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Graphic Abstract



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1 First dataset of ^{236}U and ^{233}U around the Greenland
2 coast: a 5-year snapshot (2012-2016)

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10 **ABSTRACT**

11 We report the first combined dataset of ^{236}U and ^{233}U in the Greenland marine environment during the
12 period of 2012-2016. Results are discussed in terms of time evolution and spatial distribution of ^{236}U
13 concentration, and atomic ratios of $^{236}\text{U}/^{238}\text{U}$ and $^{233}\text{U}/^{236}\text{U}$. ^{236}U concentrations along the Greenland coast
14 are distributed within a relatively narrow range of $(0.7\text{-}12.9) \times 10^7$ atom/L, corresponding to $^{236}\text{U}/^{238}\text{U}$
15 atomic ratios of $(1.1\text{-}15.5) \times 10^{-9}$. The $^{233}\text{U}/^{236}\text{U}$ atomic ratios obtained vary from 0.12×10^{-2} to 1.16×10^{-2} ,
16 with the majority distributed in the range of $(0.2\text{-}0.7) \times 10^{-2}$.

17 We applied $^{233}\text{U}/^{236}\text{U}$ and $^{236}\text{U}/^{238}\text{U}$ atomic ratios in a binary mixing model to identify possible ^{236}U
18 source terms. The results indicate that anthropogenic ^{236}U and ^{233}U in Greenland surface seawater
19 originated from the direct global fallout (GF) and the Sellafield and La Hague reprocessing plants (RP) is
20 diluted by a third endmember, mostly likely natural ocean water (NOW), containing marginal ^{236}U and
21 ^{233}U . A preliminary estimation of the source terms of ^{236}U using the $^{233}\text{U}/^{236}\text{U}$ atomic ratio indicate that for
22 both eastern and western Greenland seawater contributions from GF constitute about 30 % of ^{236}U . The
23 dominating source for ^{236}U , i.e. 70 %, is associated to reactor ^{236}U including discharges from RP and local
24 reactor input in the Arctic Ocean.

25

26 **Keywords**

27 ^{233}U , ^{236}U , Greenland coast, surface seawater, 2012-2016

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29 1. INTRODUCTION

30 Nuclear accidents (e.g., Chernobyl, Fukushima), weapons production and reprocessing plants (e.g.,
31 Sellafield (SF) in UK and La Hague (LH) in France) have released large amounts of man-made
32 radioactive contaminants to the environment. Some radioisotopes (e.g., ^3H , ^{14}C , ^{134}Cs , ^{137}Cs , ^{129}I , ^{99}Tc)
33 have relatively high mobility and bioavailability in oxic waters, making them valuable tracers in
34 oceanographic studies (Hou et al., 2002; Karcher et al., 2004; Smith et al., 2011). In recent years, the
35 long-lived minor uranium isotope ^{236}U ($T_{1/2} = 2.34 \times 10^7$ y) has been identified as an appealing new
36 oceanographic tracer (Casacuberta et al., 2014; Christl et al., 2015, 2013, 2012; Eigl et al., 2013;
37 Sakaguchi et al., 2012; Winkler et al., 2012) due to its superior properties including high solubility, long
38 residence time (10^5 y), and the well-understood steady circulation of natural uranium in the open ocean
39 (Al-qasmi et al., 2016; Bellucci et al., 2013; Boulyga et al., 2002; Casacuberta et al., 2016; Chamizo et
40 al., 2008; Desideri et al., 2002; Hotchkis et al., 2000; Lee et al., 2008; Parrish et al., 2006; Purser et al.,
41 1996; Qiao et al., 2017; Quinto et al., 2009; Sakaguchi et al., 2016; Steier et al., 2008; Wendel et al.,
42 2013; Winkler et al., 2012; Yang et al., 2016).

43 The Arctic is a key and most sensitive area of global climate change (Hansen et al., 2010), however,
44 many processes in ocean circulation in the Arctic are poorly understood due to the lack of observations.
45 The Arctic Ocean is surrounded by a disproportionately large continental shelf sea area into which several
46 large rivers flow, supplying 10 % of the global river discharge (McClelland et al., 2012). The surface
47 oceanic current, Greenland Current, is a combination of polar sea surface drift, return flow of the North
48 Atlantic Current, and Irminger Current waters. The East Greenland Current (EGC) is of major importance
49 because it directly connects the Arctic to the Northern Atlantic. The EGC is a major contributor to sea ice
50 export out of the Arctic (Woodgate et al., 1999), and it is a major freshwater sink for the Arctic
51 (Schlichtholz and Houssais, 1999).

52 Anthropogenic ^{236}U (>1000 kg) clearly dominates over its natural inventory (about 35 kg) in the
53 environment, with the primary source terms including global fallout from atmospheric nuclear weapons
54 testing (900-1400 kg) and discharges from reprocessing plants at SF and LH (115-250 kg) (Castrillejo et

55 al., 2020; Sakaguchi et al., 2009; Steier et al., 2008). Even though a growing set of ^{236}U data exists for the
56 open oceans, reported ^{236}U data in the Arctic Ocean, especially in Greenland marine system are very
57 limited (Castrillejo et al., 2018; Wefing et al., 2019). A good understanding of the input function of ^{236}U
58 is critical for establishing it as a reliable tracer. However, the ^{236}U budget in the Atlantic-Arctic Ocean is
59 still an open question due to incomplete records of the ^{236}U emission from SF and LH (Castrillejo et al.,
60 2020; Christl et al., 2015, 2013).

61 Moreover, tracer studies using solely ^{236}U suffer from methodological difficulties to distinguish
62 variations in ^{236}U source terms. Several studies have proposed the combination with other radionuclides,
63 e.g. ^{129}I , to trace different water masses (Casacuberta et al., 2018, 2016; Castrillejo et al., 2018; Wefing et
64 al., 2019). However, due to the different input functions and transport pathways of ^{129}I and ^{236}U , as
65 well as different chemical-physical behaviour between these two elements, challenges in $^{236}\text{U}/^{129}\text{I}$
66 tracer applications are still encountered.

67 ^{233}U ($T_{1/2} = 1.59 \times 10^5$ y) is also an anthropogenic uranium trace isotope like ^{236}U . Different from $^{236}\text{U}/^{129}\text{I}$
68 ratios, $^{233}\text{U}/^{236}\text{U}$ ratios are unaffected by the environmental pathways. A recently published study
69 demonstrated the high potential for $^{233}\text{U}/^{236}\text{U}$ to be used as a robust fingerprint to distinguish emission
70 sources of anthropogenic U (Hain et al., 2020). ^{233}U was mostly produced during nuclear weapons testing
71 by fast neutrons via $^{235}\text{U}(\text{n}, 3\text{n})^{233}\text{U}$ reactions or directly by ^{233}U -fueled devices, whereas almost no ^{233}U
72 is produced in thermal nuclear power reactors or reprocessing plants (Hain et al., 2020). The
73 representative $^{233}\text{U}/^{236}\text{U}$ atomic ratio was suggested to be $(1.40 \pm 0.12) \times 10^{-2}$ for global fallout. $^{233}\text{U}/^{236}\text{U}$
74 atomic ratios in LH discharges are at the level of 1×10^{-7} - 1×10^{-6} (HELCOM MORS Discharge database),
75 which is in good agreement with reactor model calculations obtained for the fuel of pressurized water
76 reactors (Naegeli, 2004). In the Irish Sea, an average $^{233}\text{U}/^{236}\text{U}$ atomic ratio of $(0.12 \pm 0.01) \times 10^{-2}$ was
77 measured (Hain et al., 2020).

78 In this work, we report data on $^{233}\text{U}/^{236}\text{U}$ in Greenland seawater for the first time and aim to draw a
79 picture about the levels, distribution pattern and source terms of ^{236}U in Greenland marine environment
80 thus to prompt the application of ^{236}U - ^{233}U as paired-tracer system in oceanographic studies.

81 **2. EXPERIMENTAL**

82 **2.1 Samples, standards, reagents and samples.**

83 Sampling around the Greenland coastal line during 2012-2016 was carried out in collaboration with
84 Greenland Institute of Natural Resources and National Institute of Aquatic Resources (DTU Aqua),
85 Denmark. Fig. 1 gives an overview of sampling locations and the main surface current systems in the
86 North Atlantic - Nordic Seas-Arctic Ocean (Aarkrog et al., 1999; Hou and Hou, 2012; Iosjpe et al., 2013;
87 Pedersen et al., 2005). The details of the currents around Greenland coast, sample information as well as
88 the overall analytical results obtained in this work (Table S-1 to S-4) are given in the supplementary
89 information. All the samples (ca. 600 L of each) were surface seawater from a depth of about 5 m, which
90 were pumped into big barrels and then taken for determination of ^{238}U by ICP-MS (50 mL), $^{236}\text{U}/^{238}\text{U}$ and
91 $^{233}\text{U}/^{236}\text{U}$ by AMS (2-5 L) and other radionuclides (e.g., ^{99}Tc , results are not shown here) in the
92 laboratory.

93 Uranium standard solution (1.000 g/L in 2 M HNO_3) was purchased from NIST (Gaithersburg, MD),
94 which was used after dilution as a standard for the inductively coupled plasma mass spectrometry (ICP-
95 MS) measurement to quantify ^{238}U in seawater. All reagents used in the experiment were of analytical
96 reagent grade and prepared using ultra-pure water (18 $\text{M}\Omega\cdot\text{cm}$). UTEVA resin (100-150 μm particle size)
97 was purchased from Triskem International, Bruz, France and packed in 2-mL Econo-Columns (0.7 cm i.d.
98 \times 5 cm length, Bio-Rad Laboratories Inc., Hercules, CA) for the chemical purification of uranium
99 isotopes.

100 **2.2 Analytical methods for determination of ^{238}U , ^{236}U , and ^{233}U .**

101 ^{238}U in the ocean is of natural origin and not related to anthropogenic emissions. The ^{238}U present in the
102 water samples was used for the normalization of the detected ^{236}U and ^{233}U in order to take into account
103 the chemical yield and the detection efficiency for U. The concentration of ^{238}U in seawater was directly

104 measured by triple-quadrupole ICP-MS (Agilent 8800 ICP-QQQ, Agilent technologies) after 50-folder
105 dilution. The ICP-QQQ instrument was equipped with a standard introduction system consisting of a
106 MicroMist nebulizer and a Scott-type double-pass spray chamber, together with a Ni sampler cone, Ni
107 skimmer cone and x-lens. The experimental details for the ^{238}U direct measurement have been given
108 elsewhere (Qiao and Xu, 2017). The combined uncertainty for ^{238}U is estimated to be 10 %, consisting of
109 statistical counting uncertainty of samples, standard and blank, dilution of samples and standards,
110 uncertainty of certified value of ^{238}U standard and matrix effect.

111 The sample preparation procedure for ^{233}U and ^{236}U determination in seawater is based on our previous
112 report (Qiao et al., 2015). In short, the seawater samples with a volume of 2-5 L were filtrated with filter
113 paper (Munktell 00K, particle retention 5-6 μm) to remove large particles and then acidified to pH 2 with
114 concentrated HNO_3 . Purified FeCl_3 solution (0.05 g/mL of Fe) was added to a final Fe concentration of
115 0.1 g/L. The sample was vigorously stirred with air bubbling for 5-10 minutes in order to decompose
116 carbonate complexes. 10 % $\text{NH}_3\cdot\text{H}_2\text{O}$ was slowly added to adjust the pH to 8-9 for the co-precipitation of
117 U with $\text{Fe}(\text{OH})_3$. The precipitate was allowed to settle for 0.5-1 h in order to decant most of the
118 supernatant. The sample slurry was centrifuged at 4000 rpm for 5 minutes and the supernatant was
119 discarded. The final residue was dissolved with 15 mL of 3 M HNO_3 and the solution was loaded onto a
120 2-mL UTEVA column which was pre-conditioned with 20 mL of 3 M HNO_3 . The UTEVA column was
121 rinsed with 40 mL of 3 M HNO_3 , followed by 20 mL of 6 M HCl . Uranium absorbed on the column was
122 eluted with 10 mL of 0.025 M HCl . The flow rate for the chromatographic separation was controlled
123 manually to 1.0-1.5 mL/min.

124 1-3 mg of Fe (as FeCl_3 solution) was added to the U eluate, and the sample was adjusted to $\text{pH}>9$ with
125 ammonia to co-precipitate U with $\text{Fe}(\text{OH})_3$. The precipitate was dried in an oven at 100 $^\circ\text{C}$ and was then
126 baked in a furnace at 800 $^\circ\text{C}$ for 12 hours so that the U is finally embedded in a Fe_2O_3 matrix. The sample
127 was pressed into aluminium sputter target holders for the AMS measurement of $^{236}\text{U}/^{238}\text{U}$ and $^{233}\text{U}/^{236}\text{U}$.
128 The AMS measurements were carried out at the 3-MV tandem accelerator facility VERA (Vienna
129 Environmental Research Accelerator) at the University of Vienna, Austria. The detailed method

130 for AMS measurements of ^{233}U and ^{236}U is reported elsewhere (Hain et al., 2020; Steier et al., 2019;
131 Winkler et al., 2015).

132 **2.3 Data quality control**

133 For each batch of seven seawater samples, a process blank was prepared and analysed with respect to
134 $^{236}\text{U}/^{238}\text{U}$ and $^{233}\text{U}/^{236}\text{U}$ ratios by AMS and ^{238}U concentration by ICP-MS, to monitor potential
135 contaminations during the AMS measurement and the chemical sample preparation. Some process blanks
136 showed occasionally elevated levels of ^{236}U and ^{233}U , which might be induced by the air-borne/particle-
137 associated contamination from the ventilation system in the building (which has been worked with high
138 radioactive waste samples from nuclear reactors). If the ^{236}U or ^{233}U count rate of the corresponding blank
139 was more than 30 % of the respective sample, this sample including process blank was re-prepared, if
140 possible, or rejected. For all the reported samples in this work, the blank level was taken into account in
141 the uncertainties of the final measurement results, in some cases exceeding the uncertainty of the AMS
142 measurement. The reference material IAEA-381 Irish seawater was analysed with respect to the $^{236}\text{U}/^{238}\text{U}$
143 ratio for quality assurance showing a very good agreement (i.e. error within $\pm 1\sigma$) with the reference value
144 (results are not shown here).

145 **3. RESULTS AND DISCUSSION**

146 **3.1 ^{236}U concentration, $^{236}\text{U}/^{238}\text{U}$ and $^{233}\text{U}/^{236}\text{U}$ atomic ratios**

147 The distributions of the detected ^{236}U concentrations and $^{236}\text{U}/^{238}\text{U}$ atomic ratios are presented
148 graphically in Fig. 2 along with some published data for ^{236}U in the Arctic Ocean. The ^{238}U concentrations
149 are in the range of 2.6-3.3 $\mu\text{g/L}$, show general positive correlation with salinity (Fig. S1). The low
150 correlation coefficient ($R^2=0.34$) is attributed to the narrow distribution range and relatively high
151 uncertainties (ca. 10%) in the ^{238}U results. The $^{236}\text{U}/^{238}\text{U}$ atomic ratios obtained vary within $(1.1-15.5) \times$
152 10^{-9} , with majority of the results falling within $(1.1-6.2) \times 10^{-9}$, which are up to 6 times higher than the
153 estimated global fallout $^{236}\text{U}/^{238}\text{U}$ background level (1×10^{-9}) in the open ocean surface (Christl et al.,
154 2012). Two samples from 2013 (2013-0538 and 2013-0537) collected near the eastern Greenland coast

155 indicate much higher $^{236}\text{U}/^{238}\text{U}$ atomic ratios (8.5×10^{-9} and 15.5×10^{-9} , respectively). The ^{236}U
156 concentrations vary within $(0.7\text{-}12.9) \times 10^7$ atom/L with majority in the range of $(0.7\text{-}4.7) \times 10^7$ atom/L.
157 Again, the two above-mentioned samples show much higher ^{236}U concentrations (6.7×10^7 atom/L for
158 2013-0538 and 12.9×10^7 atom/L for 2013-0537, respectively). The validity of the ^{236}U measurements for
159 these two samples (2013-0538 and 2013-0537) was confirmed by independently analysing 2-3 replicates
160 for each sample.

161 $^{233}\text{U}/^{236}\text{U}$ atomic ratios obtained in this work vary from 0.12×10^{-2} to 1.16×10^{-2} , with the majority
162 distributed in the range of $(0.2\text{-}0.7) \times 10^{-2}$ (except 2013-0537 $(0.12 \pm 0.01) \times 10^{-2}$ and 2012-0542 $(1.16 \pm$
163 $0.27) \times 10^{-2}$) (Fig. 3). It is noteworthy that the sample (2013-0537) with the highest ^{236}U concentration
164 shows the lowest $^{233}\text{U}/^{236}\text{U}$ atomic ratio $(0.12 \pm 0.01) \times 10^{-2}$ among all samples collected in 2013
165 indicating a reactor signal of ^{236}U . The $^{233}\text{U}/^{236}\text{U}$ atomic ratio for the 2013-0538 sample of $(0.68 \pm 0.34) \times$
166 10^{-2} shows a high uncertainty (50 %) and therefore the obtained $^{233}\text{U}/^{236}\text{U}$ atomic ratio is excluded from
167 interpretation.

168 The reported ^{236}U data in the Arctic region are limited. Casacuberta et al. (Casacuberta et al., 2016)
169 observed ^{236}U ranging from 0.8×10^7 to 3.0×10^7 atom/L, with an average concentration of $(2.3 \pm 0.6) \times 10^7$
170 atom/L for Arctic surface seawater in 2011-2012 (Casacuberta et al., 2016). Our results for ^{236}U
171 concentrations in Greenland seawater in 2012 $((0.7\text{-}3.4) \times 10^7$ atom/L, with an average of $(1.9 \pm 0.1) \times 10^7$
172 atom/L) are comparable to their reported values. Our measured ^{236}U concentrations for 2015 $((1.2\text{-}3.1) \times$
173 10^7 atom/L, with an average of $(2.2 \pm 0.6) \times 10^7$ atom/L) also show good agreement with the observations
174 by Casacuberta et al. (Casacuberta et al., 2018) for surface seawater from Barents Sea shelf, West
175 Spitsbergen, Eurasian Basin and Makarov Basin collected in 2015 $((1.1\text{-}2.7) \times 10^7$ atom/L, with an
176 average of $(1.6 \pm 0.4) \times 10^7$ atom/L). Wefing et al. (Wefing et al., 2019) reported ^{236}U concentrations of
177 $(1.2\text{-}2.2) \times 10^7$ atom/L, with an average of $(1.8 \pm 0.3) \times 10^7$ atom/L for samples from Fram Strait in 2016,
178 which agree well with our results in 2016 $((0.8\text{-}2.0) \times 10^7$ atom/L, with an average of $(1.5 \pm 0.4) \times 10^7$
179 atom/L).

3.2 The time evolution and spatial distribution

3.2.1 Time evolution.

Box-plots of yearly ^{236}U concentration, as well as $^{236}\text{U}/^{238}\text{U}$ and $^{233}\text{U}/^{236}\text{U}$ atomic ratio for samples from East Greenland coast (EG) and West Greenland coast (WG) are shown in Fig. 4. The average of $^{236}\text{U}/^{238}\text{U}$ atomic ratios around the Greenland coast during 2012-2015 are at the level of $(2-4) \times 10^{-9}$, with average ^{236}U concentrations varying within $(1-3) \times 10^7$ atom/L (Fig 4 (a) and (b)). The highest mean values of $^{236}\text{U}/^{238}\text{U}$ atomic ratio and ^{236}U concentration for both EG and WG are observed in samples from 2013 even when the two samples (2013-0537, 0538) with elevated ^{236}U levels are excluded in the calculation.

It is noteworthy that larger variation of the $^{233}\text{U}/^{236}\text{U}$ atomic ratio appears in 2013 compared to the other years (Fig 4(c)). The appearance of high values in ^{236}U concentration in Greenlandic surface seawater in the year 2013 might be attributed to various reasons including 1) existence of additional source terms; 2) redistribution / remobilization of ^{236}U via water overflow, mixing and/or ice melting 3) enrichment of ^{236}U via biogeochemical processes, e.g., redox reaction or association to fine particles (e.g., colloid, nanoparticles). This is discussed later combining the quantitative estimation of ^{236}U source terms of Greenland seawater using $^{233}\text{U}/^{236}\text{U}$ atomic ratios.

Arctic climate change affects the distribution of radioactivity in the Arctic marine environment and the pathway of radionuclides (Karcher et al., 2010). Due to the high temperature in July 2012, strong melting across almost the entire surface of the Greenland ice sheet has been reported (Nghiem et al., 2012), as can be seen from the relatively low salinities in samples from 2012 (average 31.71 ± 1.86 ‰) compared to other years (Tables S1-S4). The ice melting might somehow dilute ^{236}U in the coastal seawater in 2012, leading to relatively lower ^{236}U concentrations in the measured samples for 2012 compared to those for 2013.

3.2.2 Spatial distribution.

The ^{236}U concentration ($(1.17 \pm 0.13) \times 10^7$ atom/L) in the seawater from the Faroe Islands (2016-0474) is comparable to the majority of the Greenland samples, which is in good agreement with previous findings for other radionuclides (Dahlgaard et al., 2004). Within our measurement uncertainties, no

206 significant difference in ^{236}U concentrations (as well as $^{236}\text{U}/^{238}\text{U}$ atomic ratios) between EG and WG
207 (Fig. 4) is obtained, which is supported by the statistical data analysis (VAR-3 (Vestergaard, 1964)). This
208 is in contrast to the distribution pattern of other radionuclides (e.g., ^{90}Sr , ^{137}Cs) reported earlier (Aarkrog
209 et al., 1999, 2000; Dahlggaard et al., 2004), where higher levels show up more often in the EG compared to
210 WG because radioactive discharges from SF and LH transported by Atlantic water firstly reach EG then
211 WG (Fig. 1 and Supplementary information).

212 Among the WG sampling stations, relatively high concentrations of ^{236}U are often observed at locations
213 around 64-67°N on west Greenland (e.g., samples of 2012-0535, 2013-0533, 2015-0508 and 2016-0511).
214 Upwelling phenomena have been reported at the west Greenland shelf between 64°N and 67°N (Pedersen
215 et al., 2005), which could potentially bring the subsurface water with higher ^{236}U concentration to the
216 surface water (Castrillejo et al., 2018). To better interpret the spatial distribution pattern of ^{236}U along the
217 Greenland coast, it is important to understand potential source terms of ^{236}U and ^{233}U in this region.

218 **3.3 Source terms of ^{236}U and ^{233}U around the Greenland coast**

219 The overall sources of ^{236}U and ^{233}U in the Greenland coast seawater can be generally categorized in
220 two groups: 1) Global fallout (GF), including direct deposition of fallout from atmospheric weapons
221 testing into the ocean and indirect input via advective ocean water movement or via fresh water input
222 transporting GF signal from terrestrial area. 2) Reactor input (RT), including both discharges from
223 reprocessing plants (e.g., SF and LH) and regulatory/accidental releases from nuclear facilities (e.g. local
224 input in the Arctic).

225 **3.3.1 Global fallout.**

226 The distribution of global fallout ^{236}U is dominating (70 %) in the Northern hemisphere and variable
227 with latitude (Aoyama et al., 2006). A $^{236}\text{U}/^{238}\text{U}$ atomic ratio of about 1×10^{-9} (corresponding to 0.8×10^7
228 atom/L when apply a typical ^{238}U concentration of 3.3 $\mu\text{g/L}$) to has been estimated for the global fallout
229 signature in modern ocean surface waters (Casacuberta et al., 2016; Christl et al., 2012). This agrees with
230 published measurement (0.85×10^7 atom/L) on Northwest Pacific surface seawater (in 2014) where the
231 global fallout is the dominating source (Nomura et al., 2017a). Our results on average ^{236}U concentrations

232 $((1.5-3.8) \times 10^7$ atom/L) over 2012-2016 are 2-4 times higher than the global fallout value, indicating
233 additional input from other sources beside global fallout.

234 As mentioned earlier that ^{233}U can only be produced during nuclear weapons testing, therefore, global
235 fallout is the major source of ^{233}U in the Greenland coast. The average $^{233}\text{U}/^{236}\text{U}$ atomic ratio for global
236 fallout was obtained as $(1.40 \pm 0.12) \times 10^{-2}$ (Hain et al., 2020) and we believe this value would stay
237 constant in the open ocean with the termination of nuclear weapons test since 1980s. Therefore, any
238 actual $^{233}\text{U}/^{236}\text{U}$ atomic ratios below $(1.40 \pm 0.12) \times 10^{-2}$ for samples in the Greenland coast would
239 indicate the admixing of reactor-associated ^{236}U in the marine system.

240 Global fallout ^{236}U and ^{233}U deposited on the catchment of Greenland and the ice sheet can also be
241 transferred to the Greenlandic marine environment through freshwater input via river runoff and ice sheet
242 melting (Aarkrog, 1994; Dickson et al., 2007). The Greenland ice sheet is the largest freshwater reservoir
243 in the northern hemisphere with estimated annual fresh water flux in the range of 0.3-0.7 km³/y (Dickson
244 et al., 2007). Wendel et al. (Wendel et al., 2013) have reported $^{236}\text{U}/^{238}\text{U}$ atomic ratios of $(4 -100) \times 10^{-6}$ in
245 an Arctic ice core. Lower salinity samples (< 30 ‰) measured in this work seem not associate to high
246 ^{236}U concentrations (Fig. S1), indicating the indirect transport of global fallout ^{236}U (and ^{233}U) via
247 river/melt water is not significant.

248 **3.3.2 Reactor input**

249 **Reprocessing plants of SF and LH.** Most of the radioactive emissions from LH were transported
250 through the English Channel to the North Sea, while those from SF flow mostly north around the Scottish
251 coastline and then into the North Sea. The newly reconstructed ^{236}U release history from SF and LH show
252 a general decreasing trend since 1970s (Fig. S2) (Castrillejo et al., 2020). The $^{236}\text{U}/^{238}\text{U}$ atomic ratios
253 obtained in this work are several orders lower compared to those reported for the open North Sea, but
254 higher than other open oceans which mainly receive ^{236}U from global fallout, such as Equatorial Atlantic
255 Ocean, Sea of Japan and Northwest Pacific Ocean (Casacuberta et al., 2014; Nomura et al., 2017b;

256 Sakaguchi et al., 2012). This is in accordance with our expectation that discharges from the two major
257 European nuclear reprocessing plants are important sources of ^{236}U in Greenlandic seawater.

258 From the North Sea, the reprocessing ^{236}U can be transported to the Greenland coast by two ways: 1)
259 northward transport via the Norwegian Coastal Current (NCC) and partly the Norwegian Atlantic Current
260 (NAC) to Arctic Ocean and back from the Arctic Ocean via EGC; 2) The second branch of the NAC
261 turning back at Fram Strait and westward to Greenland merging into EGC (Raisbeck et al., 1995). These
262 two branches are further sub-divided into several loops in the Arctic ocean with different transit times
263 (Karcher et al., 2004; Smith et al., 2011).

264 **Local input.** A number of potential local sources for anthropogenic radionuclides in the Arctic region
265 have been reported, including radioactive waste dumping and submarine accidents (e.g., Komsomolets)
266 from USSR (Nies et al., 1999, 1998; Yablokov, 2001), close-in fallout from Novaya Zemlya nuclear
267 weapon test site (Aarkrog, 1994; Nies et al., 1998), Siberian river discharge and other operational
268 emission (Aarkrog, 1994; Karcher et al., 2010), nuclear powered satellite (Cosmos 954) accident (Grasty,
269 1995; Taylor et al., 1979; U.S Department of Energy, 1978), normal or accidental releases from Chalk
270 River labs (Cross, 1980) and contaminants from the Chernobyl accident (Davidson et al., 1987).
271 However, in terms of ^{236}U and ^{233}U inventories in these sources, nearly no documentation is available.
272 The only report for Cosmos 954 indicated that the nuclear reactor aboard was estimated to contain about
273 50 kg of highly enriched ^{235}U (U.S Department of Energy, 1978), which would have potentially produced
274 ^{236}U via the (n,γ) -reaction. However, due to limitations in the available measurement techniques applied in
275 radiological surveys after the accident, no data on ^{236}U concentration have been reported. Therefore, it is
276 apparent that further investigation is necessary to clarify all the above-mentioned local source terms in the
277 Arctic.

278 **3.4 Application of the $^{233}\text{U}/^{236}\text{U}$ ratio and binary mixing model for ^{236}U source**
279 **identification**

280 All measured $^{236}\text{U}/^{238}\text{U}$ and $^{233}\text{U}/^{236}\text{U}$ atomic ratios are plotted in a diagram (Fig. 5) demonstrating the
 281 binary mixing of the direct global fallout (DGF) and reprocessing plants of SF and LH (RP). The detailed
 282 parameters of the end-members are summarized in Table S5.

283 It can be seen from Fig. 5 that most of our observation data are below the binary mixing line of DGF
 284 and RP, indicating the existence of an extra endmember featured with relative low atomic ratios for both
 285 $^{233}\text{U}/^{236}\text{U}$ and $^{236}\text{U}/^{238}\text{U}$. This endmember in the mixing diagram is mostly likely natural ocean water
 286 (NOW) containing neither anthropogenic ^{236}U nor ^{233}U . The deviation from the mixing line gives the
 287 dilution of water from the two endmembers DGF and LH by the endmember NOW. In principle, any
 288 water less affected by anthropogenic U (and essentially any deep water) will lead to this dilution.

289 3.5 Quantitative estimation of the ^{236}U source term based on $^{233}\text{U}/^{236}\text{U}$ atomic ratios

290 A preliminary estimation is made to quantitatively evaluate the ^{236}U contribution from GF and RT as
 291 above-defined, based on $^{233}\text{U}/^{236}\text{U}$ atomic ratios using the following equation.

$$R_s = \frac{N_{233,f} + N_{233,r}}{N_{236,f} + N_{236,r}} = \frac{N_{236,f} \cdot R_f + N_{236,r} \cdot R_r}{N_{236,f} + N_{236,r}} = \frac{N_{233,f}/N_{236,r} \cdot R_f + R_r}{N_{236,f}/N_{236,r} + 1} \quad (1)$$

292 Where R_s , R_f and R_r respectively represent the $^{233}\text{U}/^{236}\text{U}$ atomic ratio of the Greenlandic seawater,
 293 global fallout and nuclear reactor; $N_{233,f}$ and $N_{233,r}$ refer to the atomic number of ^{233}U from global fallout
 294 and reactor, respectively; $N_{236,f}$ and $N_{236,r}$ refer to the atomic number of ^{236}U from global fallout and
 295 reactor, respectively. Therefore, the proportion (P_f) of GF can be obtained using equation (2).

$$P_f, \% = \frac{N_{236,f}}{N_{236,r} + N_{236,f}} = \frac{R_s - R_r}{R_f - R_r} \quad (2)$$

296 Assuming $R_r = 1.4 \times 10^{-2}$ and $R_f = 1 \times 10^{-6}$, the calculation results in Table 1 indicate that ^{236}U in
 297 Greenland seawater consists of $(29 \pm 5) \%$ from GF and $(29 \pm 9) \%$ from RT in both EG and WG. The GF
 298 contributions obtained here are comparable to the values calculated using representative global fallout
 299 ^{236}U concentration of 0.8×10^7 atom/L divided by the mean ^{236}U concentration measured in this work,

300 which are (32 ± 12) % for EG and (26 ± 7) % for WG, respectively. This indicates high proportion (about
301 70 %) of ^{236}U in Greenlandic surface seawater is associated to a reactor signal. A semi-quantitative
302 estimation based on transfer time and transit time (see detailed calculation in supplementary information)
303 shows that the contribution from RP is (49 ± 21) % for EG and (33 ± 4) % for WG during 2011-2016, the
304 remaining reactor contribution (e.g., local input) is obtained as (7 ± 15) % for EG and (38 ± 14) % for
305 WG (Table S5). To better quantify the contribution of reactor ^{236}U from RP and local input from the
306 Arctic, a more realistic oceanic model (e.g., transit time distribution (TTD)) is necessary (Smith et al.,
307 2011).

308 We further calculated the ^{236}U concentration associated to a reactor signal for each sample by
309 subtracting the global fallout contribution and plotted the residue (i.e., reactor ^{236}U) along the longitude
310 belt in Fig. 6. A higher reactor ^{236}U signal is observed in 2013, compared to the years before and after. It
311 seems unlikely that this additional input is related to the variation of SF and LH discharges (Fig. S2),
312 because this higher ^{236}U level is only observed within in narrow latitude/longitude belt ($61\text{-}65^\circ\text{N}$, $36\text{-}40$
313 $^\circ\text{W}$). It might be connected to local changes in ocean current branches, which potentially brought more
314 SF and LH water to the southeast of Greenland in 2013. The much higher ^{236}U level in sample 2013-0537
315 seems to be an exception and is difficult to explain in an oceanographic context. To reveal the origin and
316 the transport pathway of this additional reactor ^{236}U input further investigations are needed.

317 CONCLUSIONS

318 A first dataset of ^{236}U and ^{233}U in the Greenland marine environment during the 5-year period (2012-
319 2016) is reported. ^{236}U concentrations obtained for the Greenland surface seawater are distributed within a
320 relatively narrow range, with average ^{236}U concentrations being 2-4 times higher than the estimated value
321 from direct global fallout. Contrary to the spatial distribution pattern of other radionuclides (e.g., ^{137}Cs
322 and ^{90}Sr) reported earlier, we do not observe a significant difference in ^{236}U concentration between the
323 east and west Greenland coast, which is supported by the statistical data analysis (VAR-3). Application of
324 a binary mixing model to the correlation between $^{233}\text{U}/^{236}\text{U}$ and $^{236}\text{U}/^{238}\text{U}$ atomic ratios indicates that ^{236}U
325 contributed from the direct global fallout and the SF and LH reprocessing plants is diluted by a third

326 endmember (mostly likely natural ocean seawater) with marginal anthropogenic ^{236}U and ^{233}U . A
327 preliminary estimation of the ^{236}U source term composition using $^{233}\text{U}/^{236}\text{U}$ atomic ratios indicate that
328 global fallout contributes only about 30 %, while the majority (about 70 %) of ^{236}U is associated to
329 nuclear reactor signal in Greenland coastal water. Further investigation with focus on the analysis of ^{233}U
330 and ^{236}U in deep Greenlandic seawater is necessary to clarify the transport pathway and further use ^{236}U -
331 ^{233}U as paired-tracer system to identify different water masses.

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Table 1. Estimation of ^{236}U source composition in Greenlandic seawater by $^{233}\text{U}/^{236}\text{U}$ ratios

Year	Average $^{233}\text{U}/^{236}\text{U}$ atomic ratio, $\times 10^{-2}$		Contribution of GF, %*		Contribution of RT, %	
	EG	WG	EG	WG	EG	WG
2012	0.37 \pm 0.07	NA	26 \pm 4	NA	74 \pm 4	NA
2013	0.42 \pm 0.38	0.33 \pm 0.14	30 \pm 3	24 \pm 5	70 \pm 3	76 \pm 5
2015	0.50 \pm 0.09	0.35 \pm 0.10	36 \pm 2	25 \pm 5	64 \pm 2	75 \pm 5
2016	0.33 \pm 0.05	0.56 \pm 0.11	23 \pm 5	40 \pm 2	77 \pm 5	60 \pm 2
Mean \pm sd			29 \pm 5	29 \pm 9	71 \pm 5	71 \pm 9

*The value was calculated based on equation (2), with $R_r = 1.4 \times 10^{-2}$ and $R_r = 1 \times 10^{-6}$.

Figure 1. Map of Greenland coast showing the sampling locations (a) and general pattern of surface water currents (b). Blue arrows represent cold water currents, red arrows represent warm water currents and green arrows represent coastal water currents. NAC-Norwegian Atlantic Current, NCC-Norwegian Coastal Current, EGC-East Greenland Current, WSC-West Spitsbergen Current, WGC-Western Greenland Current, LH-La Hague nuclear reprocessing plant, SF-Sellafield nuclear reprocessing plant, FS-Faroe islands.

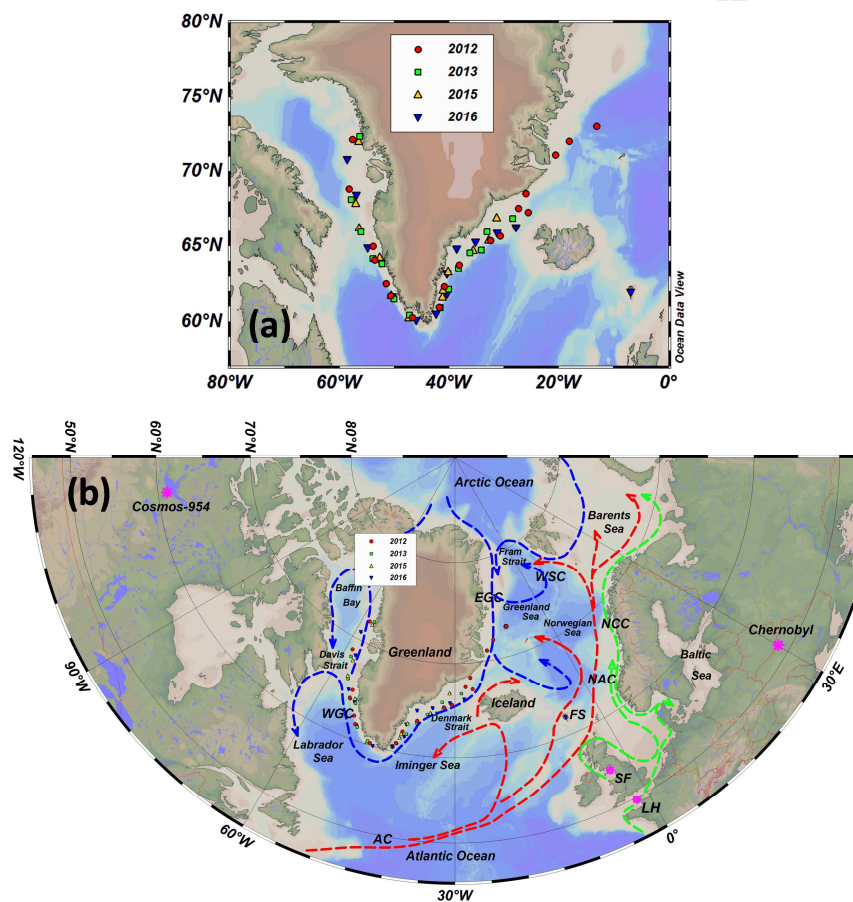


Figure 2. Time evolution and spatial distribution of ^{236}U concentration (left) and $^{236}\text{U}/^{238}\text{U}$ atomic ratio (right) in seawater around Greenland coast during 2012-2016. (Ref. a: Casacuberta et al. 2016; Ref. b: Casacuberta et al. 2018; Ref. c: Wefing et al. 2019)

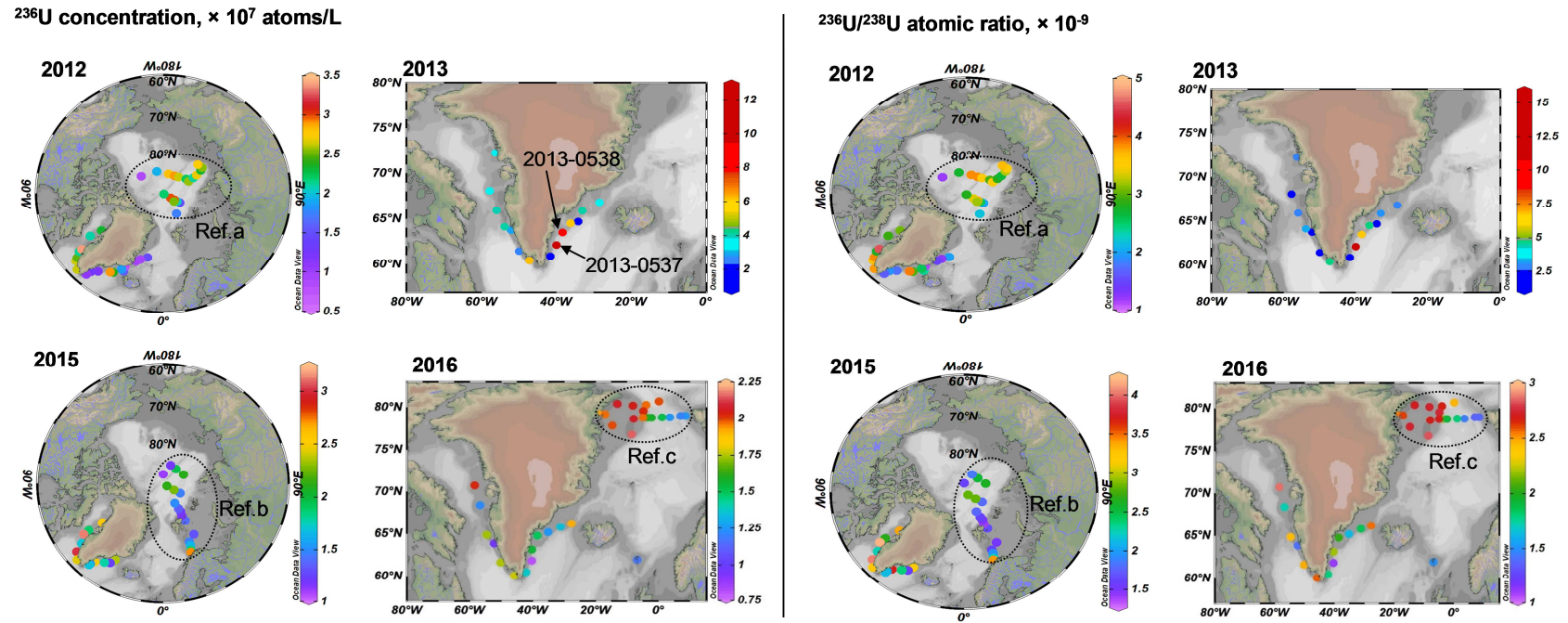


Figure 3. Time evolution and spatial distribution of $^{233}\text{U}/^{236}\text{U}$ atomic ratio in seawater around Greenland coast during 2012-2016.

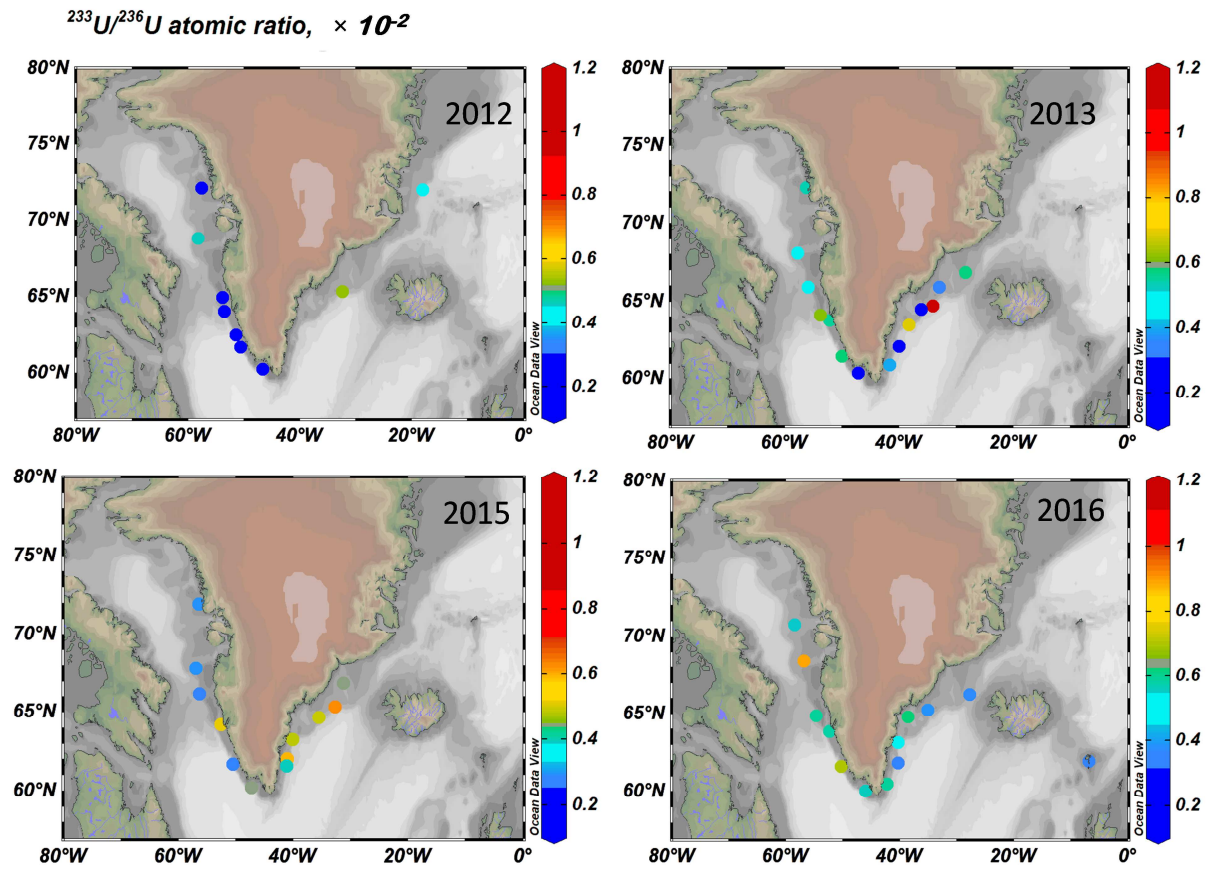


Figure. 4 Box-plots of yearly (a) $^{236}\text{U}/^{238}\text{U}$ atomic ratio, (b) ^{236}U concentration and (c) $^{233}\text{U}/^{236}\text{U}$ atomic ratio for samples from Eastern (E) and Western Greenland (W) coast (^{236}U results for sample 2013-0537 and 2013-538 are excluded in the plots). The red cross represents the arithmetic mean value of all the data for a given year, the whiskers extend from minimum (MIN) to maximum (MAX), the green box comprises of first quartile Q1 (top line), median (middle line) and third quartile Q3 (bottom line).

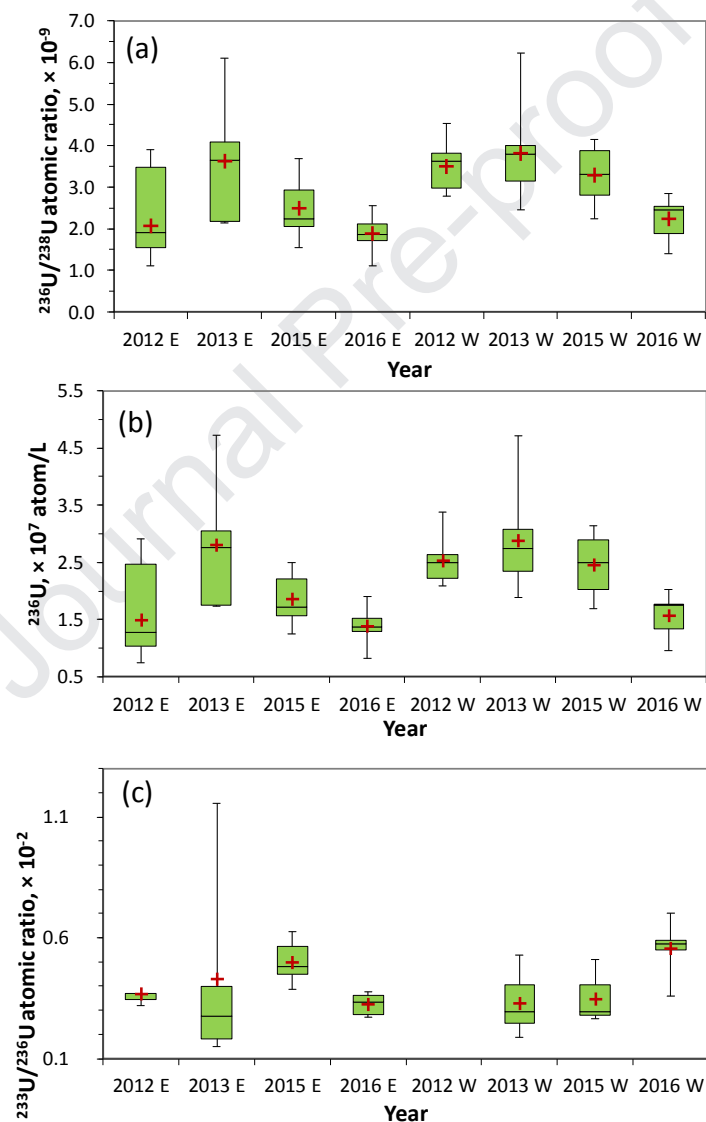


Figure. 5 Ternary mixing diagram of $^{233}\text{U}/^{236}\text{U}$ and $^{236}\text{U}/^{238}\text{U}$ atomic ratios for Greenland seawater (2012-2016).

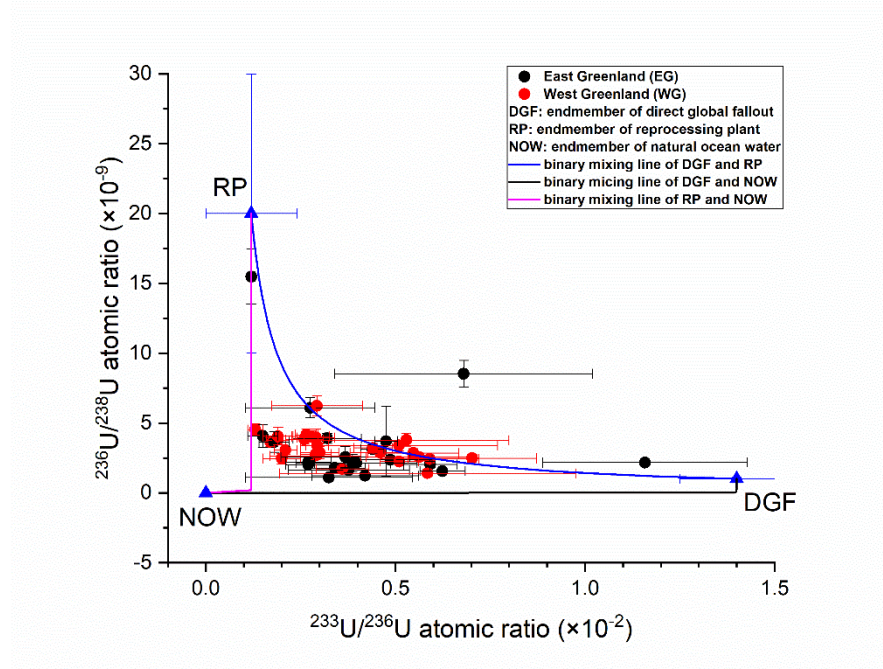
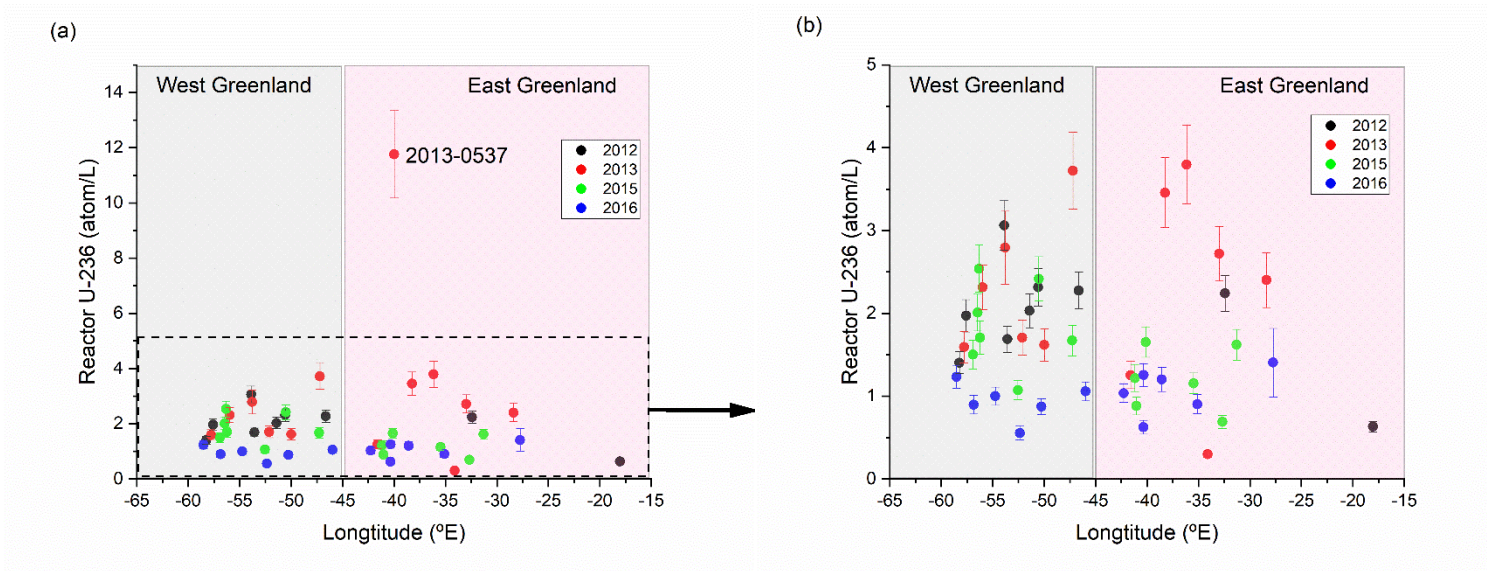


Figure. 6 Variation of calculated concentration of reactor ^{236}U for Greenland seawater (2012-2016) with longitude.



Highlights

- First results of ^{236}U and ^{233}U in the Greenland marine environment.
- ^{236}U concentration in Greenland seawater has a narrow distribution range.
- No significant difference in ^{236}U level between east and west Greenland coast.
- Binary mixing using $^{233}\text{U}/^{236}\text{U}$ and $^{236}\text{U}/^{238}\text{U}$ for interpret ^{236}U source term.
- ^{236}U source contribution is calculated based on $^{233}\text{U}/^{236}\text{U}$ atomic ratios.

Author contribution statement

Jixin Qiao: Conceptualization, Formal analysis, Methodology, Investigation, Resources, Data curation, Writing- Original draft preparation.

Karin Hain: Investigation, Methodology, Data Curation, Resources, Writing- Reviewing and Editing.

Peter Steier: Investigation, Methodology, Data Curation, Resources, Writing- Reviewing and Editing.

Journal Pre-proof

Declaration of interests

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: