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# Mercury Levels in Tissues (Cartilage, Skin, and Muscle) of the Greenland Shark (*Somniosus microcephalus*): Potential Contamination Sources and Implications for Health and Conservation

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## Abstract :

The jaws of the Greenland shark have high levels of mercury. Hg of cartilage in comparison with skin and muscle from the same specimen makes it possible to apprehend the distribution of the pollutant in the body. The level of the pollutant between jaw, skin and fresh meat (muscle) shows a strong correlation. The muscle is the most contaminated element in comparison with the skin and cartilage. The species presents the highest levels among different groups of sharks and the results are in accordance with previous studies. Marine ecosystems in the Arctic are globally contaminated by mercury (atmosphere, water, sediments, food web). The methylmercury reduces blood calcium levels, directly affecting the metabolism of cartilage cells. Even if cases of malformations could not be observed in the Greenland shark, numerous cases for other shark species have been documented in connection with heavy metals (e.g., Hg), and in particular for aplacental viviparous with potential morphological anomalies on embryos. The situation of the Greenland shark is worrying due to the conservation status, the fishing catches, the tardive sexual maturity and reproduction, the climate change and the level of mercury on its organism. The results incite to recommend ecological, environmental and fisheries management measures.

**Keywords** : Greenland shark, Mercury, Potential abnormalities, Conservation, Arctic

28        **1. Introduction**

29

30    The situation of the Greenland shark is worrying. The species holds a conservation classification of Near  
31    Threatened (NT) in Europe (Burgess et al. 2015). However, the reported catches by the Icelandic and  
32    Greenlandic fleets surged to more than 43 tonnes in 2010 and 120 tonnes in 2020 (Tonnes Live Weight; FAO  
33    areas 14, 14b, 14b2, 5a, 5a1, 5a2) (ICES, 2022).

34    The Greenland shark, inhabiting the cold waters of the Arctic and North Atlantic Oceans, including the Celtic  
35    and North Seas, faces bycatch mainly in Arctic regions, notably by halibut or shrimp fisheries (David et al.,

36 2013). Greenland Shark bycatch ranged from 34.4% for longline, 32.8% for trawl and 18.3% for gillnet  
37 respectively in NAFO subarea 0 (Bryk et al., 2018).

38 The Greenland shark displays aplacental viviparity (Nielsen et al., 2020). Genetic investigations have identified  
39 genetically mixed individuals in the Canadian Arctic, sub-Arctic, and temperate Eastern Atlantic areas,  
40 suggesting introgressive hybridization with the Pacific sleeper shark (*Somniosus pacificus*) Bigelow and  
41 Schroeder, 1944 (Walter et al., 2017). Insights into Greenland shark sexual maturity showed male and female  
42 body lengths-at-maturity (Total Length) of  $2.84\pm 0.06$  m and  $4.19\pm 0.04$  m, respectively (Nielsen et al. 2020). The  
43 species with a large long-live are particularly vulnerable to exposure to mercury biomagnified (Rumbold et al  
44 2014). Several scientific studies confirm the climate change in the Arctic and the polluted environment on  
45 mercury included food webs with potential effects of shark health (Braune et al., 2015; MacMeans, 2015;  
46 Rodríguez-Romero et al., 2018; Schaefer et al., 2020).

47 Natural phenomena such as volcanoes, rock erosion, but also forest fires, whether from natural, accidental or  
48 human causes, without forgetting industrial activities, especially the exploitation of coal and gold, emit mercury  
49 (Hg) into the environment (Veiga et al., 2006; Dabrowski et al., 2008; Witt et al., 2009). Mercury has the  
50 particularity of lasting up to two years in the atmosphere before the abiotic oxidation into  $Hg^{2+}$  (Ariya et al.,  
51 2015). Hence, the Arctic Ocean is susceptible to mercury contamination, which may stem from sources that are  
52 notably remote (Ariya et al., 2004).

53

54 Industrial activities associated with resource exploitation in the Arctic, including gold mining, have continued to  
55 expand in recent years (Mered, 2020). Gold processing involves the use of mercury and heavy metal pollution of  
56 water (Fashola et al., 2016). Kirk et al. 2008 demonstrated the methylation of Hg through analyses of Arctic  
57 waters. Concentrations of methylated Hg and Dimethylmercury (Me<sub>2</sub>Hg) were often low in surface waters  
58 ( $23.8\pm 9.9$  and  $4.7\pm 4.4$  pg L<sup>-1</sup>, respectively), but they increased with depth (maximum: 178 and 170 pg L<sup>-1</sup>,  
59 respectively; mean:  $70.3 \pm 37.3$  and  $56.8 \pm 37.8$  pg L<sup>-1</sup>, respectively). Water serves as a significant source of  
60 gaseous Hg (gaseous elemental Hg, GEM) in the atmosphere, especially during the ice-free period (Braune et al.,  
61 2015). Methylmercury (MeHg) by photodegradation phenomenon and Me<sub>2</sub>Hg on the surface can contribute to  
62 producing GEM deep in the water column (Mason et al. 1998; Chen et al 2003). Subsequently, it undergoes  
63 transformation or is reduced to elementary Hg and released back into the atmosphere. Another portion settles as  
64 sediments, while yet another part transforms into monomethylmercury (MeHg) and dimethylmercury (Me<sub>2</sub>Hg)  
65 (Braune et al., 2015). The deposition of land-based mercury on marine sediments causes increased microbial Hg

66 methylation and the transfer of MeHg to benthic and pelagic food webs (Ferreira Araujo et al., 2022). The  
67 contribution of mercury stems, in part, from the potential release of mercury from permafrost due to the erosion  
68 of rivers and coastlines resulting from Arctic warming (Schaefer et al., 2020).

69

70 The marine environment is not exempt from this concern. Different mercury forms are present, including  
71 elemental (Hg<sup>0</sup>), inorganic (Hg<sup>2+</sup>), and organic (CH<sub>3</sub>Hg<sup>+</sup>) forms (Clarkson, 1997). Among these,  
72 methylmercury (CH<sub>3</sub>Hg<sup>+</sup>), the hazardous variant of mercury, gradually accumulates within organisms of marine  
73 species including sharks and undergoes an amplification process (known as bioaccumulation) as it moves up the  
74 trophic web, from lower to higher levels (Harding et al., 2018, Biton-Porsmoguer et al., 2018; 2022). This  
75 susceptibility is linked to their extended lifespans and high trophic level (Gelsleichter and Walker 2010), which  
76 can lead to potential morphological abnormalities (Casarini et al., 1997; Rosa et al., 2004; Moore, 2015;  
77 Cabanillas-Torpoco et al., 2023).

78 The Greenland shark exhibits elevated mercury levels, among the highest among marine species analysed in the  
79 Canadian Arctic (Chételat and Braune, 2012). The presence of mercury has been demonstrated previously in  
80 different organs of the Greenland shark (only in muscle and liver) (McMeans et al., 2007, 2010, 2015; Chételat  
81 and Braune, 2012; Corsolini et al., 2014). The biological characteristics of the Greenland shark mentioned  
82 above, including longevity, make it a preferential candidate for study the level of contaminating elements in their  
83 organism. Further investigations of mercury production have a particular importance in marine arctic  
84 environment particularly in zones, which enhance uptake in the food web to evaluate source-receptor impacts  
85 (AMAP, 2021). The objective of this study are 1) to measure and confirm mercury concentration within the  
86 cartilaginous tissue of the species, specifically targeting the upper and lower jaws; 2) to analyse and compare  
87 mercury levels present in skin and muscle samples extracted from the same individual specimens; 3) to examine  
88 potential sources of mercury contamination and investigate its potential effects on the morphology and  
89 reproductive patterns of sharks; 4) to provide recommendations for conservation strategies aimed at reducing  
90 bycatch and minimizing the presence of mercury in the delicate Arctic polar ecosystem. In accomplishing these  
91 objectives, the study aims to contribute valuable insights into mercury accumulation dynamics, potential  
92 consequences for the species, and imperative conservation actions required in the Arctic environment.

93

## 94 **2. Materials and methods**

95

96 *Sampling*

97 The sampling phase was conducted aboard the bottom trawler "Steinunn SF-10." The vessel operated from  
98 Hornafjörður, Iceland, during the year 2010. On August 8, 2010, a female Greenland shark was captured in the  
99 Greenland Sea, near Denmark Strait (68°016 N - 22°764 W) within FAO fishing area 5.2 (Fig.1). Measuring 434  
100 cm in total length, the specimen was procured and retained on the ship. Samples of both upper and lower jaws  
101 muscle and skin were meticulously extracted using knives and scalpels (Fig.1). The tools underwent thorough  
102 cleaning with soapy water, disinfection using 90% ethanol, and subsequent rinsing with distilled water between  
103 each sample collection. After extraction, all samples were placed within hermetically sealed plastic bags and  
104 accurately referenced. Subsequently, the muscle sample was immersed in 70% ethanol, while the jaw  
105 (cartilaginous tissue) and skin samples were dried and were securely stored in airtight plastic bags.

106

107 *Mercury analysis*

108 The total mercury (THg) concentration of the tissue samples was determined using cold vapor atomic absorption  
109 spectroscopy (CVAAS) with thermal decomposition and gold amalgamation. The instrument used was the Hydra  
110 IIC direct mercury analyzer by Teledyne Leeman Labs (Hudson, NH). Calibration curves were constructed using  
111 the Certified Reference Materials (CRM) BCR 414 (phytoplankton), DORM-4 (fish protein), EM-CE278K  
112 (mussels) EMR-CE464 (Tuna fish) ranging from 0.276 mg.kg<sup>-1</sup> to 5.240 mg. kg<sup>-1</sup> dw of THg. Instrument  
113 linearity was established across two separate ranges; a low range (0 ng to 30 ng) and high range (75 ng to 200  
114 ng). The high calibration range has a correlation coefficient ( $r^2$ ) value of 0.9992 and the low calibration range  
115 has an  $r^2$  value of 0.9996. The limit of quantitation (LOQ) was set as 10 times the standard deviation of the y-  
116 axis intercept of the method calibration curve.

117 A portion of each tissue sample was placed in a nickel boat and weighed on an analytical balance ( $\pm 0.0001$  g).  
118 The amounts of upper and lower jaws, skin and muscle were 0.0127 g dry mass, 0.0175 g dry mass, 0.0213 g dry  
119 mass and 0.0142 $\pm$ 0.004 g dry mass, respectively. Given the unconventional method of preserving muscle in  
120 alcohol for THg analysis and the sufficient sample quantity available for analysis, a total of three separate  
121 analyses were conducted. The provided result is the average of the three measurements. Samples were first  
122 introduced into the decomposition furnace where they were dried (300 °C, 30 s) and combusted (800 °C, 150 s),  
123 then into the catalyst furnace (600 °C, 60 s), and finally through the drying tube and gold amalgamation trap  
124 (600 °C, 30 s) before entering the spectrometer. Samples were not homogenized to avoid unnecessary  
125 contamination from additional handling and due to the limited available tissue in some cases. Samples were not  
126 digested before analysis. For quality assurance, a CRM (EMR-CE464 Tuna Fish) measurement was performed at  
127 the end of the six analyses.

128

### 129 **3. Results and discussion**

130

131 Elevated THg levels were analysed in all tissues of the shark's body, ranging from 1.332 mg kg<sup>-1</sup> dw (upper  
132 jaw) to 3.225 mg kg<sup>-1</sup> dw (muscle) (Fig.2; Table 1). A linear concentration increase trend is confirmed in THg  
133 between the lower and upper jaws (cartilage), skin and fresh meat (muscle) ( $R^2=0.9996$ ) (Fig.2).

134 The distribution of mercury levels across different shark tissues reveals that muscle accumulates higher levels  
135 compared to the skin and cartilage. This disparity may be attributed to variations in the rate of cell turnover,  
136 especially notable in cartilage (Marconi et al. 2020) and skin (Meyer and Seegers, 2012), but also with the fact  
137 that mercury in fish included sharks is mostly in the methylated form and has a protein affinity (Chouvelon et al.,

138 2018). The mercury levels analysed in this study are consistent with those observed from other specimens of  
139 *Somniosus microcephalus* (Table 1). However, it appears that the varying mercury levels obtained are not  
140 correlated with the length of the individuals ( $P>0.05$ ,  $n=19$ , MacMeans et al., 2010). A multiple test on muscle  
141 tissue indicated that concentrations of mercury were higher on Greenland shark,  $5.930 \text{ mg kg}^{-1}$ ,  $n = 19$   
142 (Macnean, 2017),  $3.225 \text{ mg kg}^{-1}$ ,  $n = 1$  (this study) in comparison with specimens of Carcharhiniformes,  $1.520 \pm$   
143  $1.900 \text{ mg kg}^{-1}$ ,  $n = 1739$ ; Lamniformes,  $2.580 \pm 4.790 \text{ mg kg}^{-1}$ ,  $n = 508$ ; others species of Squaliformes,  $1.610 \pm$   
144  $1.040 \text{ mg kg}^{-1}$ ,  $n = 415$ ; and Myliobatiformes,  $0.383 \pm 0.350 \text{ mg kg}^{-1}$ ,  $n = 195$  (Tiktak et al., 2020). Several  
145 factors may explain higher THg values in Greenland shark (i) their feeding at a high trophic position (TP) and  
146 (ii) foraging on slightly more contaminated offshore resources in the Arctic (MacMeans et al., 2010). The lack of  
147 a correlation between mercury levels and the age (size) of the sharks can be attributed to (i) mercury  
148 accumulation in larger, presumably older Greenland sharks not being solely driven by bioaccumulation and (ii)  
149 larger specimens not necessarily feeding at a higher TP than smaller sharks (MacMeans et al., 2010).

150

151 The limitation of the method used

152 The measurement of mercury in a sample preserved in ethanol, such as the muscle sample in this case, is not  
153 standard. Usually, measurements are carried out on fresh (MacMeans et al., 2010) or freeze-dried samples  
154 (Biton-Porsmoguer et al., 2018; 2022). Here, the measurements were conducted 13 years after the collection, and  
155 the sample preservation method could not be controlled. Alcohol preservation can be used for maintaining tissue  
156 samples for various analyses, including mercury measurements. However, certain factors need to be considered.  
157 Ethanol itself may contain trace amounts of mercury. Hence, it is crucial to use high-purity ethanol for  
158 preservation and ensure that the ethanol used does not introduce additional mercury into the samples.  
159 Here, ethanol (70% USP-grade Isopropanol (Isopropyl Alcohol) and 30% USP-grade purified water) was used. It  
160 is also reasonable to assume that mercury concentrations could change during preservation, likely due to the loss  
161 of mass caused by protein and water loss in samples preserved in ethanol. Hence Gibbs et al. (1974) observed a  
162 slight increase in mercury concentrations in fish after a short period of preservation in ethanol. However, when  
163 preserved in ethanol, fish samples typically experience an initial loss of mass due to deshydration, followed by  
164 a stabilisation (Shields and Carlson, 1996). In this case, the muscle sample was preserved in ethanol for 13 years.  
165 It is highly likely that any initial dehydration induced by the alcohol has reached a state of stabilization. Utilizing  
166 a conversion equation holds merit in the endeavour to retrocalculate the mass of ethanol-preserved samples to  
167 their initial state under fresh conditions. Nevertheless, it is important to note that these conversion equations tend

168 to possess limited reliability, typically capable of offering only coarse approximations of the live condition  
169 (Billy, 1982). Consequently, we have chosen to refrain from their application. We acknowledge that the  
170 concentrations measured in the muscle of the ethanol-preserved shark are prone to being higher than the actual  
171 concentrations determined based on fresh weight. Shields and Carlson's study (1996) indicates that this  
172 overestimation could potentially reach up to 20%. This uncertainty adds to the measurement uncertainties.

173 The selection of analytical methods for measuring mercury in alcohol-preserved samples could vary from those  
174 employed for fresh or frozen samples. Cold vapor atomic absorption spectrometry (CVAAS) has been previously  
175 employed for mercury analysis in alcohol-preserved samples (Levengood et al., 2013), thereby confirming the  
176 applicability of the method used in this study. The authors analysed the composition of the alcohol in the jars used  
177 for specimen storage. Mercury was detected at a very low concentration in the ethanol, confirming that there are  
178 minimal mercury transfers from the samples to the alcohol.

179

180 *Sources of mercury in arctic region for sharks*

181 Several studies have highlighted that prey serves as a primary source of mercury in marine organisms,  
182 particularly for top predators like sharks, owing to the phenomenon of bioaccumulation (McMeans et al., 2015;  
183 Biton-Porsmoguer et al., 2018; Li et al., 2023). The main prey of the Greenland shark are fish (mainly Teleosts  
184 and Elasmobranch as rays), mammals (seals, whales and bears), squids and crustaceans (shrimp and crabs)  
185 (McMeans et al., 2010; Nielsen et al. 2013; 2019). The species more contaminated by mercury (muscle) in  
186 Arctic were elasmobranch and mammals (as rays and seals, 1.2-2 THg  $\mu\text{g kg}^{-1}$  dw) and teleosts (as ling, blue  
187 ling, tusk, 0.12-0.94 THg  $\mu\text{g kg}^{-1}$  dw) (McMeans et al., 2010; 2015).

188 The results of this study demonstrate 1) the presence of mercury in the cartilage and 2) a strong correlation with  
189 the level of THg in cartilage with the skin and the muscle. Numerous studies have focused on the implications of  
190 mercury for human health (Ha et al., 2017). It is noteworthy that the flesh of the Greenland shark is consumed in  
191 Iceland, known as "Hákarl," and is subjected to a distinctive fermentation process followed by drying for a  
192 duration of four to five months (Travel Food Atlas, 2020). The maximum allowable mercury level for shark meat  
193 intended for human consumption is set at 1.0 mg/kg of fresh meat<sup>1</sup>. Considering the mercury levels analysed in  
194 this study, the shark flesh appears unsuitable for human consumption and could potentially pose a health risk.  
195 (Guzzi et al., 2021). Moreover, mercury can also have effect on sharks.

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<sup>1</sup> Commission Regulation (EU) 2022/617 of 12 April 2022 amending Regulation (EC) No 1881/2006.



196

197 *Potential effects of mercury on shark morphology and reproduction*

198 The Elasmobranchs (sharks, skates and chimaeras) are characterized by a cartilaginous skeleton (Seidel et al.  
199 2021). The vertebral centra of elasmobranchs are composed of calcified cartilage, mainly comprised of the  
200 calcium phosphate mineral hydroxyapatite [Ca<sub>10</sub>(PO<sub>4</sub>)<sub>6</sub>(OH)<sub>2</sub>], which is embedded within an organic matrix of  
201 proteins, including proteoglycan and collagen (Urist, 1961; Porter et al., 2006). Mercury leads to metabolic  
202 disorders and disrupts the proper assembly of the cytoskeleton in marine species. The toxicity of mercury also  
203 causes morphological alterations on the development various fish organs, including the gills and olfactory  
204 organs. Malformations can involve the craniofacial and skeletal systems with stunted growth, curved spine, and  
205 eye deformities (Weis and Weis, 1977; Weis et al., 1981; Jagoe et al., 1996; Ribeiro et al., 1996; Oliveira  
206 Ribeiro et al., 2000, Devlin, 2006; Adams et al., 2010).

207

208 The effects of mercury on bone metabolism have also been demonstrated for teleosts and in particular for the  
209 activity of osteoclasts (cell responsible for the dissolution and absorption of bone) and osteoblasts (cell which  
210 secretes the substance of bone) (Suzuki et al., 2004). The methylmercury reduces blood calcium levels  
211 (calcemia), directly affecting the metabolism of bone cells at the scale (Suzuki et al., 2004). Osteoclasts on bone  
212 and those on cartilage also have similar basic molecular characteristics (Larroure et al., 2021). Yachiguchi et  
213 al., (2014) reported a decrease in TRAP (Tartrate-Resistant Acid Phosphatase) and ALP (alkaline phosphatase)  
214 expression in marine fish exposed to Methylmercury or inorganic mercury. The ALP and TRAP are known as  
215 marker enzymes for osteoblasts and osteoclasts, respectively (Bonuci and Nanci 2001). Mercury would therefore  
216 inhibit the activity of both osteoclasts and osteoblasts (Rodríguez and Mandalunis 2018).

217 The observation of sharks during boarding and the study of the literature did not make it possible to identify  
218 malformations in Greenland shark. However, many observations of morphological abnormalities in sharks have  
219 been documented for many years. Thirteen studies have been published since 1963 reporting malformations on  
220 the blue shark (*Prionace glauca*) (Linnaeus, 1758) in different marine areas (North and South Atlantic Ocean,  
221 Pacific Ocean, California Gulf, Mediterranean Sea) (Cabanillas-Torpoco et al., 2023). Hoenig and Walsh (1983)  
222 noted four cases of vertebral lesions in three species of sharks, Sandbar shark *Carcharhinus plumbeus* (Nardo,  
223 1827, Limon shark *Negaprion brevirostris* (Poey, 1868) and Sand tiger shark *Carcharias Taurus* Rafinesque,  
224 1810). The spines had fused centra, ribs, and neural arches, additional deposition and erosion of calcified  
225 material in the centra, and in one case compression of the centra (Hoenig and Walsh 1983). Missing fins in an

226 adult tawny nurse shark, *Nebrius concolor* (Lesson, 1831) were reported by Taniuchi and Yanagisawa (1987),  
227 abnormal bicephalism in blue shark and tope shark (*Galeorhinus galeus*) (Linnaeus, 1758) by Ramirez-Amaro et  
228 al. (2019). The complete absence of pelvic fins in a milk shark (*Rhizoprionodon acutus*) (Rüppel, 1837) was  
229 reported in the Persian–Arabian Gulf (Moore 2015). The causes of such deformities were unknown.

230 The environmental degradation or pollution (effects of contaminants) can explain the malformation  
231 presence in sharks (Casarini, 1997; Rosa et al., 2004; Mancini et al., 2006). In this context, the presence of high  
232 levels of mercury (Hg) have been confirmed by several studies (Barrera-García et al., 2012; Rodríguez-Romero  
233 et al., 2018) and genetic alterations abnormalities in a blue shark embryo related with initial stages of  
234 development (Mancini et al., 2006; Rodríguez-Romero et al., 2018). Moreover, Van Hees and Ebert (2017) have  
235 demonstrated the maternal mercury transfer on the leopard shark *Triakis semifasciata* Girard, 1855 (an  
236 aplacental viviparous species as Greenland shark). Embryos were found with potentially harmful mercury  
237 concentrations in their muscle tissues.

238

239

#### 240 4. Conclusion

241

242 Marine ecosystems in the Arctic are globally contaminated by mercury (atmosphere, water, sediments,  
243 food web). Hg analyses performed on the specimen showed a contamination in different parts of his organism  
244 (cartilage, skin and muscle) with potential risks for the individual and human health. Recent studies  
245 demonstrated the maternal mercury transfer for aplacental viviparous and connected mercury with potential  
246 morphological anomalies on sharks. Arctic temperatures will increase more than the global average  
247 (Johannessen et al., 2004). Estimated aerobic scope for polar species will indicate their adaptive capacity and  
248 predictions of food web shifts (Steiner et al., 2019). Studies on mercury levels on elasmobranch should to be  
249 encouraged to analyse the state of embryos and better understand the causes of malformations. These results  
250 could be very interesting to list the species most affected and for their survival rate and the conservation of the  
251 species.

252 However, considering the substantial amount of data that corroborates the elevated mercury levels found across  
253 various tissues of the Greenland shark (including cartilage), as well as the observed morphological anomalies in  
254 numerous other species (raising suspicions of environmental pollution), coupled with the declining population in  
255 Europe (Burgess et al., 2015), shouldn't precautionary measures be implemented as a principle of safeguard?

256 In order to protect the environment, the precautionary approach shall be widely applied by States according to  
257 their capabilities. Where there are threats of serious or irreversible damage, lack of full scientific certainty,  
258 explained in this case in part by the difficulty of studying this Green land shark at different biological stages but  
259 also by the geographical area and sampling conditions, shall not be used as a reason for postponing cost-effective  
260 measures to prevent environmental degradation (Principle 15 of the 1992 Rio Declaration States) (United Nation,  
261 1992).

262 One of the objectives of this work was to provide recommendations for conservation strategies aimed at reducing  
263 bycatch, detailed by gear previously. The proposal measures could be, 1) Islandic and Greenland fleets must take  
264 the necessary measures to develop the turtle excluder devices and Nordmore grates and release specimen still  
265 alive; 2) Protect the central Arctic Ocean beyond national jurisdiction with a precautionary fisheries moratorium;  
266 3) Contribute effectively to the fight against global warming and reduce any industrial production that is a source  
267 of mercury (and any other pollutant) in the Arctic Ocean and 4) Initiate research programs to better understand  
268 the areas of movement and distribution of the species, which will inevitably be reduced by climate change and in  
269 order to limit any fishing activity there.

270

271

#### 272 **Declaration of competing interest**

273

274 The authors have no competing interests to declare that are relevant to the content of this article.

275

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277

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## 583 FIGURES CAPTIONS

584

585 Fig. 1. Position of the Greenland shark fished and sampled off Northwest Iceland marked by red cross. A. Skin.  
586 B. Upper jaw. C. Fresh meat. D. Lower jaw. Female. Total Length 434 cm. Greenland Sea. August 2010.

587

588 Fig.2. Hg level (mg kg<sup>-1</sup> dw) in selected tissues of the Greenland shark: cartilage (upper and lower jaw), skin and  
589 muscle (fresh meat).

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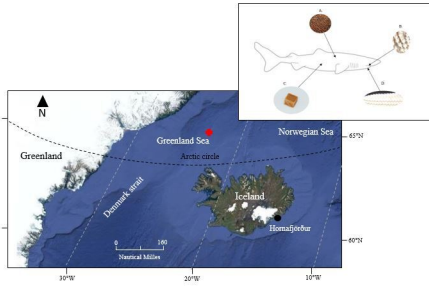
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 595 TABLES  
 596  
 597 Table 1. Range of total Hg levels ( $\text{mg kg}^{-1}$  ww) in the Greenland shark (*Somniosus microcephalus*) in the Arctic  
 598 and adjacent seas; - No data. dw: dry wet; ww: wet weight; dry weight (dw) concentration = 5 ww concentration  
 599 (Cresson et al., 2014).  
 600

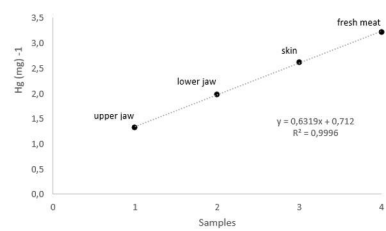
Regions	Greenland shark							References
	TL (cm)	upper jaw	lower jaw	muscle	skin	liver	ww/dw	
Greenland Sea	434	1.332	1.985	3.225	2.625	-	dw	<i>This study</i>
Cumberland Sound	236-325*	-	-	-	-	4.92±0.58	dw	McMeans et al. 2007
Islandic waters	415.6±25.2	-	-	5.93±0.59	-	5.32±2.73	dw	McMeans et al. 2010
Pangnirtung, Baffin Bay	293±1.1	-	-	1.715±0.457	-	-	ww	Chételat and Braune 2012
NE Greenland fjords	-	-	-	4.10-6.91	-	-	dw	Corsolini et al. 2014
Labrador Sea	273.3±31.5	-	-	3.54±1.02	-	-	dw	McMeans et al. 2015

601 TL = Total Length of individuals (cm); \*fork length (cm); Hg = Total Hg concentration ( $\text{mg kg}^{-1}$ )

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**Declaration of competing interest**

The authors have no competing interests to declare that are relevant to the content of this article.

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