TRITIUM IN THE EASTERN MEDITERRANEAN SEA

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A hydrographic survey conducted in the eastern Mediterranean within the framework of the POEM program during 1988/89 included the measurement of tritium in the water samples. Surface concentrations of tritium have decreased considerably relative to 1968, when values ranged up to 30 TU (ASSAF, 1968). Surface tritium values during the present survey were close to 5 TU near the Turkish coast and were even lower in the southern part of the eastern basin, ranging from 2.7-3.7 TU (Fig. 1). In the north-eastern part of the basin values of 3-5 TU persisted to a depth of 300-500 meters. In the central part of the basin, around Crete, such high tritium levels were encountered at much greater depths, down to 1000-1500 meters. A well defined step structure of the vertical profile was noted at the more southerly stations, with intermediate values of around 2 TU at a depth of 300-600 meters. Furthermore, we observed a deepening of this step-layer from east to west (from 300-400 meters at Station 9 to 400-600 meters at Stations 14 and 10).

There is little correlation between tritium values and parameters such as d18O, temperature and salinity (GAT et al., 1994), indicating that it is not simply local vertical mixing which results in penetration of tritium to deeper layers. The intermediate "step" in the tritium profiles is probably related to the Levantine Intermediate Waters (LIW) which originate in the eastern part of the basin. The relatively high tritium levels found down to the deeper waters in the area near Crete, may be indicative of the process of deep water formation at this site (ROETHER and SCHLITZER, 1991).

The tritium data in this period of declining atmospheric tritium levels have been found useful in delineating water masses and suggesting genetic relationships among them. The simultaneous measurement of helium-3 (3He) and of tritium (3H), which provides a time of closure (age) for the waters, is planned for the upcoming phase of the POEM program in December 1994. This can be expected to clarify the relationships between the different water bodies which were delineated in the eastern Mediterranean water column.

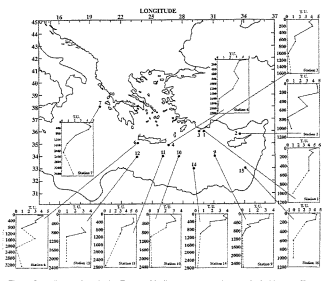


Fig. 1. Sampling stations in the Eastern Mediterranean and their vertical tritium profiles. Tritium concentrations are in tritium units (TU). Stations 1, 2 were sampled in July '88, stations 3,4,6 and 7 in September '88 and stations 9, 10,11,12, 14 and 15 in March '89.

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137CS CONCENTRATION IN BED LOAD SEDIMENTS FROM THE DANUBE RIVER AND THE BLACK SEA DURING 1993

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In a previous study mathematical modelling of man-made radionuclides transfered and transported in the Danube river was undertaken by GEORGESCU (1986). This paper presents data on the variability of 137 Cs transported by the bed load sediments of the Danube river, Danube Delta and the Black Sea during 1993

Sampling of the bed load sediments was performed simultaneously with hydrological and solid discharge measurements made during specific hydrological periods, i.e. during spring, summer and autumn at high and relatively low flow rates in the following cross-sections: Svinitza, Orsova, before the Turnu-Severin dam, Bechet (in front of the Kozlodui Nuclear Power Plant), Giurgiu, Ceatal-Izmail (beginning of the Danube Delta), Chilia and Sulina (Danube Delta) and Portiza and East Constantza in the Black Sea. About 2 kg of each sediment sample were dried in an electric oven at 105°C, homogenized and analyzed by gamma spectrometry with

a HPGe detector for 18-20 h. The ¹³⁷Cs activity in the sediments sampled is presented in Table 1. Concerning the presence of ¹³⁴Cs, it was only identified at locations with higher ¹³⁷Cs activities, e.g. up to 24 ± 1 Bq/kg ¹³⁴Cs at Sulina.

Table 1. Contamination by ¹³⁷Cs of bed load sediments from the Danube river, Danube Delta and Black Sea during 1993.

Location and date of sampling		Activity Bq/kg (dry)	Location and date of sampling		Activity Bq/kg (dry)
Svinitza	8.04 8.08	114 ± 3 107 ± 5	Ceatal-Izmail	24.04 30.08	1.6 ± 0.5 13.5 ± 1.9
Orsova	8.04 18.08	81 ± 2 193 ± 10	Chilia km. 3	5.09	9.6 ± 0.8
Turnu-Severin	9.05 19.08	3.4 ± 0.5 < 0.6	Sulina* Sulina**	16.07 30.05	17.0 ± 0.7 171 ± 5
Bechet km. 705	17.04 31.08	4.8 ± 0.4 3.1 ± 0.3	Portitza	18.04 19.07	50 ± 2 22.4 ± 0.7
Bechet km. 678	$17.04 \\ 21.08$	$\begin{array}{c} 2.8 \pm 0.3 \\ 2.0 \pm 0.5 \end{array}$	East Constantza	5.06 13.07	125 ± 6 253 ± 5
Giurgiu	$\begin{array}{c} 13.04 \\ 23.08 \end{array}$	$\begin{array}{c} 2.1 \pm 0.3 \\ 1.0 \pm 0.2 \end{array}$			

*) 2.5 nautical miles from Sulina **) 26 m depth in the Black Sea in front of Sulina Port

With respect to the ¹³⁷Cs radioactivity levels, the Danube river and Black Sea coast can be divided into the following zones: Svinitza-Orsova (1st zone), Turnu-Severin dam - Ceatal-Izmail (2nd zone), Danube Delta (3rd zone) and Black Sea (4th zone). The lowest and nearly constant values were observed in the second zone where there are no important pollution sources of ¹³⁷Cs. In the first zone which includes the entrace of the Danube into Romania, ¹³⁷Cs

activities are about 50-100 times higher than those observed in the second zone. The highest ¹³⁷Cs activities were measured at the mouth of Danube river (Sulina) as well as south of the Danube Delta at Portitza and Constantza on the Black Sea. This may be explained by contaminated waters being transported in a southerly direction by the northeast marine currents.

To calculate radionuclide transport by the bed load sediments between two time intervals, the following relation has been used :

$$C_{1} = Q_{b}^{i} \cdot C_{b}^{i}, \quad i = 1,2 \text{ (time periods)} \quad (1)$$

$$C_{e} = \frac{C_{1} + C_{2}}{2} \cdot \Delta t \quad (2)$$

where Q_b is solid discharge (kg/s) C_b is activity (Bq/kg) and Δt is the time interval between the two measurements. For example, at the Giurgu cross section with $Q_b^1 = 18.4$ kg/s, $Q_b^2 = 10$ kg/s, $C_b^1 = 2.1$ Bq/kg, $C_b^2 = 1.0$ Bq/kg (see table 1), the total transported ¹³⁷Cs activity, during 132 days is 2.7 x 10⁸ Bq. Spatial and temporal variation of the natural radioactive series U-Ra and Th will be the relativity of a program (CECOURSCU).

be the subject of a separate paper (GEORGESCU, in prep.).

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