Tritium transfer between sea and atmosphere in the English Channel (North Cotentin and Bay of Seine)

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Abstract. Oceans, seas, estuaries and rivers represent a vast sink for many substances of anthropic origin (metals, radionuclides, etc…). Depending on their chemical form, artificial radionuclides discharged into the sea by the nuclear industry can be carried onto land in marine aerosols, as well as by degassing seawater as is the case for tritium (³H) or for radiocarbon (¹⁴C). Three oceanographic cruises in the English Channel: TE-SEA, TRANSAT 1 and 2 have been performed on R.V. "Côtes de la Manche" to quantify the flux of ³H in gaseous form, from the sea into the atmosphere, following particular discharge into the sea from the COGEMA spent fuel reprocessing plant at La Hague (North West France). During cruises, the maximum concentration measured in the air was 10.6 Bq.L⁻¹, which is distinctly higher than the background of 1 Bq.L⁻¹ thus demonstrating the transfer of ³H between water and atmosphere. The mean flux of ³H between the water and the atmosphere, calculated during these cruises was 2.4 10⁷ Bq.km⁻².d⁻¹. For the Seine Bay area (4400 km²) this flux represents 39 TBq.yr⁻¹ and hence less than 0.3 % of ³H discharged into the ocean from the COGEMA spent fuel reprocessing plant at La Hague.

1. INTRODUCTION

Oceans, seas, estuaries and rivers represent a vast sink for many substances of anthropic origin (metals, radionuclides, etc…). Depending on their chemical form, artificial radionuclides discharged into the sea by the nuclear industry can be carried onto land in marine aerosols, as well as by degassing seawater as is the case for tritium (³H) or for radiocarbon (¹⁴C) [1, 2]. Knowledge of these exchanges should lead to quantification of the radioactivity discharged into the atmosphere then transferred into the soil and plants and to man. In June 2000 and February and August 2002, the Laboratoire de Radioécologie de Cherbourg-Octeville (LRC, Cherbourg, France) in association with the Laboratoire des Sciences du Climat et de l’Environnement (LSCE, Gif sur Yvette, France) and the French Navy Groupe d’Etudes Atomiques (GEA, Cherbourg, France), carried out oceanographic cruises TE-SEA, TRANSAT 1 and 2 on the R.V. "Côtes de la Manche", to quantify the flux of gaseous ³H from the sea into the atmosphere, between the point of discharge into the ocean from the COGEMA spent fuel reprocessing plant at La Hague and the Seine estuary. This paper presents the results of these cruises.
2. EQUIPMENT AND METHOD

2.1 Method

Discharge into the ocean from the COGEMA spent fuel reprocessing plant at La Hague (Nord Cotentin, France) is the main source of $^3$H in the English Channel [3]. For example, in 2002, this plant discharged $1.2 \times 10^{16}$ Bq yr$^{-1}$ of $^3$H into the sea and during the same year, atmospheric discharge was $6.3 \times 10^{15}$ Bq yr$^{-1}$ of $^3$H [4].

To evaluate the transfer of $^3$H between water and atmosphere due to discharge into the sea by COGEMA La Hague, the flux of water vapour (L m$^{-2}$ d$^{-1}$) is calculated first, using Rohwer's formula [5]:

$$\Phi(\text{H}_2\text{O}) = 0.372 \times (1-0.000374 \times \text{Pa}) \times (1+0.6 \times V_w) \times (e_s-e_a)$$

where:
- $\Phi(\text{H}_2\text{O})$: Flux of water vapour (L m$^{-2}$ d$^{-1}$),
- Pa: Atmospheric pressure (mbar),
- $V_w$: Wind speed (m s$^{-1}$),
- $e_s$: Saturated vapour pressure (mbar),
- $e_a$: Vapour pressure in the atmosphere (mbar).

Then, knowing the concentration of $^3$H in the water, the flux of $^3$H (Bq m$^{-2}$ d$^{-1}$) is determined using equation 2:

$$\Phi(\text{H}_2\text{O}) = \Phi(\text{H}_2\text{O}) \times [^3\text{H}_w]$$

where:
- $\Phi(\text{H}_2\text{O})$: Flux of $^3$H between water and atmosphere (Bq m$^{-2}$ d$^{-1}$),
- $[^3\text{H}_w]$: Concentration of $^3$H in the water (Bq L$^{-1}$).

2.2 Sampling of seawater and air

In 2000 and 2002, three oceanographic cruises were carried out with the INSU/CNRS Oceanographic Vessel "Côtes de la Manche" (Figure 1) between 1 and 4 June 2000 (TE_Sea) and between the 24 and 28 February 2002 (TRANSAT 1) and 27 to 31 August 2002 (TRANSAT 2).

Figure 1. Location of sampling stations during TE_Sea, TRANSAT 1 and TRANSAT 2.
The sampling periods reflect roughly the different meteorological patterns encountered during the year.

To avoid being under the atmospheric plume of the COGEMA La Hague plant, samples were taken upwind of the plant. During the cruises, close to COGEMA La Hague (< 50 km), the activity of krypton 85 (85Kr), a gas discharged by the plant into the atmosphere at the same time as 3H, was also measured continuously at one second intervals in order to verify the presence of atmospheric plume. The presence of 85Kr discharged from the plant stack was detected immediately [6]. Samples for measuring 3H in the air and seawater were therefore taken when the influence of the plant plume was undetectable.

Seawater was sampled after filtration at 0.45 µm on the surface, in one litre polyethylene flasks. The water vapour was condensed on a cold trap (PREVAIR system) developed specifically by IRSN (LRC) for this study in association with the French Navy (GEA). This sampling system allows the limit of detection to be lowered when measuring 3H by a factor of 3000 compared with sampling systems using bubble tubes. This sampling, which takes about fifteen minutes, was performed two metres above the surface of the water.

Water temperature and salinity were measured with a Seabird SBE 19-03 probe. Air temperature, humidity, atmospheric pressure, wind speed and direction were measured using the “Côtes de La Manche” meteorological station (Young equipment).

2.3 Measurement of 3H

The seawater was purified by distillation to eliminate any radionuclides which could interfere with measuring 3H. The distilled seawater samples and water vapour samples from the air were mixed with a scintillating liquid in a standard metering flask before being transferred to a liquid scintillation counter for measurement. Measurements were made in the French Navy's low background laboratory at Cherbourg (Roule Laboratory).

3. RESULTS AND DISCUSSION

3.1 Concentrations of 3H in seawater and atmospheric water vapour

The concentrations of 3H in the seawater and atmospheric water vapour measured during the TE-SEA and TRANSAT 1 and 2 cruises are presented in figures 2 to 4.

The mean concentrations measured in seawater were 14.8 Bq.L⁻¹ (maximum 20.9 Bq.L⁻¹) during TE-SEA, 5.7 Bq.L⁻¹ (maximum 13.3 Bq.L⁻¹) during TRANSAT 1 and 4.5 Bq.L⁻¹ (maximum 6.4 Bq.L⁻¹) during TRANSAT 2. In comparison with the background in 3H, around 0.15 Bq.L⁻¹ at the entrance to the Channel, these measurements indicate industrial seawater labelling by mainly the COGEMA La Hague spent fuel reprocessing plant [7]. For the same points, the mean concentrations measured in atmospheric water vapour were 7.0 Bq.L⁻¹ (maximum 10.1 Bq.L⁻¹) during TE-SEA, 2.8 Bq.L⁻¹ (maximum 4.8 Bq.L⁻¹) during TRANSAT 1 and 2.4 Bq.L⁻¹ (maximum 3.8 Bq.L⁻¹) during TRANSAT 2. In comparison with the 3H background, below 1.0 Bq.L⁻¹ in air, these measurements indicate air labelling [3]. Since the samples were taken upwind, outside the atmospheric plume from the COGEMA La Hague plant, these results demonstrate that 3H is transferred from the water into the atmosphere by evaporating seawater.

The highest concentrations in the air were measured during the TE-SEA cruise in an area close to the COGEMA La Hague plant ocean discharge point. In general, for all the cruises, the evolution of concentrations in the air follows the evolution of concentrations in water. The observed differences are due to different meteorological conditions at the time, wind speed and air hygrometry.
**Figure 2.** Variation of $^3$H activities (Bq.L$^{-1}$) in surface seawater and air versus distance from COGEMA outlet pipe during TE-SEA cruise.

**Figure 3.** Variation of $^3$H activities (Bq.L$^{-1}$) in surface seawater and air versus distance from COGEMA outlet pipe during TRANSAT 1 cruise.

**Figure 4.** Variation of $^3$H activities (Bq.L$^{-1}$) in surface seawater and air versus distance from COGEMA outlet pipe during TRANSAT 2 cruise.
3.2 Flux of $^3$H between water and atmosphere

The water vapour and $^3$H flux determined during the TE-SEA, TRANSAT 1 and 2 cruises are given in figures 5 to 7.

The mean water vapour flux measurements obtained during these cruises was $2.7 \times 10^6$ kg.km$^{-2}$.d$^{-1}$ (TE-SEA), $3.2 \times 10^6$ kg.km$^{-2}$.d$^{-1}$ (TRANSAT 1) and $2.8 \times 10^6$ kg.km$^{-2}$.d$^{-1}$ (TRANSAT 2), corresponding to a mean flux of $^3$H from water to atmosphere of $4.0 \times 10^7$ Bq.km$^{-2}$.d$^{-1}$ (TE-SEA), $2.0 \times 10^7$ Bq.km$^{-2}$.d$^{-1}$ (TRANSAT 1) and $1.3 \times 10^7$ Bq.km$^{-2}$.d$^{-1}$ (TRANSAT 2). Generally, $^3$H flux between water and atmosphere falls with distance from the discharge point, reflecting the dilution of liquid waste in the Channel waters. The mean flux was slightly higher during TE-SEA in the area close to the COGEMA La Hague point of discharge into the sea. During TRANSAT 1 the mean flux was higher than during TRANSAT 2 because the windspeed was higher (factor 2). The mean flux calculated for all the cruises was $2.4 \times 10^7$ Bq.km$^{-2}$.d$^{-1}$. Over an area corresponding to Seine Bay (4400 km$^2$) the quantity of $^3$H transferred into the atmosphere per year can be evaluated at 39 TBq i.e. 0.3 % of the quantity of $^3$H discharged into the sea in 2002 by the COGEMA La Hague plant.

![Figure 5](image1.png)

**Figure 5.** Variation of water vapour fluxes (a) and $^3$H fluxes (b) versus distance from COGEMA outlet pipe during TE-SEA cruise.

![Figure 6](image2.png)

**Figure 6.** Variation of water vapour fluxes (a) and $^3$H fluxes (b) versus distance from COGEMA outlet pipe during TRANSAT 1 cruise.
Figure 7. Variation of water vapour fluxes (a) and $^3$H fluxes (b) versus distance from COGEMA outlet pipe during TRANSAT 2 cruise.

4. CONCLUSION

The oceanographic cruises TE-SEA, TRANSAT 1 and 2 performed in the English Channel revealed atmospheric labelling with $^3$H following the evaporation of seawater labelled with tritiated waste from the COGEMA La Hague spent fuel reprocessing plant. The mean flux of water vapour and $^3$H between water and atmosphere was evaluated respectively at $2.9 \times 10^6$ kg km$^{-2}$ d$^{-1}$ and $2.4 \times 10^7$ Bq km$^{-2}$ d$^{-1}$, which corresponds an annual quantity of 39 TBq transferred into the atmosphere by this process, i.e. 0.3% of the quantity of $^3$H discharged into the sea in 2002 by COGEMA La Hague. This method must be generalised to provide a global estimate of the exchanges of $^3$H between water and atmosphere for the whole of the English Channel.

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