# Intermediate water from the Greenland Sea in the Faroe Bank Channel:

spreading of released sulphur hexafluoride

K. Anders Olsson<sup>a</sup>,\*, Emil Jeansson<sup>a</sup>, Leif G. Anderson<sup>a</sup>, Bogi Hansen<sup>b</sup>, Tor Eldevik<sup>c,d</sup>, Regin Kristiansen<sup>b</sup>, Marie-José Messias<sup>e</sup>, Truls Johannessen<sup>f,d</sup>, Andrew J. Watson<sup>e</sup>

<sup>a</sup> Department of Chemistry, Göteborg University, SE-412 96 Göteborg, Sweden
<sup>b</sup> Faroese Fisheries Laboratory, Box 3051, FO-110 Tórshavn, Faroe Islands
<sup>c</sup> Nansen Environmental and Remote Sensing Center, Thormøhlensgate 47, NO-5006 Bergen, Norway
<sup>d</sup> Bjerknes Centre for Climate Research, Allégaten 55, NO-5007 Bergen, Norway
<sup>e</sup> School of Environmental Sciences, University of East Anglia, Norwich NR4 7TJ, United Kingdom
<sup>f</sup> Geophysical Institute, University of Bergen, Allégaten 70, NO-5007 Bergen, Norway

\* Corresponding author, now at: Bjerknes Centre for Climate Research and Geophysical Institute,
University of Bergen, Allégaten 55, NO-5007 Bergen, Norway. Phone: +47-555-89817, fax: +47-555-84330
*E-mail address:* anders.olsson@bjerknes.uib.no (K.A. Olsson).

#### Abstract

The Faroe Bank Channel is the deepest passage for dense water leaving the Nordic Seas into the North Atlantic. The contribution to this part of the Greenland-Scotland Overflow by intermediate water from the Greenland Sea is investigated by the tracer sulphur hexafluoride  $(SF_6)$  that was released into the central Greenland Sea in summer 1996. Continuous monitoring has since traced it around the Nordic Seas and into the connecting areas. It was for the first time observed close to the Faroe Islands in early 1999, indicating a transport time from the Greenland Sea of around 2.5 years. This study estimates that approximately 16 kg of  $SF_6$  had passed the Faroe Bank Channel by the end of 2002; that is 5 % of the total amount released. Both the arrival time and the amount of exported  $SF_6$  deduced from the observations are consistent with the results from a numerical ocean model simulating the tracer release and spreading.

*Keywords:* Tracers, Overflow, Intermediate water masses, Thermohaline circulation, Sulphur hexafluoride, Nordic Seas, Greenland Sea, Faroe Bank Channel

## 1. Introduction

The Nordic Seas are of great climatic importance as they are the northern terminus of the oceanic conveyor belt (e.g. Broecker, 1991). The relatively high salinity of the warm Atlantic water that enters the Nordic Seas is a crucial ingredient in the formation of the deep-water masses which return as dense water from the Nordic Seas into the North Atlantic where it again mixes and transforms into North Atlantic Deep Water.

The circulation pattern in the Nordic Seas (Fig. 1a) involves the north-flowing Norwegian Atlantic Current (NwAC) and Faroe Current (FC), which together carry ~7 Sv (1 Sverdrup =  $10^6 \text{ m}^3 \text{ s}^{-1}$ ) of Atlantic water into the Nordic Seas. A much smaller inflow of Atlantic water, the North Icelandic Irminger Current (NIIC) enters north-west of Iceland (Fig. 1a). The main outflow is the East Greenland Current (EGC) that flows to the south along the Greenland continental margin and can be divided up in a shallow and a deep fraction of about the same magnitude (~3 Sv) (Hansen and Østerhus, 2000). There is also energetic circulation within the Nordic Seas. Cold surface water is deflected from the EGC to the east both at about 72°N by the Jan Mayen Current (JMC) as well as into the Iceland Sea. Intermediate and deep water of the Greenland Sea pass over the Mohns Ridge and through the Jan Mayen Channel into the Norwegian Sea (e.g. Østerhus and Gammelsrød, 1999) and intermediate water enters the southern Norwegian Sea from the Iceland Sea through the East Iceland Current (EIC). A schematic illustration of the current system is shown in Fig. 1a.

Totally there are four paths for cold overflow from the Nordic Seas to the deep North Atlantic; in addition to the one between Greenland and Iceland (the Denmark Strait), these exit between Iceland and the Faroe Islands, across the Wyville-Thomson Ridge, and through the Faroe Bank Channel (FBC). The total overflow between Iceland and Scotland is of similar magnitude to that through Denmark Strait and the deepest exit is through the Faroe Bank Channel after passing the Faroe-Shetland Channel between the Faroe and Shetland Islands (Fig. 1b). The present volume flux through the FBC, of water colder than 3°C, has been estimated to 1.9 Sv, but it is suggested that this flux has decreased by about 0.5 Sv during the last  $\sim$ 50 years (Hansen *et al.*, 2001). The pure overflow, defined as water below 0.3°C, is estimated to be 1.2 Sv (Hansen and Østerhus, 2000) and is estimated to have decreased as much as 2-4 % annually 1995-2000 and in total at least 20 % during the last 50 years (Hansen et al., 2001). This latter definition of the overflow is used throughout this work. Results using a synoptic-forced numeric model (Nilsen et al., 2003) indicate substantial decadal variability both in the north-flowing and south-flowing volume flux over the Greenland – Scotland Ridge. The overflow has most likely also decreased across the Iceland-Faroe Ridge and the Wyville-Thomson Ridge (Hansen et al., 2003). Time series also reveal a change in water properties of the FBC overflow which has became fresher (Turrell et al., 1999) mainly by an elevated incorporation of intermediate water due to a decrease in deepwater formation. Both the inflow and the overflow are thoroughly discussed by Hansen and Østerhus (2000).

To investigate the formation of water contributing to the overflows into the North Atlantic, a tracer release experiment was initiated in summer 1996 (see Watson *et al.*, 1999). In total 320 kg of sulphur hexafluoride (SF<sub>6</sub>) was injected to the central Greenland Sea Gyre (see Fig. 1b) at the density surface,  $\gamma_{\theta} = 28.049$  kg m<sup>-3</sup> at an average depth of about 300 m<sup>-1</sup>. Sulphur hexafluoride is a compound that is almost entirely man-made (Harnisch and Eisenhauer, 1998) and it has been utilised as a deliberately released tracer in a range of oceanographic studies (Watson and Ledwell, 2000). In addition to this, SF<sub>6</sub> has been used as a transient tracer in recent investigations (Law and Watson, 2001; Tanhua *et al.*, 2004) in the same way as the chlorofluorocarbons (CFCs).

The Greenland Sea tracer release has made it possible to follow intermediate water from the Greenland Sea around the Nordic Seas and into connecting areas. The spreading of the tracer has been monitored ever since the release and the tracer is now found in most parts of the Nordic Seas and a has also overflown the Greenland-Scotland Ridge and entered the North Atlantic (Messias *et al.*, 2004; Olsson *et al.*, 2004).

# 2. Data and Methods

#### 2.1. Data collection

The data reported in this work have mainly been collected within the EU projects "European Subpolar Ocean Programme, phase 2" (ESOP-2) and "Tracer and Circulation in the Nordic Seas Region" (TRACTOR). The experiment with the release of  $SF_6$  was performed in ESOP-2 that was dedicated to investigate the thermohaline circulation mainly by studies in the Greenland Sea (Messias *et al.*, 1999). One of the goals of the TRACTOR project has been to follow the  $SF_6$  tracer as it spreads in time and space from the Greenland

<sup>&</sup>lt;sup>1</sup> Often reported as  $\sigma_{\theta} = 28.049$  with or without units, which is not consistent with the JPOTS standard (JPOTS editorial panel, 1991).

Sea to the surrounding basins. Further results from the experiment are presented elsewhere (Watson *et al.*, 1999; Gascard *et al.*, 2002; Olsson *et al.*, 2004).

Samples have been collected around the Faroe Islands regularly since 1998, mainly north of the islands and in the FBC (Fig. 1b). Information on the station activity in the FBC is given in Table 1. Samples for  $SF_6$  were collected with rosettes mounted on SeaBird CTDs and temperature and salinity measurements accompany the samples. Most of the sampling sites are located on standard sections, which are occupied by R/V Magnus Heinason at least four times a year. From these cruises, a large number of CTD profiles have been acquired from the areas around the sampling sites. On each CTD station, a double set of water samples is collected at one depth and analysed by salinometer for salinity calibration.

The determination of SF<sub>6</sub> was performed using a gas chromatograph with electron capture detection (ECD) coupled to a purge-and-trap pre-treatment system (Law *et al.*, 1994; Tanhua *et al.*, 2004). The standard deviation of a set of samples from the same depth is less than 2 %. The standardisation of sulphur hexafluoride was performed using gas calibrated against standards from Plymouth Marine Laboratory (UK) that in turn was calibrated against standards from University of Heidelberg (Germany). The concentrations of all samples in this study were well above the detection limit of about 0.1 fmol kg<sup>-1</sup>.

#### 2.2. Calculations

The transient atmospheric source of  $SF_6$  must be considered also in a deliberately-released tracer experiment. The atmospheric background signal of  $SF_6$  in a seawater sample can be estimated from the observed CFC concentrations. The atmospheric history is obtained from Walker *et al.* (2000) for the CFCs and from Maiss and Brenninkmeijer (1998) for  $SF_6$ . The solubility of the gases in oceanic surface water is computed by equations that are functions of

potential temperature ( $\theta$ ) and salinity (S) (Warner and Weiss, 1985; Bu and Warner, 1995; Bullister *et al.*, 2002). Time series for a typical upper water in the Greenland Sea are illustrated in Fig. 2a.

Mixing of water masses of different ages can result in apparent ages that are quite different for CFCs and SF<sub>6</sub> and hence make it problematic to calculate the atmospheric background. To minimise this uncertainty it is necessary to evaluate which source waters that are mixed. From the cruises where CFCs were available (see Table 1), the overflow water in the FBC was divided into three different types, upper, middle and lower, based on the properties of the observed samples, (average  $\theta$  and S are shown in Table 2). In the next step the mixing histories of these water types were trying to be solved. All three water types have evidently a portion of water from the Greenland Sea (Greenland Sea Arctic Intermediate Water, GSAIW), as shown by the elevated levels of SF<sub>6</sub>. The other contributing water masses are believed to be: Norwegian Sea Deep Water (NSDW), and Northeast Atlantic Water (NEAW) and Modified East Icelandic Water (MEIW) (Fogelqvist et al., 2003). The water mass commonly defined as Norwegian Sea Arctic Intermediate Water (NSAIW), believed to be a major contributor to the Iceland-Scotland Overflow Water (Hansen and Østerhus, 2000; Fogelqvist et al., 2003), is not included here since it is to a large extent made-up from what is called GSAIW (Blindheim, 1990) herein. This name is used to make it clear that this is the water mass containing the tracer from the Greenland Sea. The properties of GSAIW differ slightly from those of NSAIW, which is affected by NSDW situated underneath and the mixing between these two occurring in the boundary layer. This can be the reason why this study gives a larger portion of NSDW in the overflow compared to (Fogelqvist et al., 2003) who use NSAIW as source water instead of GSAIW. The properties of the intermediate water from the Greenland Sea containing the released tracer are taken from observations made

during the ESOP-2 project, 3-4 months after the release (Tanhua and Olsson, unpublished data) and the properties of the other involved water masses are obtained from (Fogelqvist *et al.*, 2003).

The temporal evolution of the concentrations of CFC-12 and SF<sub>6</sub> in the four source waters was computed using the solubility and  $\theta$  and S data and assuming equilibrium with the atmosphere. Furthermore, as the time from contact with the atmosphere to their appearance in the FBC varies between the source waters, this has to be considered when computing the background concentration. The observed CFC-12 concentration in GSAIW is comparable to a time delay of 15 years from ventilation to observation. This does not mean that the "age" of the water mass is 15 years since an eventual under-saturation is also included here. If a two-year transit time from the Greenland Sea to the FBC is used (which agrees well with the tracer observations) we get a total time delay of 17 years for GSAIW. The observed CFC-12 concentration in NSDW is comparable to a time delay of 31 years, which together with a transit time of 1 year gives a total delay of 32 years. MEIW and NEAW are saturated with respect to CFC-12 but a delay time of 1 year is used. Thus, the time evolution in the three water types of the FBC is computed by adding together the contributions by the different source waters and taking the time delay into consideration according to Equation 1.

$$X_{year=i}^{wt} = f^{GSAIW} \cdot X_{year=i-17}^{GSAIW} + f^{NSDW} \cdot X_{year=i-32}^{NSDW} + f^{NEAW} \cdot X_{year=i-1}^{NEAW} + f^{MEIW} \cdot X_{year=i-1}^{MEIW}$$
(1)

Here *X* represents the concentration of any tracer and *f* the fraction of source water contributing to the water type (wt). The fractions of the four source waters are computed to fit  $\theta$ , S and CFCs (Table 2) observed in the three water types. When computing the background for a given year *i*, the concentration of the tracer in the source water is taken for the year *i* 

minus the transit time as indicated by the subscript for any of the source waters given by the superscript. The results of the computations are shown in Fig. 2b. For comparison the computed relationships for 1997 and 2001 in the three water types are indicated. No direct comparison with observations can be made since there are no one made with only background  $SF_{6}$ .

For each observed sample, one of the three water types are chosen to be the most representative, decided by its  $\theta$  and S properties. The corresponding water mass fraction (see Table 2) and year are put into Equation 1 and the resulting background concentration is subtracted from the observed concentration of each sample.

#### 3. Results

The typical water mass distribution on a section crossing the FBC is illustrated by an example in Fig. 3. The deepest part of the channel is always occupied by dense overflow water on its way into the Atlantic Ocean, passing the section from the southeast to the northwest, and usually water with density exceeding  $\gamma_{\theta} = 28.049$  kg m<sup>-3</sup> (the release density) dominates the bottom layer (Fig. 4). Thus, all of the 23 CTD profiles acquired at the deepest standard station, V06, in the period 1998-2001, reached or exceeded this density at the deepest measurement. Unfortunately, the SF<sub>6</sub> samples have not always reached this layer, but on most cruises, this has been the case.

All water below about 600 m had  $\gamma_{\theta}$  >28 kg m<sup>-3</sup>, and showed increasing SF<sub>6</sub> concentrations from at least year 2000 (Fig. 5). However, the SF<sub>6</sub> concentration does not show a continuous increase with time, but the variability is to some extent reflected in the variability in water properties. The water shallower than about 500 m (not shown) has had less relative increase in SF<sub>6</sub> concentration (1.4 fmol kg<sup>-1</sup> at 300 m already in June 1999) and this signal is of atmospheric origin only.

When the measured SF<sub>6</sub> concentrations are corrected for the background, and all data below 600 m are plotted versus time a clearer pattern is seen (Fig. 6). In the autumn of 1998 the SF<sub>6</sub> concentration is close to the background values, after which it increases almost linearly until spring/summer 2000. After this the excess concentration varies around  $0.9\pm0.1$ fmol kg<sup>-1</sup>, except for the last observations in 2002 when it had once again increased. At some occasions the concentrations show larger variations, mainly a result of the presence of water that was not well represented by any of the three water types, e.g. more saline water and hence containing a larger portion of Northeast Atlantic Water. The reason for the increase in June 2002 (Fig. 6) is not known but might indicate that the overflow is supplied by more than one pathway, and that the transit time of the second caused it to appear 4 years later. Alternatively the overflow could have a quite different composition with a larger fraction of water originating in the Greenland Sea.

# 4. Discussions

#### 4.1. Time evolution

As illustrated in Fig. 6, the SF<sub>6</sub> signal from the tracer release did not appear in the FBC until winter 1998-99, which is slightly more than 2 years after the tracer was released in the Greenland Sea Gyre (summer of 1996). This should then reflect the transit time of a water parcel, at the density level of the release, from the central Greenland Sea to the FBC. The build-up time, from fall 1998 to spring/summer 2000, is ~1.5 years and should be the result of mixing both within the Greenland Sea and during the transit to the FBC. Observations

indicate that the SF<sub>6</sub> signal was homogeneously mixed ( $\pm 10$  %) in the Greenland Sea gyre after 1.5-2 years (Messias *et al.*, 1999). The level in the Greenland Sea has continued to decrease ever since although at a slower rate.

The escape from the central Greenland Sea occurs presumably both direct and indirect; to the west diffusive mixing probably dominates the spread into the East Greenland Current bordering the Greenland Sea, while advection likely plays a larger role when exiting to the east through the Jan Mayen Channel and into the Norwegian Sea. The close to stable excess tracer signal in the FBC could be a combination of mixing of water taking two routes from the Greenland Sea, one more directly and one e.g. passing further into the Norwegian Basin. It should be noted that the strength and relative importance of the different flow paths are not constant in time. The essential role of the Jan Mayen Current in the simulated tracer spreading presented in section 4.3 underlines this. Also, it is becoming more and more evident that the variability of the atmospheric forcing has an impact on the Nordic Sea current regimes, the FBC region included (e.g. Nilsen *et al.*, 2003).

#### 4.2. Transport estimates

The concentration evolution in the FBC combined with literature values of the volume fluxes in different depth layers (e.g. Hansen *et al.*, 2001) gives an estimate of the SF<sub>6</sub> transport. The volume flux below the 0.3°C isotherm shows a seasonal signal with highest values in fall and lowest in spring and a decreasing trend of 2-4 % per year from 1996 to 2000 (Hansen *et al.*, 2001). The mean flux in 2000 was ~1.2 Sv, which would give a mean transport of SF<sub>6</sub> during the phase of constant excess (~0.9 fmol kg<sup>-1</sup>) of ~5 kg SF<sub>6</sub> yr<sup>-1</sup>. Adding together the annual transports from the fall 1998 to the winter of 2002/2003, would give a total transport of 1.5 years x 5 (kg yr<sup>-1</sup>)/2 (during the build up phase) plus 2.5 years x 5 (kg yr<sup>-1</sup>) (during the constant phase), or 16 kg. This is ~5 % of the SF<sub>6</sub> released in the central Greenland Sea. This means that although the concentrations are low, significant amounts of the tracer are exported thanks to the large fluxes and this indicates that the importance of different regions can hence easily be underestimated. It is assumed that the amount of tracer exported above the  $0.3^{\circ}$ C isotherm is insignificant since the water properties indicate that very little water from the Greenland Sea is included here. The  $0.3^{\circ}$ C limit is defined to include NSDW and NSAIW and hence GSAIW (see Hansen *et al.*, 2003).

#### 4.3. Comparison with a numerical ocean model

Further understanding of the pathways of SF<sub>6</sub> within the Nordic Seas and subsequent export through the FBC may be gained from a numerical ocean model. For this purpose, we use an advective-diffusive model of tracer transport and mixing within the intermediate waters of the Nordic Seas (Eldevik *et al.*, 2004). The model was originally set up by Straneo *et al.* (2003) to describe the spreading of Labrador Sea Water. The stationary horizontal velocity field prescribed to it should ideally be based on *in situ* observations (which is the case for the Labrador Sea study), but adequate current data are not available for the Nordic Seas. Eldevik *et al.* (2004) deduce their flow field (on a 10 km x 10 km grid) from the output of the high resolution (20 km) general circulation model (GCM) of Hátún *et al.* (2004). The GCM is a version of the Miami Isopycnic Coordinate Ocean Model (MICOM, Bleck *et al.*, 1992) covering the North Atlantic and the Nordic Seas for the period 1951-2000. This regional setup, as well as its corresponding global (and coarser) versions focused on the Arctic Mediterranean, should be state of the art for GCM systems covering the Nordic Seas. The GCMs' spreading of active (Furevik *et al.*, 2002) and passive (Gao *et al.*, 2004) tracers, and Atlantic-Nordic Seas exchanges (Nilsen *et al.*, 2003; Hátún *et al.*, 2004) have all been evaluated favourably against observations.

The velocity field used by the advection-diffusion model is the average of the GCM horizontal velocity over the intermediate water (IW) column. The IW is here defined to be the water between 500 and 1500 m depth (where the ocean depth is less than 1500 m, the average velocities are weighted consistent with the reduced thickness). The SF<sub>6</sub> is generally observed in the IW part of the water column in the Nordic Seas (cf. Messias et al., 2004). For the case at hand, the mean 1997-2000 GCM flow field is used. The corresponding streamlines of the advective-diffusive solver are displayed in Fig. 7, where the black rectangle centred at 1.5°W, 75.25°N is the August 1996 release site. It does show the general IW patterns of Fig. 1a, but there are features more emphasized in the model current field. Particularly, the streamlines following the Jan Mayen Current, then diverting into the Norwegian Sea east of Jan Mayen, are more important contributors to the export of the simulated IW towards the FBC than those of the deep East Greenland Current. Note that the model domain does not really include the Faroe Bank Channel. The southern open boundary is the slightly upstream Faroe-Shetland Channel. Comparing the observed and modelled tracer concentration at the two different sites should nevertheless be consistent. What leaves the Nordic Seas through the Faroe-Shetland Channel is exported to the Atlantic predominantly through the FBC (about 90%), with the flow over the Wyville-Thomson Ridge closing the budget (cf. Hansen and Østerhus, 2000).

The modelled outflows through the Denmark Strait (0.9 Sv) and the Faroe-Shetland Channel (1.7 Sv) are realistic. Corresponding observationally based estimates are 0.6 Sv (Girton *et al.*, 2001) and 1.7 Sv (Hansen and Østerhus, 2000). There is no exchange between Iceland and the Faroes as the Iceland-Faroe Ridge is shallower than 500m. The net IW inflow through the Fram Strait, in the conservative model being the sum of the Denmark Strait and Faroe-Shetland Channel fluxes, is thus 2.6 Sv. The model's Laplacian eddy diffusivity is assumed constant, and set to the nominal value  $100 \text{ m}^2 \text{s}^{-1}$ . Eldevik *et al.* (2004) also do simulations where they vary the magnitude of the diffusion, as well as the degree of anisotropy and spatial inhomogeneity, and find that the results are rather insensitive to this particular choice. The reader is referred to Eldevik *et al.* (2004) for further details on the model setup and evaluation.

A snapshot from the numerical simulation of the SF<sub>6</sub> release and spreading is seen in Fig. 8a. From the release site, the tracer is advected along and mixed across the streamlines of Fig. 7 for six model years to produce the summer 2002 distribution in Fig. 8a. A synthesis of the corresponding *in situ* observations (column inventories), taken from (Messias *et al.*, 2004), is shown in Fig. 8b. The crosses indicate the hydrographic stations. The agreement between model and field data is fair with respect to patterns, and good on concentrations. The observations support the model prediction of the importance of the pathway directly from the Greenland Sea to the Norwegian Sea via the Jan Mayen Current. Consistent high concentration anomalies are found roughly following the 2000-m isobath in Fig. 8b: from the north to the south-east of Jan Mayen, then to the western Norwegian Basin, and then to the north of the Faroes. The model seems to be off in the central Norwegian and Lofoten basins, where it predicts two high concentration anomalies. This suggests that the model recirculation in the two basins (cf. Fig. 7) is somewhat too strong, although a similar pattern is present in the flow climatology of Nøst and Isachsen (2003).

The concentration of  $SF_6$  in the FBC predicted by the model is compared with the observations in Fig. 6. The model concentration, given in tracer mass per unit area, has been converted to mass per unit volume by dividing it by the thickness of the IW. There is a remarkable agreement in arrival time, build-up time, and 'end' concentration between the

two. The modelled total export through FBC by winter 2002/2003 is 15 kg, consistent with the 16 kg estimated from observations.

The comparison of the model results and the observations shows that the rather simple advective-diffusive model is quite skilful. The key component in reproducing the observed export of  $SF_6$  is the direct pathway east of Jan Mayen from the Greenland Sea to the Norwegian Sea. It suggests that this 'shortcut' is more important than previously assumed (cf. Fig. 1a), at least as the IW circulation of the late 1990s and early 2000s is concerned.

# 4.4. Uncertainties

The uncertainties in estimating the tracer transport are the results of at least four different components, the actual observations, the background subtraction, the time evolution and the volume fluxes. The errors originating in sampling, analysis and calibration of the tracer data are only a few percent and insignificant compared to the total.

The largest uncertainty is associated with the establishment of the background-corrected  $SF_6$  concentration. This is dependent on how well all the observed water samples fit into the defined water types and how representative and constant the source water compositions of these water types are. It is also dependent on the properties of the defined source waters, e.g. the assumed tracer saturation, and how variable they are.

The common variability in time and space affects the FBC water types and the source waters as well as the sampled locations in the FBC. The mean excess  $SF_6$  for all samples in the FBC between June 2000 and February 2002 is 0.9 fmol kg<sup>-1</sup> with a standard deviation of 0.1 fmol kg<sup>-1</sup>. The variations are relatively small and hence we assume that it can be relatively representative for the last 30 months and for the whole FBC.

How well the water types are able to represent the water samples is hard to tell.

However, some estimates can be made from the observed CFC concentrations based on the ranges within each water type and differences between the water types. From this approach a rough estimate 0.1 fmol kg<sup>-1</sup> is obtained. It might be higher for the more saline samples, e.g. above 34.91 although these are so few that they do not considerably affect the estimates.

The offset between the mixing proportions in Table 2 and the actual mixing history of each water type is dependent on a couple of things. The four selected source waters are those believed to contribute to the overflow (Hansen and Østerhus, 2000; Fogelqvist et al., 2003). The error from the mixing proportions is limited by the fact that no source water dominates the total SF<sub>6</sub> contribution, each one stands for a fraction between 9 and 43 %. This implies that a more than 20 % change in any of the water masses is needed to receive a markedly different background concentration. A direct comparison between the fraction of GSAIW and the SF<sub>6</sub> concentration cannot be made since the concentration in the Greenland Sea has been decreasing with time and was patchy to start with. Initially, part of the GSAIW passing the FBC might have left the Greenland Sea before the experiment and be free of released tracer which makes a straightforward approach difficult. Based on the mean water mass composition of all overflow samples in this study, as deduced from the water types, the fraction of GSAIW in the overflow would be 17 %. If this is true the 0.9 fmol  $kg^{-1}$  in the FBC would have been about 5.5 fmol kg<sup>-1</sup> in the Greenland Sea. As mentioned above this value has however been changing considerably with time although as an average it is reasonable compared to the observations (Messias et al., 2004).

An estimate of the extreme offset from the assumed  $SF_6$  levels in the source waters can be done. The uncertainty is mainly due to the degree of saturation of the tracer. Since the same processes determine the uptake of CFCs and  $SF_6$  from the atmosphere to the ocean it is

assumed that these gases are saturated to the same degree in surface water. The saturation of SF<sub>6</sub> in the two recently ventilated water masses, MEIW and NEAW, is assumed to be 100 % which is consistent to the saturation of CFCs observed by Fogelqvist et al.(2003). In the Greenland Sea however, the saturation for CFCs has been reported to be around 80 % (Bullister and Weiss, 1983; Rhein, 1991; Anderson et al., 2000). Such an undersaturation implies that the actual atmospheric concentration was higher than the value directly obtained from the observed tracer concentration in the water and compensation for an assumed degree of under-saturation gives hence a lower age estimate. Assuming a saturation of 80 % in GSAIW instead of 100 % results in a CFC age being five years lower and the corresponding SF<sub>6</sub> concentration, five year later but saturated to 80 %, would be higher, cf. 0.63 instead of 0.46 fmol kg<sup>-1</sup>, an effect of the different temporal evolution of the two compounds (Fig. 2a). The alternative that GSAIW in the Greenland Sea would be newventilated and the relatively low observed concentration of CFCs was caused only by low saturation (65 %), would give a corresponding SF<sub>6</sub> concentration of 1.15 instead of 0.46 fmol  $kg^{-1}$ . If any of these two scenarios are true, the excess-SF<sub>6</sub> is over-estimated with between 0.02 and 0.24 fmol kg<sup>-1</sup>. Although such an extremely low saturation as the latter has been observed in the Labrador Sea (Azetsu-Scott et al., 2003) it seems not to be a valid number in the Greenland Sea and the numbers are only mentioned here for comparison. Only the first scenario of 80 % saturation will be included in the uncertainty estimates and this indicates an offset of less than 0.1 fmol kg<sup>-1</sup>. The concentration of SF<sub>6</sub> in NSDW is not markedly off-set since observations both in 1997 and 2001 show levels close to those used as the source function and the levels in MEIW and NEAW are at least not much underestimated since these are based on newly ventilated water with 100 % saturation.

In summary the mean excess  $SF_6$  during the period of relatively constant level (years 2000-2002) would with the mentioned uncertainties be somewhere between 0.6 and 1.1 fmol  $kg^{-1}$ .

When estimating the uncertainty in flux of the released SF<sub>6</sub> through the FBC also the uncertainty in the temporal evolution has to be considered. This uncertainty appears mainly during two phases, the initial build-up and an eventual increase the last year. The arrival time of excess SF<sub>6</sub> in the FBC occurred at the earliest around November 1998 and at least before June 1999. The build-up phase is estimated to be 1.5 years long but with an extreme start and end this could vary  $\pm$  6 months. This also means that the remaining time will be between 2.5 and 2.75 years. If we on the other hand assume that the last observation (which in average is 0.3 fmol kg<sup>-1</sup> higher) is representative for the last nine months (from immediately after the second last observation) we will have three scenarios of the time evolution. Applying the interval in concentration presented above, the total export would range between 10 and 23 kg.

### 4.5. Overflow export in the other regions

Additional outflow of  $SF_6$  east of Iceland, i.e. across the Iceland-Faroe Ridge or the Wyville-Thomson Ridge is believed to be low since the major portion of water passing these is not dense enough to contain the released tracer. A smaller amount might exit over the Wyville-Thomson Ridge although the agreement between the observations and the model indicates that this should be of less importance since the modelled Faroe-Shetland Channel estimate is only slightly higher than the observational FBC estimate. No observation-based estimate of the export through the Denmark Strait has yet been made since no similar time series exists there although the first observation of the tracer at the western overflow has been

reported by Olsson *et al.* (2004). The model estimate for the Denmark Strait is presented by Eldevik *et al.* (2004).

### 5. Conclusions

The first arrival of the released sulphur hexafluoride in the Faroe Bank Channel is estimated to early 1999 both based on observations and on numerical modelling. This gives a transit time from the central Greenland Sea to the FBC of about 2.5 years. The total transport of released SF<sub>6</sub> through the FBC into the North Atlantic by the end of 2002 is calculated to be 16 kg from the observations and 15 kg based on the numerical model. This means that of the 320 kg released in the Greenland Sea in 1996, approximately 5 % had passed through the FBC 6.5 years later.

In the numerical model the main pathway of this water is the direct route of the Jan Mayen Current from the Greenland Sea into the Norwegian Sea. This is also consistent with repeated observations of the tracer on the eastern side of the Jan Mayen Ridge (Olsson *et al.*, 2004). The East Greenland Current's main role during this period is to 'feed' tracer from the Greenland Sea to the Denmark Strait (Eldevik *et al.*, 2004).

It has not been estimated how large fraction of the overflow through the Faroe Bank Channel that has been formed in the Greenland Sea although a considerable part of the tracer released there has exited the Nordic Seas through this passage. The transit can be accomplished in as short time as 2.5 years or less. However, the transit time might be different for different portions and only a fraction of the intermediate water leaves the Greenland Sea each year which is seen in the well retained SF<sub>6</sub> concentration (Messias *et al.*, 2004). A change in the ventilation of the Greenland Sea can hence be transferred to the surrounding basins rather quickly although its full effect is not seen until after some decades. Nevertheless, the Greenland Sea is seen to be important for the overflow through the Faroe Bank Channel and perhaps more comprehensive investigations of the tracer evolution in the Greenland Sea can give estimates also of the volume export of Greenland Sea Arctic Intermediate Water within the overflows. The composition and changes of the overflow are of great importance in examining how it can be affected by climate change and hence also its role for the thermohaline circulation and the climate of Northern Europe.

## Acknowledgements

We are grateful to Marie Persson and Johanna Balle for their effort in analysing the sulphur hexafluoride and Toste Tanhua for his contribution to the tracer data. This research has been supported by grants from the European Union's 5<sup>th</sup> Framework Programme project TRACTOR (contract EVK2-CT-2000-00080), the Swedish Research Council and the Norwegian Research Council project ProClim (155923/700). The crucial initiation by the release experiment was financed inside ESOP-2 and parts of the earlier investigations inside VEINS, both EU/MAST III projects. Finally we thank the three reviewers for their valuable comments which improved the manuscript.

#### References

- Anderson, L.G., Chierici, M., Fogelqvist, E., Johannessen, T., 2000. Flux of anthropogenic carbon into the deep Greenland Sea. Journal of Geophysical Research 105 (C6), 14339-14345.
- Azetsu-Scott, K., Jones, E.P., Yashayaev, I., Gershey, R.M., 2003. Time series study of CFC concentartions in the Labrador Sea during deep and shallow convection regimes (1991-2000). Journal of Geophysical Research 108 (C11), 3354, doi: 10.1029/2002JC001317.
- Bleck, R., Rooth, C., Hu, D., Smith, L.T., 1992. Salinity-driven thermocline transients in a wind- and thermohaline-forced isopycnic coordinate model of the North Atlantic. Journal of Physical Oceanography 22 (12), 1486-1505.

Blindheim, J., 1990. Arctic Intermediate Water in the Norwegian Sea. Deep-Sea Research A 37 (9), 1475-1489. Broecker, W.S., 1991. The great ocean conveyor. Oceanography 4, 79-89.

- Bu, X., Warner, M.J., 1995. Solubility of chlorofluorocarbon 113 in water and seawater. Deep-Sea Research I 42 (7), 1151-1161.
- Bullister, J.L., Weiss, R.F., 1983. Anthropogenic chlorofluoromethanes in the Greenland and Norwegian seas. Science 221 (4607), 265-268.
- Bullister, J.L., Wisegarver, D.P., Menzia, F.A., 2002. The solubility of sulfur hexafluoride in water and seawater. Deep-Sea Research I 49 (1), 175-187.

- Eldevik, T., Straneo, F., Sandø, A.B., Furevik, T., 2004. Pathways and export of Greenland Sea Water. In: Drange, H., Dokken, T.M., Furevik, T., Gerdes, R., Berger, W. (Eds.), Climate Variability in the Nordic Seas. Geophysical Monograph Series. American Geophysical Union, in press.
- Fogelqvist, E., Blindheim, J., Tanhua, T., Østerhus, S., Buch, E., Rey, F., 2003. Greenland-Scotland overflow studied by hydro-chemical multivariate analysis. Deep-Sea Research I 50 (1), 73-102.
- Furevik, T., Bentsen, M., Drange, H., Johannessen, J., Korablev, A., 2002. Temporal and spatial variability of the sea surface salinity in the Nordic Seas. Journal of Geophysical Research 107 (C12), doi: 10.1029/2001JC001118.
- Gao, Y., Drange, H., Bentsen, M., Johannessen, O.M., 2004. Simulating transport of non-Chernobyl <sup>137</sup>Cs and <sup>90</sup>Sr in the North Atlantic-Arctic region. Journal of Environmental Radioactivity 71, 1-16.
- Gascard, J.-C., Watson, A.J., Messias, M.-J., Olsson, K.A., Johannessen, T., Simonsen, K., 2002. Long-lived vortices as a mode of deep ventilation in the Greenland Sea. Nature 416 (6880), 525-527.
- Girton, J.B., Sanford, T.B., Käse, R.H., 2001. Synoptic sections of the Denmark Strait Overflow. Geophysical Research Letters 28 (8), 1619-1622.
- Hansen, B., Østerhus, S., 2000. North Atlantic Nordic Seas exchanges. Progress in Oceanography 45 (2), 109-208.
- Hansen, B., Turrell, W.R., Østerhus, S., 2001. Decreasing overflow from the Nordic seas into the Atlantic Ocean through the Faroe Bank Channel since 1950. Nature 411 (6840), 927-930.
- Hansen, B., Turrell, W.R., Østerhus, S., 2003. Indications and consequences of weakened Iceland-Scotland overflow. ICES Marine Science Symposia 219, 102-110.
- Harnisch, J., Eisenhauer, A., 1998. Natural CF<sub>4</sub> and SF<sub>6</sub> on Earth. Geophysical Research Letters 25 (13), 2401-2404.
- Hátún, H., Sandø, A.-B., Drange, H., Bentsen, M., 2004. Seasonal to decadal variations in the Faroe-Shetland inflow waters. In: Drange, H., Dokken, T.M., Furevik, T., Gerdes, R., Berger, W. (Eds.), Climate Variability in the Nordic Seas. Geophysical Monograph Series. American Geophysical Union, in press.
- JPOTS editorial panel. Processing of oceanographic station data, 1991. UNESCO, Paris, France, 138 pp.
- Law, C.S., Watson, A.J., 2001. Determination of Persian Gulf Water transport and oxygen utilisation rates using SF<sub>6</sub> as a novel transient tracer. Geophysical Research Letters 28 (5), 815-818.
- Law, C.S., Watson, A.J., Liddicoat, M.I., 1994. Automated vacuum analysis of sulphur hexafluoride in seawater: Derivation of the atmospheric trend (1970-1993) and potential as a transient tracer. Marine Chemistry 48 (1), 57-69.
- Maiss, M., Brenninkmeijer, C.A.M., 1998. Atmospheric SF<sub>6</sub>: Trends, sources, and prospects. Environmental Science & Technology 32 (20), 3077-3086.
- Messias, M.-J., Watson, A.J., Fogelqvist, E., Van Scoy, K.A., Tanhua, T., Olsson, K.A., 1999. The tracer release experiment. In: The Thermohaline Circulation in the Greenland Sea. ESOP-2, Final Scientific Report, Bergen, Norway.
- Messias, M.-J., Watson, A.J., Johannessen, T., Olsson, K.A., Smethie, W.M., Jr, Anderson, L.G., Bacon, S., Gascard, J.-C., Olafsson, J., Oliver, K.I.C., Swift, J., Rey, F., Simonsen, K., Ledwell, J.R., Budeus, G., 2004. The Greenland Sea Tracer Experiment: Formation, spreading and dispersion of intermediate waters of the Greenland Sea, Manuscript in preparation.
- Nilsen, J.E.Ø., Gao, Y., Drange, H., Furevik, T., Bentsen, M., 2003. Simulated North Atlantic-Nordic Seas water mass exchanges in an isopycnic coordinate OGCM. Geophysical Research Letters 30 (10), 1536, doi:10.1029/2002GL016597.
- Nøst, O.A., Isachsen, P.E., 2003. The large-scale time-mean ocean circulation in the Nordic Seas and Arctic Ocean estimated from simplified dynamics. Journal of Marine Research 61, 175-210.
- Olsson, K.A., Jeansson, E., Tanhua, T., Gascard, J.-C., 2004. The East Greenland Current studied with CFCs and released sulphur hexafluoride. Journal of Marine Systems, in press.
- Østerhus, S., Gammelsrød, T., 1999. The abyss of the Nordic seas is warming. Journal of Climate 12 (11), 3297-3304.
- Rhein, M., 1991. Ventilation rates of the Greenland and Norwegian Seas derived from distributions of the chlorofluoromethanes F11 and F12. Deep-Sea Research A 38 (4), 485-503.
- Straneo, F., Pickart, R.S., Lavender, K., 2003. Spreading of Labrador sea water: an advective-diffusive study based on Lagrangian data. Deep-Sea Research I 50 (6), 701-719.
- Tanhua, T., Olsson, K.A., Fogelqvist, E., 2004. A first study of SF<sub>6</sub> as a transient tracer in the Southern Ocean. Deep-Sea Research II, in press.
- Turrell, W.R., Slesser, G., Adams, R.D., Payne, R., Gillibrand, P.A., 1999. Decadal variability in the composition of Faroe Shetland Channel bottom water. Deep-Sea Research I 46 (1), 1-25.

- Walker, S.J., Weiss, R.F., Salameh, P.K., 2000. Reconstructed histories of the annual mean atmospheric mole fractions for the halocarbons CFC-11, CFC-12, CFC-113 and carbon tetrachloride. Journal of Geophysical Research 105 (C6), 14285-14296.
- Warner, M.J., Weiss, R.F., 1985. Solubilities of chlorofluorocarbons 11 and 12 in water and sea water. Deep-Sea Research 32 (12), 1485-1497.
- Watson, A.J., Ledwell, J.R., 2000. Oceanographic tracer release experiments using sulphur hexafluoride. Journal of Geophysical Research 105 (C6), 14325-14337.
- Watson, A.J., Messias, M.J., Fogelqvist, E., Van Scoy, K.A., Johannessen, T., Oliver, K.I.C., Stevens, D.P., Rey, F., Tanhua, T., Olsson, K.A., Carse, F., Simonsen, K., Ledwell, J.R., Jansen, E., Cooper, D.J., Kruepke, J.A., Guilyardi, E., 1999. Mixing and convection in the Greenland Sea from a tracer-release experiment. Nature 401 (6756), 902-904.

#### **Figure captions**

Fig. 1. (a) Main features of the surface currents (thick grey arrows) and intermediate-deep water currents (thin black arrows) in the Nordic Seas. The acronyms used are: EGC - East Greenland Current, EIC – East Icelandic Current, FC – Faroe Current, JMC – Jan Mayen Current, NIIC – North Icelandic Irminger Current, NwAC – Norwegian Atlantic Current.

(b) A map of the Nordic Seas with some key geographic features indicated with the following acronyms: DS – Denmark Strait, FBCh – Faroe Bank Channel, FI – Faroe Islands, FS – Fram Strait, FSCh – Faroe Shetland Channel, GS – Greenland Sea, IS – Iceland Sea, JM – Jan Mayen, JMCh, Jan Mayen Channel, JMR – Jan Mayen Ridge, MR – Mohns Ridge.

Fig. 2. Graphs showing correlation of transient tracer concentrations in seawater. (a) Functions of CFC-12 (solid line) and SF<sub>6</sub> (dashed line) versus time in a surface water of S = 34.89 and T = -0.7°C in equilibrium with the atmosphere.

(b) Function of CFC-12 versus SF<sub>6</sub> in the different water types resulting from mixing of the source waters according to Table 2 and Equation 1. The relationship in the years 1997 and 2001 in the three water types are indicated ( $\circ$  = upper,  $\blacktriangle$  = middle, + = lower).

Fig. 3. Potential temperature in °C (a), salinity (b), and potential density  $\gamma_{\theta}$  in kg m<sup>-3</sup> (c) on a section across the FBC in June 2000. Station locations are shown in Fig. 4. The positions of the two standard stations, V05 and V06, from which most of the SF<sub>6</sub> samples derive, are indicated.

Fig. 4. Map of the FBC (a), showing the standard CTD stations and  $\theta$ -S diagram (b) from CTD profiles acquired in the vicinity of standard stations V05 and V06 in the period 1998-2001. In (b), the  $\gamma_{\theta} = 28.049$  kg m<sup>-3</sup> isopycnal is identified. The overflow passes the section from the southeast to the northwest and also the additional sampled stations are located along this section but at slightly different positions than those showed.

Fig. 5. Observed SF<sub>6</sub> concentration in fmol kg<sup>-1</sup> versus time in the FBC at (a) V05 and (b) V06 respectively. The depth interval was chosen to include all observations of overflow water which are shown by dots, while observations from outside the overflow (> $0.3^{\circ}$ C) are marked by asterisks.

Fig. 6. The time evolution of excess SF<sub>6</sub> in the Faroe Bank Channel based on observations (markers) and model (solid line), in units of fmol kg<sup>-1</sup>. All observations with  $\theta < 0.3^{\circ}$ C, taken between November 1998 and June 2002 are included. It also shows lines representing the average estimated excess of the constant phase utilised to give the estimated outflow as well as the increase trend during the build-up phase.

Fig. 7. The model domain and IW streamlines. The contour interval corresponds to 0.5 Sv. The greyscales show the topography at 500 m intervals and the rectangle marks the site of the tracer release.

Fig. 8. Modelled (a) and observed (b) distribution of released  $SF_6$  integrated over the water column (nmol m<sup>-2</sup>) six years after the tracer release. The observed distribution is received by subtracting the  $SF_6$  estimated to be of atmospheric origin from the total observed and is from Messias *et al.* (2004) where the spreading is presented in more detail. The rectangles in both figures mark the site of the tracer release.

# Tables

Table 1. The sampled stations in the FBC including timing, location and determined tracers.

Year	Month	Ship	Positions <sup>1</sup>	Tracers <sup>2</sup>
1997	Aug	Aranda	61°20'N, 8°16'W; 61°22'N, 8°13'W, 61°25'N, 8°10'W; 61°28'N, 8°08'W & 61°30'N, 8°05'W	CFCs
1998	Nov	Magnus Heinason	V06	$SF_6$
1999	Feb	Magnus Heinason	V05 & V06	$SF_6$
	June	Magnus Heinason	V05 & V06	$SF_6$
2000	Feb	Magnus Heinason	V05 & V06	$SF_6$
	June	Magnus Heinason	V05 & V06	$SF_6$
2001	Feb	Magnus Heinason	V05 & V06	$SF_6$
	June	Håkon Mosby	V05, V06, 61°15'N, 8°01W; 61°18'N, 7°57'W & 61°21'N, 7°50'W	SF <sub>6</sub> , CFCs
	Sep	Magnus Heinason	V05 & V06	$SF_6$
2002	Feb	Magnus Heinason	V05 & V06	$SF_6$
	June	Magnus Heinason	V05 & V06	$SF_6$
2003	Feb	Magnus Heinason	V05 & V06	$SF_6$

<sup>1</sup> The positions for V05 and V06 are 61°20'N, 07°53'W and 61°16'N, 08°00'W respectively (see Fig. 4).

<sup>2</sup> In addition to the listed tracers T and S were determined on all cruises.

Table 2. Properties of the three different water types in the FBC as follows: potential temperature, salinity and fractions of the four source waters. The properties of the source waters are taken from (Fogelqvist *et al.*, 2003) except for those of GSAIW which are taken from observations during ESOP-2.

				Source waters		
Water types in FBC	Θ	Salinity	GSAIW,	NSDW,	MEIW,	NEAW,
	°C		fractions	fractions	fractions	fractions
Upper	-0.016	34.900	0.35	0.50	0.10	0.048
Middle	-0.482	34.902	0.20	0.72	0.060	0.023
Lower	-0.638	34.907	0.11	0.83	0.040	0.020

The following acronyms are used: GSAIW – Greenland Sea Arctic Intermediate Water, NSDW – Norwegian Sea Deep Water, MEIW – Modified East Icelandic Water, NEAW – North East Atlantic Water.

















