Does the Fukushima NPP disaster affect the caesium activity of North Atlantic Ocean fish?

G. Kanisch and M.-O. Aust
Thünen Institute of Fisheries Ecology, Marckmannstraße 129b, 20539 Hamburg, Germany

Correspondence to: G. Kanisch (guenter.kanisch@ti.bund.de)

Received: 18 December 2012 – Published in Biogeosciences Discuss.: 5 March 2013
Revised: 30 June 2013 – Accepted: 8 July 2013 – Published: 13 August 2013

Abstract. Fillet samples of marine fish collected from the East/West Greenland currents (GC) and from the Baltic Sea (BS) have been investigated by gamma-ray spectrometry within the regular German monitoring programme. In samples of the second half of 2011, $^{134}$Cs traces have been detected that are suggested to originate from the Fukushima fallout that was deposited in March/April 2011 over the northern North Atlantic and accumulated by fish. The radionuclide $^{134}$Cs (half-life 2 yr) was indeed detected with quite small activities at about 0.0036 Bq kg$^{-1}$ w.w. Existing box models describing the transport of Cs within seawater boxes of the northeast Atlantic allowed for estimation of $^{134}$Cs contributions from other sources, i.e. from the Chernobyl fallout and from discharges by the two major European nuclear reprocessing plants; both were negligible around Greenland, while for the Chernobyl fallout a small $^{134}$Cs background contribution to BS fish was estimated. Model results confirmed the level of $^{134}$Cs measured in BS fish and showed its maximum to have occurred in winter 2011/2012 followed by a continuous decrease. It was also determined that $^{134}$Cs activity, but not that of $^{137}$Cs, showed a significant negative correlation with sampling depth (150–400 m) of GC fish; this strengthens our Fukushima fallout assumption. As a result, the Fukushima fallout in these sea areas only marginally enhanced (GC: 4%; BS: 0.1%) pre-Fukushima levels of individual dose rates received by human fish consumers; the addition was around 0.001 μSv following the consumption of 10 kg of fish per year, which is not expected to cause concern according to present guidelines for radiation protection.

1 Introduction

On 11 March 2011, a Tsunami hit the Fukushima Daiichi nuclear power plant (FD-NPP), which caused the loss of cooling capacity in four of its six nuclear reactors and led to the release of radionuclides into the environment. It was expected that between 6 and 47 PBq (1 PBq = $10^{15}$ Bq) of $^{137}$Cs (half-life 30.17 yr) was directly discharged into the Pacific Ocean (e.g. Bailly du Bois et al., 2012) in the aftermath of the tragedy. More recent publications using more detailed data estimated amounts between 4 and 6 PBq $^{137}$Cs leaked into the Pacific Ocean (Povinec et al., 2013; Miyazawa et al., 2013; Estournel et al., 2012). Due to the determined $^{134}$Cs/$^{137}$Cs ratio of around 1, about the same amount of $^{134}$Cs (half-life 2.07 yr) was discharged into the Pacific. Initially, discharge was assumed to represent the larger fraction of total Cs releases. Therefore, many researchers and TEPCO have focussed on the determination and estimation of the behaviour of Cs in Pacific waters and its behaviour in the environment, especially the uptake by biota (e.g. Buesseler et al., 2011, 2012; Honda et al., 2012; Madigan et al., 2012; Behrens et al., 2012).

The explosions of units 1 to 4 of FD-NPP also released radionuclides into the atmosphere, with an amount of about 12 to 15 PBq for each of $^{137}$Cs and $^{134}$Cs, respectively (Povinec et al., 2013; Estournel et al., 2012); this was detectable around the world (e.g. Hsu et al., 2012; Stohl et al. 2012; Jakobs, 2011). From these investigations the wet and dry deposition of caesium isotopes in marine areas where German monitoring was carried out after the accident at FD-NPP occurred – the North Sea, the Baltic Sea (BS), and the coastal West and East GCs – were estimated to be between 0.1 and 100 Bq m$^{-2}$.
The deposition of Fukushima fallout radionuclides occurred during the last week in March 2011 until about the beginning of May 2011. An average deposition of 2 Bq m\(^{-2}\) for each of \(^{134}\)Cs and \(^{137}\)Cs was measured at three locations in Denmark by Risø DTU (Nielsen, 2011). Furthermore, in Finland, depositions of 1.75 Bq m\(^{-2}\) for each of \(^{134}\)Cs and \(^{137}\)Cs was measured by STUK in Helsinki (Ikäheimonen, 2011) during 21 April and 14 May in 2011 using 1-week sample periods. From Greenland, to the best of our knowledge, no such determinations are available.

Putting FD-NPP deposition estimates and the resulting concentrations measured in Greenland into perspective, they are comparable with those of the Chernobyl accident in the Baltic Sea (BS), which was characterized by a mean deposition of 12000 Bq \(^{137}\)Cs m\(^{-2}\), and led to an activity concentration between 50 and 5200 Bq m\(^{-3}\) in seawater and up to 270 Bq \(^{137}\)Cs kg\(^{-1}\) wet weight (w.w.) in fish (HELCOM, 1995). The resulting concentration factors of up to 400 imply that fish is the most reasonable test organism for monitoring the input of caesium isotopes from FD-NPP into different oceans and the separation of FD-NPP from historic Cs inputs, namely through global fallout and the catastrophe at Chernobyl nuclear power plant.

From this point of view, the aims of our study were (i) to determine the activity concentration of caesium isotopes in cod and redfish as economically relevant fish species in the North Atlantic Ocean and its marginal seas, (ii) to separate the FD-NPP input from that from Chernobyl and global fallout and (iii) to validate the determinations and estimate the future behaviour of caesium isotopes in fish of the BS using the first-order kinetics box model previously developed within a HELCOM Working Group MORS-EG to model Chernobyl \(^{137}\)Cs in seawater and fish of the BS.

2 Materials and methods

2.1 Sampling and sample preparation

Biota were caught during research and monitoring cruises using the RV Walther Herwig III carried out between October 2010 and December 2011 using bottom trawls or collected from commercial catches from trawlers operating within ICES (International Council for the Exploration of the Sea) subdivisions in the BS. Sampling positions are indicated in Fig. 1. Round fish samples selected from the catches, consisting of 2 to 31 individual animals (Table 1), were stored at \(-20^\circ\)C after sampling of 5 kg small fish or 15 kg larger fish. Larger fishes (especially cod) were filleted on board before freezing. The dry mass of the samples was determined in the laboratory by drying at 110 \(^\circ\)C for 2 days. Afterwards, dried samples were dry-ashed at a maximum temperature of 420 \(^\circ\)C. The retrieved ash, which was between 50 and 70 g, was homogenized and transferred to 200 mL cylindrical PVC containers for gamma spectrometric analysis, resulting in variable filling heights and bulk densities of about 25–40 mm and 0.44–0.56 g cm\(^{-3}\), respectively. The fish samples

![Fig. 1. Sites for sampling of cod, redfish and whiting (see Table 1 for sample details) for analysis of radioactivity in the Labrador Basin and the BS (scale of the large map is 1 : 42,000,000; scale of the small map is 1 : 10,000,000). The small map of the BS contains the borders and nomenclature of the water boxes (numbers in boxes: 71 = Kattegat, surface; 75 = Belt Sea, surface; 81 = BS west, surface; 83 = BS east, surface) for estimation of the Cs behaviour using the modified HELCOM box model (see HELCOM, 1995, or Kanisch et al., 2000, for details).]
Table 1. Characterization of fish samples (cod: *Gadus morhua*; redfish: *Sebastes marinus*; whiting: *Merlangius merlangius*)

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<th>average fish length (cm)</th>
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selected for this investigation are characterized in Table 1. Fish age was not determined.

2.2 Gamma-ray spectrometric analysis

2.2.1 Spectrometry basics

For the analysis of gamma-emitting radionuclides three high-purity germanium detectors (2 × n-type; 1 × p-type) with relative efficiencies between 36 and 55% and energy resolutions of around 1.9 keV, both at 1.33 MeV. The detectors are protected against environmental radiation by standard low-level lead shieldings of 10 cm lead and inner layers of copper, cadmium and Plexiglas. Two multi-channel analyser (MCA) modules were used for recording the spectra with 4096 channels. The MCA spectra were transferred into an external spectrum database used by the GamW software for spectrum evaluation, which is an in-house development. Some details of the spectrum analysis techniques used in this program are given in Appendix A, in which also some measures for improving the detection of $^{134}$Cs are described. Radionuclide activity is always referred to in wet weight (w.w.).

The full-energy peak efficiency curves were re-calibrated in January 2011 using a liquid multi-radionuclide standard solution 7503ML (Eckert & Ziegler, Braunschweig, Germany; 11 radionuclides including $^{210}$Pb and $^{241}$Am) covering the energy range of 46 to 1836 keV; relative standard uncertainties of the activity concentrations were about 1%. Two-dimensional efficiency curves, depending on gamma energy and filling height in the container were fitted to the measured efficiency data for four heights between 12 and 56 mm). Calculations of true coincidence summing corrections (TCS) were included during efficiency curve fitting; see Appendix A for more information about TCS corrections.

The standard counting durations for screening on caesium isotopes was 160 h. When there was some evidence of the presence of $^{134}$Cs, the measurements of those samples were prolonged to 320–410 h for improving the counting statistics and lowering the detection limit. The ISO 11929 decision threshold was used as a criterion for the detection of $^{134}$Cs (see Appendix A for details).

2.2.2 Quality control

The stability of the detection efficiencies is controlled by counting every few weeks a $^{152}$Eu point source on the detectors. The detector background is measured two times a year by collecting two or three 1-week spectra; the peak net counting rates are evaluated statistically, also by comparing the results with those of the sum spectra obtained by summing up these 2–3 spectra per detector. For external quality control the lab participates in national (Federal Office for Radiation Protection (BfS), Berlin, Germany; water samples) as well as IAEA-EL intercomparison exercises with marine...
origin samples. The latest IAEA intercomparison test sample measured with the new calibration from January 2011 was a seaweed sample (code IAEA-446). Our results for mean activity concentrations (i.e. activities per unit mass) of $^{40}$K and $^{137}$Cs differed only 1.1 % from the preliminary IAEA reference values (under evaluation). More information about the intercomparison exercises our lab participated in, as well as about GamW built-in utilities for routinely testing evaluations of complex spectra, can be found in HELCOM (2012).

2.3 Modelling Cs in fish

For validation of the activity concentrations for $^{134}$Cs and $^{137}$Cs measured in BS fish and giving an estimation of the future behaviour of the Cs isotopes in fish, the HELCOM box model for the BS case has been used (HELCOM, 1995; Kanisch et al., 2000). In this compartment model, the BS and the North Sea are separated into several water boxes in which the transport of radionuclides between water boxes, sorption and remobilization of radionuclides to/from the sediment, the physical decay of radionuclides and the input from anthropogenic sources is described by first-order differential equations. Modelling results have recently been presented in a HELCOM Thematic Assessment Report (HELCOM, 2009); see also Nielsen et al. (2010). The model takes time-dependent uptake by biota into account. For the special case of the fallout deposited to the BS, it has been extended to a two-compartment model by considering piscivorous fish (or predatory fish in the following) feeding on smaller fish (food fish). We refer to Rowan and Rasmussen (1996, 1997), as well as Smith (2006), Fievet and Plet (2003) and Wang et al., (2015), for the underlying bioenergetics-based assumptions. A direct uptake of Cs from seawater has been included according to Brown et al. (2006). The link between the model’s seawater activity concentrations, $\alpha$ $c_w$ $(t)$ in Bq L$^{-1}$, and the activity concentrations in food fish and the predating fish, $\alpha$ $c_f(t)$ and $\alpha$ $c_p(t)$ in Bq kg$^{-1}$ w.w., respectively, is given by a system of two coupled first-order differential equations solved by a Runge-Kutta method (Press et al., 1992). Including specific growth of the predating fish, the equations are

$$\frac{d\alpha c_f(t)}{dt} = K_{fd}(\lambda_{bio, fd} + \lambda_p)c_w(t) + k_wc_w(t) - (\lambda_{bio, fd} + \lambda_p)c_f(t), \quad (1)$$

$$\frac{d\alpha c_p(t)}{dt} = k_F(\alpha c_f(t) + k_wc_w(t)) - (\lambda_{bio, F} + \lambda_p + \lambda_G)c_p(t). \quad (2)$$

with rate constants $\lambda_p$, $\lambda_{bio, fd}$ and $\lambda_{bio, F}$ (d$^{-1}$) linked to the physical half-life of the radionuclide and the biological half-lives of food and fish, respectively; $\lambda_G$ is the specific growth rate (g g$^{-1}$ d$^{-1}$ or % d$^{-1}$) of the predator fish; $K_{fd}$ and $k_F$ are the concentration factors of food and predator fish, respectively, in L kg$^{-1}$; $\alpha$ is the food assimilation efficiency of the predator; for $k_w$ denoting the uptake rate of Cs from water a value of 0.04 d$^{-1}$ was used; $k_F$ denotes the predating fish’s rate of ingesting food (g g$^{-1}$ d$^{-1}$). The general relation between $T_{bio}$ and $\lambda_{bio}$ is $\lambda_{bio} = \ln2/T_{bio}$. An average value of 0.0015 g g$^{-1}$ d$^{-1}$ was estimated as $\lambda_G$ for BS cod, with a typical (total) length of about 50 cm, from the weight-based von Bertalanffy growth parameters, which are available in the FishBase database (Froese and Pauly, 2011). The quantities $k_p$ and $\alpha$ can be eliminated by expressing by analogy the first term of Eq. (2) like that of Eq. (1) and taking the relation $\alpha F = K_{fd}c_w(t)$ into account:

$$k_F\alpha F = K_F(\lambda_{bio, F} + \lambda_p + \lambda_G). \quad (3)$$

By implementing this in the model, assuming a single-time input to the Baltic seawater box surfaces, activity concentrations of fish were modelled by using a caesium concentration factor $K_F$ of 164 L kg$^{-1}$ (HELCOM, 2009; due to lower salinity in the BS it is larger than the value of 100 L kg$^{-1}$ used otherwise for marine fish; IAEA, 2004a) and an assumed value of 150 days for the biological half-life $T_{bio, F}$ (corresponding to the rate constant of the Cs elimination from fish). For food fish (smaller fish), assumed values of 50 L kg$^{-1}$ and 80 days were used for $K_{fd}$ and $T_{bio, fd}$, respectively.

Brown et al. (2006) used in their simulation study a range of $T_{bio, F}$ from 69 to 385 days for marine fish and a “representative” value of 116 d. We used a value of 100 days for modelling the behaviour of Sellafield-derived $^{134}$Cs in GC fish with a single-fish-compartment version; this case can be considered as very close to a steady-state situation; thus a value more appropriate than 100 days is not considered necessary. We did not try to model the Fukushima-derived $^{134}$Cs in Greenland fish, because we lack an appropriate sub-model describing how the $^{134}$Cs, deposited on the surface, would be mixed within the vertical water column. However, in the BS, with its significantly lower salinity compared to the Atlantic Ocean, the biological half-lives may be significantly larger for fresh-water fish than for pure marine fish (Rowan and Rasmussen, 1996, 1997). Therefore, a value of 150 days used for $T_{bio, F}$ in the BS is only a rough estimate.

The fish samples originating from the BS were collected from the water boxes 75, 81 and 83, which represent the Belt Sea and the Baltic Proper (Fig. 1). The water depths and the salinities are quite variable, with gradients from west to east: between about 30 and 160 m (depth), between 15 and 7.5 m (surface salinity) and between 20 and 8 m (bottom salinity) (HELCOM, 2010). Therefore, each of the three water boxes included a surface and a bottom layer; within the model, the surface layer thickness is about one-third of the total depth. For calculating the activities in fish, the modelled activity concentrations in seawater were averaged over the selected six box layers by weighting with their associated box-layer volumes. A value of 2 Bq m$^{-2}$ (see introduction) for each of the caesium isotopes was used for the Fukushima fallout deposited to the BS surface boxes.
Table 2. Decay-corrected activity concentrations $A$ of $^{134}$Cs, $^{137}$Cs and $^{40}$K and associated relative uncertainties ($u_{\text{rel}}$) in fish fillet samples from the Greenland and BS sites; the decision threshold DT (not shown) is well approximated by the value DL/1.548; SF is a significance factor, i.e. the ratio $A/DT$; “non-detects” are documented as $<DL$.

<table>
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<td>$\gamma$</td>
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<td>17679</td>
<td>cod</td>
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<td>0.0034</td>
<td>1</td>
</tr>
<tr>
<td>17720</td>
<td>cod</td>
<td>0.0045</td>
<td>0.0035</td>
<td>2</td>
</tr>
</tbody>
</table>

For the Greenland case, another box model was used which extends the HELCOM box model with respect to the seas in and around the Arctic (Nielsen et al., 1995); with regards to the BS, the results from this extended model agreed very well with results of the other model. The extended box model was used for clarifying possible $^{134}$Cs background sources in the Greenland case. However, it was not used to model the short-term behaviour of deposited Fukushima $^{134}$Cs in seawater and in fish around Greenland. If one were to do this anyway, it would lead to unrealistically small estimated concentrations in seawater and fish. This is caused by the compartment model’s specific property of instantaneously perfectly mixing the analyte within the overall box volume. This means in the case of the Greenland boxes that complete mixing is assumed over a large depth of 330 m in the surface layer, while the surface boxes are shallower in the BS. In reality one would expect a decreasing vertical activity concentration profile after fresh deposition on the seawater surface.

3 Results and discussion

3.1 Greenland seas – Cs-activity concentrations and source identification for fish

The results of the Cs measurements of Greenland fish are given in Table 2. The $^{137}$Cs activities showed a quite similar distribution pattern of four samples each between 2010 and 2011. It is found that $^{134}$Cs was clearly detected in the two 2011 fish samples (17704 and 17705) collected at the end of October 2011 in the West GC with ISO 11929-related significance factors $SF > 2.3 \gg 1$; SF is the ratio ($^{134}$Cs activity) / ($^{134}$Cs decision threshold), the latter being calculated according to ISO 11929. The activity concentrations found, around 0.005 Bq kg$^{-1}$, are extremely low. In contrast, in the two fish samples from the East GC (17702 and 17703), $^{134}$Cs was about two times smaller and thereby hardly detectable, SF $< 1$. The activity ratio $^{134}$Cs/$^{137}$Cs was about 0.013 for sample 17703, while it showed the identical value of 0.022 for the two West GC samples. As the $^{134}$Cs/$^{137}$Cs ratio of FD-NPP depositions was described to be close to 1, the FD-NPP-derived $^{137}$Cs must also be expected in fish of...
the same size for $^{134}\text{Cs}$. Therefore, for the two West GC samples of 2011, the ratio $^{137}\text{Cs}_{\text{Fuku}} / ^{137}\text{Cs}_{\text{pre-Fuku}} = ^{134}\text{Cs}_{\text{Fuku}} / ^{134}\text{Cs}_{\text{measured}} = 0.023$, which means that the Fukushima deposition added at most about 2% to the existing pre-Fukushima $^{137}\text{Cs}$ in the West GC, while it was about or less than 1% in the East GC samples. Whether the surface seawater activity concentrations in the two regions were significantly different cannot be deduced, because due to other sampling strategy reasons, the fish were sampled in stratified layers; average depths were about around 200 m and about 400 m in the West Greenland and East Greenland site, respectively (see Table 1). For the rather fresh fallout situation the Cs isotopes with FD-NPP origin are not fully mixed over depths down to nearly 400 m; therefore, the difference of $^{134}\text{Cs}$ values in fish between the two sites may simply be a consequence of different sampling/water depths when we assume that the fish stay for a longer time in these layers. The $^{134}\text{Cs}$ and $^{137}\text{Cs}$ values of fish were plotted versus sampling depth (Fig. 4). This reveals that the depth dependence of $^{137}\text{Cs}$ is rather weak, which means that slight $^{137}\text{Cs}$ differences found between the two sampling sites probably do not depend on sampling depth. However, the sampling depth dependency is quite strong for $^{134}\text{Cs}$. Thus, this observed sampling depth dependency, showing an incomplete vertical mixing of $^{134}\text{Cs}$ in the water column, supports our assumption that $^{134}\text{Cs}$ originates from the deposition of FD-NPP fallout.

For the coastal East GC, Dahlgaard et al. (2004) found surface $^{137}\text{Cs}$ concentrations of about 3–4 Bq m$^{-3}$ in 2001; they measured concentrations decreasing from 5 to 2.4 Bq m$^{-3}$ in this Current, from north to south in the eastern branch and in the coastal West GC up to the north again. They also determined $^{137}\text{Cs}$ in two fish samples collected in 1999 in the East GC with values of 0.35 and 0.47 Bq kg$^{-1}$ w.w. After decay correction with an effective half-life of 18.6 yr, by which the $^{137}\text{Cs}$ concentration in the North Atlantic seawater and in fish decreases (IAEA, 2005; according to data for the box 28 given therein), their average then amounts to 0.27 Bq kg$^{-1}$ in 2010. This is rather close to our 2010 and 2011 averages of about 0.20 Bq kg$^{-1}$ (from Table 2) for the two sites in the East GC. Extrapolating a $^{137}\text{Cs}$ seawater time trend for the corresponding water box number 28 taken from IAEA (2005) and using a concentration factor of 100 L kg$^{-1}$ leads to a value of around 0.2 Bq kg$^{-1}$ in fish. Our samples from the West GC show only slightly larger values of about 0.25 Bq kg$^{-1}$.

Dahlgaard (1995) estimated a transport time of 7–10 yr of such discharges to the East GC. For testing possible contributions of the Sellafield-derived $^{134}\text{Cs}$ background to our measurements in fish, the box model extended with respect to the Arctic Sea (see Sect. 2.3) was used. $^{137}\text{Cs}$ values around 0.1 Bq kg$^{-1}$ were obtained from this model using the surface layer (thickness 330 m) of the relatively large box “Irminger Sea” and a fish concentration factor of 100 L kg$^{-1}$. This is only 2.5 times lower than our measured values, i.e. not unrealistic. However, the corresponding values modelled for $^{134}\text{Cs}$ in fish, around $4 \times 10^{-6}$ Bq kg$^{-1}$, were 1000 times below our measured $^{134}\text{Cs}$ values. This finding assures us that the $^{134}\text{Cs}$ measured in fish originates from the FD-NPP fallout deposition. Due to reasons similar to those given at the end of Sect. 2.3, but also because the Chernobyl Cs deposition (Bq m$^{-2}$) to the GC is assumed to be much smaller than in the BS, it can be assumed that Chernobyl-derived $^{134}\text{Cs}$ in fish around Greenland would be much smaller than that estimated by modelling in the BS, and therefore would not contribute to the measured $^{134}\text{Cs}$ values.

### 3.2 Baltic Sea – Cs-activity concentrations and source identification for fish

Compared to the seas around Greenland with larger water flows, the rate of BS water exchange with the adjacent North Sea is rather small. In the Baltic the salinities therefore are lower due to freshwater inflows. The Chernobyl $^{137}\text{Cs}$ seawater inventory in the southern part of the BS (Baltic Proper) is decreasing, with an effective half-life of around 15 yr (HELCOM, 2009).

Results of measurements are included in Table 2; the samples from the additional retrospective screening of many other spectra for the existence of $^{134}\text{Cs}$ are included. In the BS, due to the impact of the Chernobyl fallout, the $^{137}\text{Cs}$ activity concentrations in fish are around 7 Bq kg$^{-1}$, which is about 35 times larger than in the GCs. $^{134}\text{Cs}$ was found in only three of the six samples with larger counting durations. The ISO 11929-related significance factors SF of the $^{134}\text{Cs}$ values where closer to 1, with about 1.36 at maximum, but for the other samples at around or even below 1. Thus, when including the four values from screening values, $^{134}\text{Cs}$ was definitely detected in only 50% of the samples. Therefore, methods for left-censored data (Helsel and Cohn, 1988; Helsel, 2005) were applied for obtaining descriptive statistics; mean and median values were about 0.0036 Bq kg$^{-1}$. The BS model was used for clarifying the origin of $^{134}\text{Cs}$. Figures 2 and 3 show the curves modelled in fish and measured values of $^{137}\text{Cs}$ and $^{134}\text{Cs}$, respectively. The lower and upper boundary (dashed) curves ($\pm 2\sigma$) were calculated from the standard deviation of the six individual box activity concentrations.

It had been tested how setting up the fish kinetic sub-model affects the shape of the modelled curve of $^{134}\text{Cs}$. Within the curves (Fig. 3), mainly their increasing part and the height of the maximum were found to be more variable, while the decrease following the maximum is determined by the decrease of modelled $^{134}\text{Cs}$ in seawater (near equilibrium between water and fish). Lowering the biological half-life as well as adding specific fish growth to the single-compartment sub-model resulted in a steeper increase of the $^{134}\text{Cs}$ activity concentration before reaching the maximum. Introducing the food fish compartment has the effect of some delay in this increase. Figure 3 shows the results of two model...
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Fig. 2. Curves of $^{137}$Cs activity per unit mass in fish using the two-compartment fish model, volume-averaged over the surface boxes 75, 81 and 83 and their associated bottom-layer boxes, and measured values (circles) of $^{137}$Cs in BS fish. Individual measured values are given with ±$2\sigma$ bars; the dashed lines represent the 95% confidence curves of the model (see text).

variants: (i) a single-compartment model neglecting specific growth and Cs intake from water (upper graph), and a (ii) two-compartment model taking both of these effects into account (lower graph). Therefore, the result obtained with a simple single-compartment sub-model is considered reasonable taken as a first guess.

The model curve for $^{137}$Cs in fish, which is mainly caused by the still large contribution of $^{137}$Cs deposited during the Chernobyl accident, is unaffected by the FD-NPP deposition; the increase is as small as the line thickness in Fig. 2a. For $^{134}$Cs, however, the effect of FD-NPP fallout is clearly visible in the model, but with very low values of the order of 0.004 Bq kg$^{-1}$ w.w., which is about 2000 times smaller than those of $^{137}$Cs. Note that the $^{134}$Cs modelled for the pre-FD-NPP time is the one still originating from the Chernobyl accident; its activity continues to decrease with an effective half-live $t_{eff}$ of 1.9 yr (calculated from the $t_{eff}$ value of 15 yr for $^{137}$Cs; HELCOM, 2009). Additionally, it has been tested that possible $^{134}$Cs contributions originating from small annual liquid discharges of the BS area nuclear power plants (see HELCOM, 2009, chapter (2)) would hardly be identifiable in the graph for $^{134}$Cs.

The good agreement between modelled and measured $^{137}$Cs values suggests that the modelled $^{134}$Cs values are as reliable as for $^{137}$Cs. As the two Cs isotopes have a different origin, one might argue that the very recently introduced $^{134}$Cs behaves differently from $^{137}$Cs with respect to mixing in the seawater. However, apart from one measured value from 19 April 2011, the measured $^{134}$Cs values from the post-Fukushima time are well within the lower and upper model boundaries (Fig. 2); this also holds for $^{137}$Cs. When continuing the upper $^{134}$Cs boundary curve from the beginning of 2011, i.e. when assuming no Fukushima input, this curve (pure Chernobyl $^{134}$Cs) during 2011 would also safely lie beneath the measured values. Therefore, we conclude that the major contribution to $^{134}$Cs values measured in fish, on average about 0.0026 Bq kg$^{-1}$, is related to the FD-NPP fallout deposition. The addition of $^{137}$Cs from FD-NPP to existing $^{137}$Cs in BS fish corresponds to an increase of only 0.04%.

3.3 Dose considerations about the Fukushima contribution

The contribution of the FD-NPP fallout deposition to effective dose rates received by human individuals through consumption of fish is estimated in relation to the corresponding pre-FD-NPP dose rates. Assuming an annual human consumption of 10 kg of fish fillets and estimating the FD-NPP contribution from averages of measured $^{134}$Cs values in fish using the activity ratio $^{134}$Cs/$^{137}$Cs of about 1 in March 2011 (neglecting the $^{134}$Cs decay), dose rate contributions
Table 3. Average activity concentrations (Bq kg\(^{-1}\)) and effective dose rate (µSv yr\(^{-1}\)) contributions to human individuals from consumption (10 kg yr\(^{-1}\)) of fish from 2011; dose conversion factors (Sv Bq\(^{-1}\)) used were 1.9 \times 10^{-8} and 1.3 \times 10^{-8} for \(^{134}\)Cs and \(^{137}\)Cs, respectively; note that \(^{134}\)Cs pre-Fukush, not occurring in the table, can be neglected.

<table>
<thead>
<tr>
<th>Radionuclide/Source</th>
<th>Greenland Bq kg(^{-1})</th>
<th>Greenland µSv yr(^{-1})</th>
<th>Baltic Sea Bq kg(^{-1})</th>
<th>Baltic Sea µSv yr(^{-1})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fukushima</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(^{134})CsFukush</td>
<td>0.0035</td>
<td>6.7E-04</td>
<td>0.0026</td>
<td>4.9E-04</td>
</tr>
<tr>
<td>(^{137})CsFukush \approx (^{134})CsFukush</td>
<td>0.0035</td>
<td>4.6E-04</td>
<td>0.0026</td>
<td>3.4E-04</td>
</tr>
<tr>
<td>Total F</td>
<td></td>
<td>1.1E-03</td>
<td>8.3E-04</td>
<td></td>
</tr>
<tr>
<td>(^{137})Cs measured, as measured</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(^{137})Cs pre-Fukush \approx (^{134})CsFukush</td>
<td>0.21</td>
<td>0.027</td>
<td>6.8</td>
<td>0.88</td>
</tr>
<tr>
<td>dose rate ratio: total F/(^{137})Cs pre-Fukush \approx relative increase of the dose rate by Fukush.</td>
<td>0.042</td>
<td>0.00094</td>
<td></td>
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</tbody>
</table>

Fig. 4. Activity concentrations of \(^{134}\)Cs (open circles, left scale, 2011) and of \(^{137}\)Cs (closed circles, right scale, 2010 + 2011) in fish fillets plotted versus sampling depth; measurement uncertainties given are ±\(\sigma\); Note: the smallest of the \(^{134}\)Cs values represents a “non-detect” with DL of 0.0032, for which a replacement value was plotted.

4 Conclusions

During our study, we determined caesium isotopes in fish samples from Greenland and the BS originating from FD-NPP fallout deposited over the North Atlantic as suggested by other publications on modelling its airborne deposition. \(^{134}\)Cs became detectable by using large amounts of fish fillets and an extended counting duration as compared to our standard method. In the Greenland Sea, which is much deeper compared to the BS, we observed a significant negative correlation (\(p = 0.0018\)) of \(^{134}\)Cs activities in fish with sampling depths (150–400 m), while this was not true for \(^{137}\)Cs. This indicates that \(^{134}\)Cs was not yet mixed in the water column which substantiates the Fukushima fallout as its origin, while a major part of \(^{137}\)Cs originates from global fallout and is well mixed over this depth. It also indicates the general feature of \(^{134}\)Cs as an ideal tracer that it is well suited for studying processes of vertical mixing of fallout in an ocean when the \(^{134}\)Cs levels prior to the fallout are extremely low.

The additional contribution through FD-NPP fallout to the effective dose received by humans through consumption of fish was calculated to be extremely low, i.e. 4 orders of

0.94 Bq kg\(^{-1}\) w.w. in BS fish (HELCOM, 2009), a human fish consumer receives a dose of 11 µSv per year after consumption of 10 kg fish per year; a \(^{210}\)Po dose conversion coefficient of 1.2 \(10^{-8}\) Sv Bq\(^{-1}\) has been considered (ICRP, 1996). In comparison, this dose rate is larger than that from the Fukushima contribution by more than a factor of 10\(^4\). The data in Table 3 also imply that even the dose rate received from Chernobyl-derived \(^{137}\)Cs in BS cod (0.88 µSv yr\(^{-1}\)) is nearly 100-fold larger than that of the Fukushima contribution. Within a dose assessment for consumption of sea fish (Aarkrog et al., 1997), a \(^{210}\)Po value in fish of 2.4 Bq kg\(^{-1}\) was used which, taking the meanwhile larger ICRP dose conversion coefficient into account, would result in a dose rate of 29 µSv yr\(^{-1}\) for consuming 10 kg per year.
magnitude lower than the effective dose rate received at the same time by the naturally occurring $^{210}$Po, which is persistent in fish. Due to the short duration of deposition at sea, it is expected that levels of caesium isotopes from FD-NPP will continue to decrease in seawater by vertical and horizontal mixing and by physical decay. Therefore, our results indicate that, although based on a limited set of data, the improved monitoring of $^{134}$Cs in fish, also in other northern areas such as the Barents Sea, in combination with the northeast Atlantic waters box model, are essential tools to maintain consumer safety.

Appendix A

Details of gamma-ray spectrometric analysis with GamW

A numerical procedure for geometry-dependent self-attenuation corrections, accounting also for different elemental compositions of fish ash and the calibration solution, is included in GamW. Calculated self-attenuation corrections at 605 keV were between 0.91 and 0.94.

The method used for the TCS calculations including X-ray contributions was that of Novčović et al. (2007); see also Kanisch et al. (2009). The extension from point to volume sources was achieved by the "LS-curve" method (Vidmar and Korun, 2006; Vidmar and Kanisch, 2010; Lépy et al., 2012). Approximate total efficiency curves required by such calculations were obtained with the gamma-ray spectrometry simulation tool GESPECOR (Sima et al., 2001). The linear energy calibration is re-calculated for each sample spectrum according to well-known radionuclide energies in the spectrum.

The spectrum evaluation then starts with a peak search followed by nuclide-library-based complementation of the list of peaks; peak multiplies are fitted by non-linear weighted least squares (Press et al., 1992; Levenberg–Marquardt subroutines MRQMIN). The peak shape function used (Koskelo et al., 1981) consists of a Gaussian function with low-energy tailing, extended by adding a step function to the 1–3 parameter background-continuum polynomial. The method described by Uher et al. (2010), which is in fact a penalized weighted least-squares method, was implemented into the MRQMIN package for stabilizing the fits, especially in the case of small peaks, which is achieved by adding a penalty term to the chi square to be minimized for the non-linear parameters (width, location, relative step height and the tailing parameter of a peak). Thus, such parameters are allowed to vary, thereby accounting for slight deviations from calibration values as they may occur during long-term measurements, and, on the other hand, they are prevented from leaving safe value domains, which guarantees convergence of the fit.

Net background peak counting rates are subtracted from the corresponding gross peak counting rates. Then, decay-corrected massic radionuclide activities are calculated by solving a linear matrix equation system by the linear least-squares method according to the standard ISO 11929 (2010; Annex C.5 therein) taking peak efficiency correlations into account (Glavic-Cindro et al., 2004). Major relative standard uncertainties were about 1.5% (peak efficiency around 662 keV), 2.9% (preparing the fish ash as counting source including the sample wet/dry weight ratio uncertainty), about 0.5% (gamma emission probabilities), 1% (self-attenuation corrections) and around 2% (TCS).

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Decision thresholds and detection limits were calculated for each sample measurement by applying the ISO 11929 concept to the matrix-based least-squares procedure; that is, not only for the easier-to-solve single-peak case of $^{137}$Cs but also for $^{134}$Cs for which one or two lines (at 604.8 and 796.0 keV) have been used; in the latter case the detection limit value is lower than for a single peak. Treating this by using a weighted mean rather than by least squares has been considered by Vivier et al. (2012). Values of $k_{1-\alpha} = 3$ and $k_{1-\beta} = 1.645$ were used for the quantiles of the normal distribution; the larger $k_{1-\alpha}$ value was used in order to prevent finding too many non-significant peaks (De Geer, 2004).

A1 Special measures for improving the detection of $^{134}$Cs

For reducing the uncertainty of the fitted areas of the background continuum polynomial, the multiplet regions of interest (ROI) around 605 and 796 keV were extended to the ranges of about 560 keV to 630 keV and from 782 keV to 825 keV, respectively; the first ROI often was enlarged to include also the $^{137}$Cs peak at 661.7 keV. Peaks from naturally occurring radionuclides, also found in the separately measured background spectra, appeared at 609.3 keV ($^{214}$Bi) and at 794.9 keV ($^{228}$Ac), the latter lying quite close to the $^{134}$Cs peak at 795.9 keV. Especially the spectra measured with the larger detector 4 showed more clearly an extra gamma-ray peak at around 596 keV as well as an additional broad background continuum contribution with a slow exponential decay towards energies above about 620 keV. This is due to neutrons caused by cosmic ray interactions in the shielding materials, which, then, by interactions within the detector, produce special gamma-ray peaks (e.g. at 596 keV), and, amongst others, this broad “neutron peak” above 596 keV due to inelastic scattering with Ge (Heusser, 1995; Jovančević et al., 2010); it is much more easily recognized in background spectra in which the $^{40}$K related Compton background is much smaller than in fish ash samples. The functional representation of this “neutron-induced peak” given by Siiskonen and Toivonen (2005) has been implemented here in a simplified way as a further background component in the total fitting function. Figure A1 shows an example for the fitting region containing the 605 and 662 keV peaks in the spectrum of sample 17705 (Greenland site).

The upper limit of relative standard uncertainties of single peaks entering the matrix equation mentioned above was raised to 50%; however, according to ISO 11929, final activity values were considered as being detected only if they were above the decision threshold. TCS corrections for $^{134}$Cs being comparable between its two major lines were estimated at between about 14 and 24 % for the three detectors. For the fish measurements, uncertainty budgets of activity concentrations (IAEA, 2004b) were dominated by contributions of sample preparation, peak efficiency and self-attenuation correction for $^{40}$K and $^{137}$Cs, while the counting rates contributed most dominantly in the case of $^{134}$Cs.

Acknowledgements. We are grateful for the valuable comments given by the reviewers H. Nies and G. Pröhle and an anonymous reviewer, which significantly improved the quality of the paper. For the opportunities of receiving samples from RV Walther-Herwig III (Baltic Sea), from commercial Baltic Sea catches and for samples from around Greenland we thank our colleagues Thomas Lang, Joachim Dröse and Heino Fock from the Thünen Institute.

Edited by: H. Nies

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