</td><td>BDL</td><td>n.d.</td><td></td><td></td><td></td></t<>	55.506 3950 349 995 73 0.07 0.099 995 13 BDL BDL m.d. 55.506 3170 771 1015 43 0.05 0.062 892 10 BDL BDL m.d. 58.843 3470 1379 93 0.06 0.050 937 1120 99 m.d. 58.843 1087 697 1120 39 0.06 0.050 937 12 BDL BDL m.d. 58.843 1087 697 1120 39 0.025 1120 99 10 m.d. 58.843 1166 a 1002 1045 993 12 BDL BDL m.d. 58.843 136 106 0.045 1004 21 0.021 m.d. 0.04 10 10 m.d. 0.04 10 10 m.d. 0.02 0.02 0.021 0.021 0.021 0.021 0.02<						С	1089			0.041	1089	12	BDL	BDL	n.d.			
55.506 3170 771 894 43 0.05 0.062 0.042 101 BDL BDL BDL BDL 58.843 643 770 1015 48 0.05 0.042 1015 13 BDL BDL m.d. 58.843 1389 1080 0.05 0.05 0.05 1120 99 10.0	55.506 3170 771 894 43 0.05 0.062 892 10 BDL BDL n.d. 58.843 643 770 1015 48 0.05 0.042 1015 13 BDL m.d. 58.843 840 1389 0.06 0.050 937 16 0.017 0.021 59 58.843 1087 697 1120 39 0.025 112 9DL BDL n.d. 58.843 136 1160 9 0.025 10.03 12 BDL BDL n.d. 58.843 136 1160 9 1002 12 BDL BDL n.d. 58.843 136 1063 0.042 1004 1002 1021 n.d. 32 18 58.843 136 106 0.042 1004 1002 1021 n.d. 32 18 59.623 134 0.01 0.025 0.023<		-26.710		3050			995			600.0	995	13	BDL	BDL	n.d.			
58.843 643 770 1015 48 0.05 0.042 1015 11 BDL B	58.843 643 770 1015 48 0.05 0.042 1015 13 BDL BDL BDL m.d. 58.843 840 1389 937 59 0.05 0.050 937 16 0.017 0.021 59 58.843 186 1120 39 0.05 0.045 99 12 BDL m.d. 58.843 1316 1160 a 0.045 0.045 190 1120 9 BDL m.d. 58.843 1316 1160 a 0.045 0.045 104 BDL m.d. 32 18 58.843 1316 1160 a 0.042 1042 104 BDL BDL m.d. 32 18 59.623 274 1374 15 0.02 0.043 104 27 0.028 0.025 18 104 104 27 0.028 0.025 0.024 104 104 107		-26.710	55.506	3170			894		0.02	0.062	892	10	BDL	BDL	n.d.			
58.843 840 1389 937 59 0.05 0.050 937 16 0.017 0.021 0.021 0.037 0.03 0.052 0.03 0.04 0.03 0.03 0.04 0.03 <t< td=""><td>58.843 840 1389 937 59 0.06 0.050 937 16 0.017 0.021 59 58.843 1087 697 1120 39 0.025 1120 9 BDL m.d. n.d. 58.843 1235 656 47 0.03 0.045 993 12 BDL m.d. n.d. 58.843 1316 1160 a 1002 0.042 1002 14 BDL BDL m.d. 32 18 58.843 1316 1160 a 1002 0.042 1002 1021 0.021 0.031 n.d. 32 18 59.623 134 0.03 0.043 1179 29 0.025 0.027 37 37 36 59.623 2880 838 1016 65 0.02 0.023 0.02 0.02 0.02 0.02 0.02 0.02 0.02 0.02 0.02 0.02 <td< td=""><td></td><td>-31.267</td><td></td><td>643</td><td>770</td><td></td><td>1015</td><td></td><td>0.05</td><td>0.042</td><td>1015</td><td>13</td><td>BDL</td><td>BDL</td><td>n.d.</td><td></td><td></td><td></td></td<></td></t<>	58.843 840 1389 937 59 0.06 0.050 937 16 0.017 0.021 59 58.843 1087 697 1120 39 0.025 1120 9 BDL m.d. n.d. 58.843 1235 656 47 0.03 0.045 993 12 BDL m.d. n.d. 58.843 1316 1160 a 1002 0.042 1002 14 BDL BDL m.d. 32 18 58.843 1316 1160 a 1002 0.042 1002 1021 0.021 0.031 n.d. 32 18 59.623 134 0.03 0.043 1179 29 0.025 0.027 37 37 36 59.623 2880 838 1016 65 0.02 0.023 0.02 0.02 0.02 0.02 0.02 0.02 0.02 0.02 0.02 0.02 <td< td=""><td></td><td>-31.267</td><td></td><td>643</td><td>770</td><td></td><td>1015</td><td></td><td>0.05</td><td>0.042</td><td>1015</td><td>13</td><td>BDL</td><td>BDL</td><td>n.d.</td><td></td><td></td><td></td></td<>		-31.267		643	770		1015		0.05	0.042	1015	13	BDL	BDL	n.d.			
58.843 1087 697 1120 99 1120 9 BDL BDL BDL m.d. 58.843 1235 656 993 27 0.03 0.045 993 12 BDL BDL m.d. 58.843 1316 1166 a 1002 47 0.05 0.034 1002 14 BDL BDL m.d. 32 18 58.843 1316 106 a 1002 0.042 1004 21 0.021 0.031 0.04 10	58.843 1087 697 1120 39 0.03 0.055 1120 9 BDL BDL m.d. 58.843 1235 656 993 27 0.03 0.045 993 12 BDL m.d. 58.843 1316 1166 a 1002 47 0.042 1004 21 0.021 0.03 18 59.623 136 0.01 0.042 1004 21 0.021 0.03 1026 18 59.623 136 91 1 1 2 0.023 971 2 0.028 0.027 0.028<		-31.267		840	1389		937		90.0	0.050	937	16		0.021	59			
58.843 1235 656 993 27 0.045 0.045 993 12 BDL B	58.843 1235 656 993 27 0.045 993 12 BDL BDL m.d. 32 18 58.843 1316 1166 a 1002 47 0.05 0.034 1002 14 BDL BDL m.d. 32 18 59.623 1316 1166 a 1003 0.042 1004 21 0.021 0.031 26 37 59.623 1374 3 0.02 0.023 0.023 0.023 0.029 0.028 0.027 0.028 0.027 0.028 0.024 0.028 0.028 0.028 0.028 0.028 0.029 0.028 0.029 0.028 0.029 0.028 0.029 0.028		-31.267	58.843	1087	269		1120			0.025	1120	6	BDL	BDL	n.d.			
58.843 1316 1166 a 1002 47 0.03 0.034 1002 14 BDL BDL n.d. 32 18 59.623 1316 1003 51 0.03 0.042 1004 21 0.021 0.031 26 18 59.623 1374 179 34 0.03 0.043 1179 29 0.025 0.029 37 79 </td <td>58.843 1316 1166 a 1002 47 0.05 0.034 1002 14 BDL B</td> <td></td> <td>-31.267</td> <td></td> <td>1235</td> <td></td> <td></td> <td>666</td> <td>27</td> <td>0.03</td> <td>0.045</td> <td>666</td> <td>12</td> <td>BDL</td> <td>BDL</td> <td>n.d.</td> <td></td> <td></td> <td></td>	58.843 1316 1166 a 1002 47 0.05 0.034 1002 14 BDL B		-31.267		1235			666	27	0.03	0.045	666	12	BDL	BDL	n.d.			
59.623 176 107 51 0.05 0.042 1004 21 0.021 0.031 59.623 175 0.1 179 34 0.03 0.043 1179 29 0.025 0.027 59.623 2746 1374 15 0.02 0.033 971 27 0.028 0.005 59.063 1716 169 a 1016 65 0.064 1016 12 BDL BDL 59.068 1716 1169 a 1027 34 0.03 0.052 1026 0.034 59.068 2056 205 0.050 0.019 0.050 0.018 BDL BDL 59.068 205 0.05 0.050 0.051 111 BDL BDL	59.623 1136 901 1003 51 0.05 0.042 1004 21 0.021 0.031 26 59.623 1136 901 1179 34 0.03 0.043 1179 29 0.025 0.027 37 59.623 1374 137 15 0.02 0.023 971 27 0.028 79 59.623 2746 1374 1016 65 0.06 0.064 1016 12 BDL BDL n.d. 59.068 1716 1169 a 1027 34 0.052 1025 0.026 0.034 34 59.068 206 52 0.052 102 0.050 0.01 0.04 0.050 0.01 0.02 0.03 0.03 0.01 0.050 0.03 0.01 0.04 0.04 0.050 0.01 0.04 0.04 0.050 0.01 0.01 0.01 0.04 0.050 0.01 0.01		-31.267		1316			1002			0.034	1002	14		BDL	n.d.	32	18	58
59.623 1136 901 97 15 0.03 0.043 1179 29 0.025 0.027 59.623 1136 901 15 0.02 0.023 971 27 0.028 0.005 59.623 2746 1374 134 0.0 0.06 0.043 191 27 0.028 0.018 59.068 1716 116 a 1075 34 0.05 0.052 11 BDL BDL 59.068 2060 25 0.05 0.052 0.052 0.034 BDL BDL 59.068 2069 26 0.061 0.050 0.016 BDL BDL 59.068 2060 23 0.03 0.051 111 8 BDL BDL	59.623 1136 901 971 15 0.03 0.043 1179 29 0.025 0.025 0.027 37 59.623 1374 901 971 15 0.02 0.023 971 27 0.028 0.005 79 59.623 2746 1374 938 50 0.03 0.038 938 19 0.020 0.018 54 59.623 2880 838 1016 65 0.064 1016 12 BDL BDL n.d 59.068 1716 1169 a 1027 34 0.052 1026 0.036 0.034 34 59.068 2060 523 0.03 0.051 1112 8 BDL BDL n.d						Р	1003		0.02	0.042	1004	21		0.031	26			
59,623 1136 901 971 15 0.02 0.023 971 27 0.028 0.005 59,623 2746 1374 938 50 0.05 0.038 938 19 0.020 0.018 59,063 1716 1169 a 1016 65 0.064 0.064 1016 12 BDL BDL 59,068 1716 1169 a 1027 34 0.03 0.052 102 0.034 59,068 2060 23 1112 35 0.03 0.051 1112 8 BDL BDL	59.623 1136 901 971 15 0.02 0.023 971 27 0.028 0.005 79 79 59.623 2746 1374 938 50 0.03 0.038 938 19 0.020 0.018 54 59.623 2880 838 1016 65 0.06 0.064 1016 12 BDL BDL n.d 59.068 1716 1169 a 1027 34 0.052 11 BDL BDL n.d 59.068 2060 523 0.03 0.051 111 8 BDL n.d n.d						၁	1179		0.03	0.043	1179	29		0.027	37			
59,623 2746 1374 938 50 0.03 0.038 938 19 0.020 0.018 59,623 2880 838 1016 65 0.06 0.064 1016 12 BDL BDL 59,068 1716 1169 a 1027 34 0.03 0.052 1027 27 0.026 0.034 59,068 2060 523 1112 35 0.03 0.051 1112 8 BDL BDL	59,623 2746 1374 938 50 0.05 0.038 938 19 0.020 0.018 54 59,623 2880 838 1016 65 0.06 0.064 1016 12 BDL m.d. 59,068 1716 1169 a 1027 34 0.03 0.052 1027 27 0.026 0.034 34 59,068 2060 523 1112 35 0.051 1112 8 BDL BDL n.d.		-38.954		1136			971		0.02	0.023	971	27		0.005	79			
59.623 2880 838 1016 65 0.06 0.064 1016 12 BDL BDL 59.068 1716 1169 a 1027 34 0.03 0.052 1027 27 0.026 0.034 59.068 171 BDL BDL BDL BDL BDL BDL 59.068 2060 523 1112 35 0.03 0.051 1112 8 BDL BDL	59.623 2880 838 1016 65 0.06 0.064 1016 12 BDL BDL BDL n.d. 59.068 1716 1169 a 1027 34 0.03 0.052 1027 27 0.026 0.034 34 59.068 2060 40 0.04 0.050 990 11 BDL BDL n.d. 59.068 2060 523 1112 35 0.03 0.051 1112 8 BDL BDL n.d.		-38.954		2746			938		0.05	0.038	938	19		0.018	54			
46.083 59.068 1716 1169 a 1027 34 0.03 0.052 1027 27 0.026 0.034 46.083 59.068 2060 40 0.04 0.050 990 11 BDL BDL 46.083 59.068 2060 523 1112 35 0.03 0.051 1112 8 BDL BDL	46.083 59.068 1716 1169 a 1027 34 0.03 0.052 1027 27 0.026 0.034 34 46.083 59.068 2060 523 1112 35 0.03 0.051 1112 8 BDL BDL DDL 1112 8 BDL BDL DDL n.d.		-38.954		2880			1016		90.0	0.064	1016	12	BDL	BDL	n.d.			
46.083 59.068 2060 523 1112 35 0.03 0.051 1112 8 BDL BDL	46.083 59.068 2060 523 1112 35 0.03 0.051 1112 8 BDL BDL n.d.		-46.083		1716			1027	34	0.03	0.052	1027	27		0.034	34			
46.083 59.068 2060 523 1112 35 0.03 0.051 1112 8 BDL BDL	46.083 59.068 2060 523 1112 35 0.03 0.051 1112 8 BDL BDL n.d.						Р	066	40	0.04	0.050	066	11		BDL	n.d.			
	r no pomining on m		-46.083	59.068	2060			1112	35	0.03	0.051	1112	8	BDL	BDL	n.d.			

Station	Station Longitude Latitude Depth Volume Replicates Cartridge A	Latitude	Depth	Volume	Replicates	Cartridge 1	A			Cartridge B	3						
						Runtime ²¹⁹ Rn		²¹⁹ Rn :	²¹⁹ Rn ²²⁷ Ac on Mn- cartridge A	Runtime ²¹⁹ Rn		²¹⁹ Rn	²¹⁹ Rn ²²⁷ Ac on Mn- cartridge B	Extraction efficiency	Replicates Mean	Replicates SD (10)	Replicates RSD (10)
	(°E)	(N°)	(m) (L)	(T)		(min) (counts)		(cpm) (dpm)	(mdp)	(min)	(min) (counts) (cpm) (dpm)	(cpm)	(mdp)	%	%	%	%
64	- 46.083			1365		1032	46	0.04).053	1032	12		BDL	n.d.			
64	-46.083	59.068	2443	276		933	47	0.05	0.076	933	30		0.033	57			
69	-48.093			418	В	1147		0.05	0.073	1147	18 (0.018	75	64	17	26
					þ	962		0.05	0.081	962	22 (0.039	52			
69	-48.093		2953	277		296		0.04	0.050	296			BDL	n.d.			
69	-48.093	55.841	3440	1169		1148	107	0.09	0.123	1148	15 1	BDL 1	BDL	n.d.			
69	-48.093	55.841	3616	738	а	1393		0.03	0.050	1393			0.034	31	35	10	29
					þ	1038		0.02	7.034	1038			0.028	17			
					c	962		0.03	0.042	962	23 (0.018	26			
													Weighted average		47	12	

by excluding either samples displaying relatively small seawater volumes (i.e. below 1000 L), or samples displaying low $^{227}\mathrm{Ac}$ activities yield to the same values within errors. This extraction efficiency (E) of 47% was applied to all samples (including samples collected using single Mn-cartridges and samples collected using Mn-cartridges in series). Finally, the $^{227}\mathrm{Ac}$ activities of the seawater samples, A_{227Ac} , were estimated as follows:

$$A_{227_{AC}} = \frac{^{227}Ac_{Cart A}}{E} \tag{2}$$

The extraction efficiency of 47 \pm 12% is lower than those reported by previous studies. In a study conducted in the Pacific Ocean, Kipp et al. (2015) assumed an ²²⁷Ac extraction efficiency for Mn-cartridges of 70%. However, this value was not determined by analyzing ²²⁷Ac directly in the Mn-cartridges. In this latter study, the extraction efficiencies have been determined for Ra and Th isotopes. A mean value was thus determined from these Ra and Th extraction efficiencies (i.e. 70%) and was actually considered to be representative of ²²⁷Ac Mncartridge extraction efficiency. Geibert and Vöge (2008) and Geibert et al. (2002) reported extraction efficiencies for 227 Ac of 77 \pm 13% (n = 14) and $69 \pm 11\%$ (n = 31), while Kemnitz (2018) and Hammond et al., (pers. comm.) reported mean extraction efficiencies for ²²⁷Ac of 54% and 65%, respectively (Table 5). A significant variability in the ²²⁷Ac extraction efficiencies is thus observed. Such variability may be due to the composition of the Mn-cartridges (acrylic vs polypropylene) or to the size of the Mn-cartridges used in the different studies or to a combination of the two (Table 5). Leaks may also reduce the extraction efficiency, in case the sealing between the cartridge holder and the Mn-cartridge is not optimal; such a problem cannot be completely excluded in the present study, since the relatively small Mncartridges that we used were placed into large cartridge holders. Geibert and Vöge (2008) used the largest Mn-cartridges (25 cm long polypropylene cartridges) and obtained relatively high extraction efficiencies. (Kemnitz, 2018) used smaller Mn-cartridges (12.7 cm long acrylic Mn-cartridges) and the extraction efficiency was significantly reduced compared to Hammond et al., (pers. comm.) who used the same length of Mn-cartridges but used cellulose instead of acrylic. The Mn-cartridges used in the present studies were even shorter (7.7 cm long acrylic cartridges) and the extraction efficiency is further reduced. Both the size and the formulation of the Mn-cartridge may thus impact the yield of ²²⁷Ac extraction. Note that we impregnated the Mn-cartridges using a continuous flow of KMnO₄ through the cartridge (see section 2.1.1.). This step does not seem to improve significantly the extraction efficiency.

In a parallel study, 226 Ra activities were measured in seawater samples collected during the GEOVIDE cruise at the same depth, using Mn-fibers and radon emanation technique (Le Roy et al., 2018). Several Mn-cartridges were also analyzed for 226 Ra by gamma spectrometry (same samples as for 227 Ac, unpublished data). These data allowed us to estimate the Mn-cartridge extraction efficiencies for 226 Ra in a similar manner as described above. The extraction efficiencies for 226 Ra were found on average to be $60 \pm 16\%$ (1 SD, n = 15). The 226 Ra extraction efficiency thus obtained is in good agreement with the extraction efficiency of 226 Ra on acrylic Mn-cartridges of 54% reported by (Henderson et al., 2013).

The 226 Ra extraction efficiency of the Mn-cartridges is thus significantly higher than the 227 Ac extraction efficiency estimated in the same Mn-cartridges. This result shows that assuming the same extraction efficiency between Ra and Ac could lead to significant biases. The systematic determination of the 227 Ac extraction efficiency seems therefore required. Finally, note that there were no significant correlations between efficiency and other parameters such as volume, flow rate, station, water depth, activity on the A Mn-cartridge or the final activity. The extraction efficiency determined in this study will thus be applied to all the samples collected along the GEOVIDE section to provide an entire section of dissolved 227 Ac activities.

Table 5Comparison of the extraction efficiencies for ²²⁷Ac on Mn-cartridges determined by different studies. The analytical method used, the composition and the size of the Mn-cartridges are also reported.

Reference	Mean	SD	Range	9		n	Analytical method	Formulation	size	Method
This study Kemnitz (2018)	47% 54%	12% 6%	31%	-	78%	9	Delayed coincidence counting Delayed coincidence counting		7.7 12.7	2 Mn-cartridges in serie 2 Mn-cartridges in serie
Hammond et al. (Pers. Comm.) Geibert et al. (2002) Geibert and Vöge (2008) Kipp et al. (2015)	65% 69% 77% 70%	11% 13%	45% 41%	-	90% 95%		Delayed coincidence counting Alpha spectrometry Alpha spectrometry Delayed coincidence counting	Polypropylene Polypropylene	12.7 25 25 12.7	2 Mn-cartridges in serie 2 Mn-cartridges in serie 2 Mn-cartridges in serie Mean between Ra and Th extraction efficiency

3.2.2. $^{227}\mathrm{Ac}$ determined from Mn-cartridges vs $^{227}\mathrm{Ac}$ determined from Mn-fibers

Six discrete seawater samples were collected and were passed by gravity through Mn-fibers on board. Because Mn-fibers quantitatively adsorb ²²⁷Ac (Reid et al., 1979), this allowed us to validate the ²²⁷Ac activities determined from the Mn-cartridges. Note, however, that although Mn-fibers have the advantage of adsorbing 100% of ²²⁷Ac, the collection of reduced volumes of seawater (120 to 541 L) with Niskin bottles yields to low activities on the Mn-fibers, and the associated uncertainties, therefore, are relatively high. Consequently, in the present paper, we did not estimate the extraction efficiency of the Mn-cartridges by comparing the ²²⁷Ac activities on the Mn-Fibers and those on the Mn-cartridges. Here, we use the ²²⁷Ac activities determined using Mn-fiber as a comparison.

Results are shown in Table 6. Fig. 3 shows the seawater 227 Ac activities determined from Mn-cartridges (corrected for the extraction efficiency) versus the seawater 227 Ac activities measured from the Mn-fibers. The 1:1 line is also plotted on Fig. 3. Although the two deepest samples (station 32–3051 m and 3171 m depth) are slightly outside of the 1:1 line, possibly due to lower volume (only 89 L ont Mn-cartridge) at 3051 m. The 227 Ac activities determined with the two methods are in relatively good agreement with each other. Overall, this Mn-cartridge vs Mn-fiber comparison gives confidence to the extraction efficiency determined here (47 \pm 12%).

3.3. Comparison with historical data

There are a limited number of studies that have reported ²²⁷Ac activities in the open ocean to date (Geibert et al., 2008, 2002; Kipp et al., 2015; Koch-Larrouy et al., 2015; Nozaki, 1984). Fig. 4 includes these ²²⁷Ac profiles determined in the open ocean in different oceanic basins together with station 32 and 38 from GEOVIDE cruise (this study). Above 2000 m, ²²⁷Ac activities from the GEOVIDE cruise in the North Atlantic are within the same range of those determined in the Arctic Ocean and the Pacific Ocean. Below 2000 m, ²²⁷Ac activities from the GEOVIDE cruise are lower than in other basins except for the

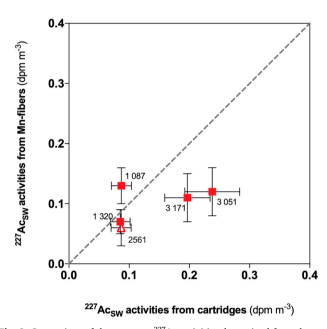


Fig. 3. Comparison of the seawater ²²⁷Ac activities determined from the measurement of Mn-fibers (water sampled using Niskin bottles) and of Mn-cartridges (deployed on in situ pumps). Labels indicate the depths of the samples. One of these Mn-fibers (triangle) has no matching depth Mn-cartridge. For this sample, a mean of the ²²⁷Ac activities from Mn-cartridges located below and under the Mn-fiber sample is used here. The grey line represents the 1:1 line.

Arctic Ocean. The ²²⁷Ac activities from North Atlantic (GA03; Kipp et al., 2015) were measured in samples collected in a hydrothermal plume over the mid-Atlantic ridge in the Atlantic Ocean, which explain the higher ²²⁷Ac compare to our study (GEOVIDE). The low ²³¹Pa activities along the GEOVIDE section, compare to other basins, are due to ventilated overflow waters in the North Atlantic, particularly in the Labrador and Irminger Seas (Deng et al., 2018). The low ²³¹Pa activities

Table 6Comparison between ²²⁷Ac activities determined from Mn-fibers and from Mn-cartridges A (uncorrected for extraction efficiency). The dissolved ²²⁷Ac activities in seawater estimated using Mn-cartridges (corrected for extraction efficiency) are also reported. The standard deviation (SD) is determined from repeated measurement of the same samples as described in section 3.1.3.

Station	Depth	Mn-fiber					Mn-cartrid	ges				
		Runtime	²¹⁹ Rn	²¹⁹ Rn	²²⁷ Ac	SD (1σ)	Runtime	²¹⁹ Rn	²¹⁹ Rn	²²⁷ Ac on Mn-cartridge A	²²⁷ Ac in SW	SD (1σ)
	(m)	(min)	(counts)	(cpm)	(dpm m	⁻³)	(min)	(counts)	(cpm)	(dpm m ⁻³)		
32	2267						972	33	0.034	0.03	0.072	0.013
32	2561	4153	29	0.007	0.058	0.028					0.078	0.014
32	2854						1089	34	0.031	0.04	0.084	0.015
32	3051	4153	47	0.011	0.118	0.037	995	73	0.073	0.10	0.212	0.039
32	3171	4153	37	0.009	0.108	0.037	894	43	0.048	0.08	0.175	0.032
32	3217	4153	61	0.015	0.184	0.049						
38	1087	4090	79	0.019	0.126	0.031	1120	39	0.035	0.04	0.078	0.014
38	1320	4060	142	0.035	0.067	0.017	1061	44	0.042	0.03	0.074	0.014

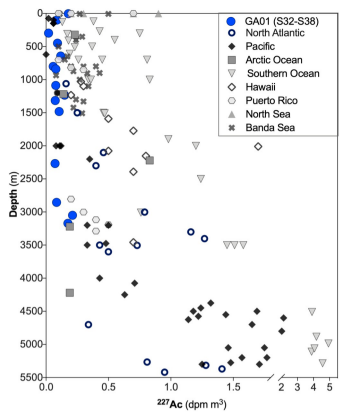


Fig. 4. ²²⁷Ac activities versus depth published to date in the water columns of different oceanic basins (Geibert et al., 2008, 2002; Kipp et al., 2015; Koch-Larrouy et al., 2015; Nozaki, 1984), together with the ²²⁷Ac activities determined at station 32 and 38 along the GEOVIDE section.

in the North Atlantic lead to a small production of 227 Ac which explain the relatively low 227 Ac activities compare to the Southern Ocean or in deep water of the Pacific Ocean.

3.4. Seawater ²²⁷Ac vertical profiles and comparison with ²³¹Pa

In the aim of further validating the protocol, vertical profiles of ²²⁷Ac—determined from Mn-cartridges and from Mn-fibers—are compared to the vertical profiles of ²³¹Pa at stations 32 and 38 (Fig. 5; ²³¹Pa

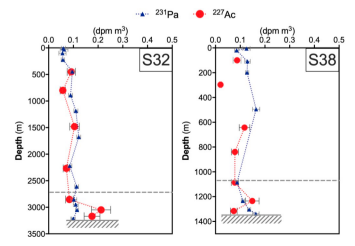


Fig. 5. Vertical profiles of dissolved ²²⁷Ac (from Mn-cartidges and Mn-fibers) determined at stations 32 and 38 along the GEOVIDE section (GEOTRACES GA01) in the Iceland Basin. For comparison, the dissolved ²³¹Pa activities vertical profiles are also reported (black dots, data from Deng et al., 2018).

data from Deng et al., 2018). 231 Pa and 227 Ac have different chemical behavior in the water column. 231 Pa adsorbs onto particle surfaces and is thus scavenged by settling particles. In contrast, 227 Ac is less reactive and is supposed to remain in the dissolved phase. In the first 2000 m, 227 Ac is produced by the decay of the remaining 231 Pa in the water column. Therefore, 227 Ac and 231 Pa activities are expected to be at secular equilibrium in the upper 2000 m, away from external sources of 227 Ac. 227 Ac and 231 Pa activities show a good agreement within error bars above 2000 m where 227 Ac and 231 Pa are expected to be in secular equilibrium because 227 Ac is produced in the water column from the 231 Pa decay (Nozaki et al., 1990). This is an additional validation of the overall protocol described here. The error bars reported here for the dissolved 227 Ac activities (repeatability experiment, 19% 1 SD) suggest the validation of the overall protocol and its accuracy, including its trueness (absence of bias, cf. good agreement with 231 Pa) and its precision (the internal precision of 19%, cf. error bars on the plot).

At station 38, the ²³¹Pa activities increase close to the bottom. The ²²⁷Ac activities do not seem to follow such a pattern. The ²²⁷Ac activities are thus significantly below ²³¹Pa activities close to bottom at 1300 m. This is an unexpected feature, as one may expect that ²²⁷Ac and ²³¹Pa are at secular equilibrium over such water columns, or in excess close to bottom. This pattern could be associated to the presence of nepheloid layers (Gourain et al., 2018). Presence of MnO2 resuspended from the sediments may adsorb ²²⁷Ac, while ²³¹Pa may be released from resuspended particles. At station 32, below 2000 m, the ²²⁷Ac activities are significantly higher than the ²³¹Pa activities. These excess ²²⁷Ac activities likely reflect input of unsupported ²²⁷Ac from the deep-sea sediments as previously observed in other ocean basins (Geibert et al., 2002; Nozaki, 1984; Nozaki et al., 1990). The entire section of dissolved ²²⁷Ac activities will be reported in a separate paper (Le Roy et al., in prep.), where it will be compared to the section of dissolved ²³¹Pa activities (Deng et al., 2018), and also to several other trace elements.

4. Conclusion

This study shows that ²²⁷Ac can be measured accurately in the open ocean using Mn-cartridges mounted on in situ pumps and using the RaDeCC system. This study is the second one using Mn-cartridges to extract ²²⁷Ac from seawater followed by their analysis using RaDeCC and is the first one that aims to quantify the extraction efficiencies of ²²⁷Ac on Mn-cartridges using Mn-cartridges placed in series. The low background of the RaDeCC systems leads to a limit of quantification of 0.014 cpm (10 times the SD). Extraction efficiencies of ²²⁷Ac from seawater using Mn-cartridges were measured to be 47 \pm 12% (1 SD, n = 9) for sample sizes from 418 to 1565 L. For seawater with low ²²⁷Ac activities such as in the North Atlantic, a minimum of 800 L per samples would be recommended so that significant activities on the Mn-cartridge B can be quantified, which in turn allows us to estimate the extraction efficiency. For future studies, Mn-cartridges in series should also be placed at all water depths so that the ²²⁷Ac extraction efficiency can be determined for each sample.

Replicate analyses of several samples collected during the GEOVIDE cruise allowed us to estimate that the Mn-cartridge $^{227}\mathrm{Ac}$ activity measurement uncertainty is $19\pm14\%$ (1 SD, n=20), that is significantly more precise than what would have been calculated through error propagation following Garcia-Solsona et al., 2008; i.e., 33%). This precision is sufficient to clearly detect and quantify $^{227}\mathrm{Ac}$ variations in the North Atlantic Ocean, thus opening new perspectives to better exploit $^{227}\mathrm{Ac}/^{231}\mathrm{Pa}$ ratios (or excess $^{227}\mathrm{Ac}$ activities) as oceanic tracers, notably for studying deep water mixing.

The advantages of this method are the low backgrounds of the RaDeCC detectors and reduced chemical procedures compared to other techniques (Geibert and Vöge, 2008; Shaw and Moore, 2002). An additional advantage of the delay coincidence counting method is that it is possible to analyze radium and thorium isotopes at the same time.

However, extraction efficiency for radium and thorium would be different from ²²⁷Ac extraction efficiency.

In the aim of further validating the protocol, two vertical profiles from the GEOVIDE cruise were reported. Above 2000 m ²²⁷Ac is in equilibrium with its parent ²³¹Pa, which do validate the protocol. Below 2000 m, ²²⁷Ac excesses compared to ²³¹Pa may be used to estimate sedimentary fluxes of ²²⁷Ac or—as was suggested by previous studies—to study vertical mixing in the deep ocean on time scales of years to decades, especially in association with other radionuclides such as ²²⁸Ra. The low ²²⁷Ac activities reported along the GEOVIDE section, compare to other basins, suggest recently ventilated overflow waters in the North Atlantic. Overall, the technique is validated and offers significant advantages over previous techniques. This opens up opportunities for large sections in the future, particularly in the context of the GEOTRACES programme.

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