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Radioactivity from Lisboa urban wastewater discharges in the Tejo River Estuary *

Radioactividade no estuário do Tejo proveniente das descargas de águas residuais de Lisboa

Margarida Malta 1, João M. Oliveira 1, Lídia Silva 1, Fernando P. Carvalho @1

Abstract

Sediments, water, mussels and fish were collected monthly during a 19 month period at eight sampling stations along the North bank of the Tejo estuary (Lisboa, Portugal). Fresh samples were immediately analysed for short lived gamma emitting radionuclides. Amongst these, 131I and 99mTc were detected in most samples of mussels (Mytilus galloprovincialis) and mullets (Liza ramada). Spatial distribution of samples containing these radionuclides consistently indicated contamination at Alcântara and Praça do Comércio, and, on occasion, at Chelas, all sites inside the estuary. Round-the-year, generally there was no contamination at Algés, near the estuary mouth, and Beirolas, upstream by the Vasco da Gama Bridge. The estuarine areas found contaminated by radionuclides correspond to the areas receiving urban wastewater discharges from the city of Lisboa. The artificial radionuclides 131I and 99mTc are used in nuclear medicine practices and originate from hospital liquid effluent discharges into the city sewage system. Their presence near Praça do Comércio was related to the nearby discharges of a main sewer discharging untreated urban wastewater. Near Alcântara, the sewer outlet discharged treated urban wastewater from the Alcântara wastewater treatment plant and the presence of short-lived radionuclides indicated that sewage treatment and the time delay introduced before discharge of the treated effluent were not sufficient to allow for radioactive decay of those radionuclides. The highest concentration values of 131I and 99mTc determined in mussels and fish were 12±2 and 136±20 Bq kg⁻¹ wet weight (w.w.), respectively. These concentrations did not exceed the concentrations of naturally-occurring radionuclides, such as 40K that averaged 66±13 Bq kg⁻¹ (w.w.) in mussels and 618±21 Bq kg⁻¹ (w.w.) in fish. Nevertheless, artificial radionuclides accumulated in biota increase the radiation dose received by organisms from natural radiation sources and may also increase the risk of radionuclide transfer to humans. Reduction of radioactivity in effluent discharges towards improved estuarine water and sediment quality is debated as a requirement for sustainable use of the estuary.

Keywords: Technetium-99m, Iodine-131, Cesium-137, Natural radionuclides, Radioactive contamination, Urban wastewater

Resumo

Amostras de sedimento, água, mexilhão e peixe foram colhidos mensalmente durante um período de 19 meses em várias estações ao longo da margem norte do estuário do Tejo (Lisboa, Portugal). As amostras foram imediatamente analisadas para radionuclídeos emissores gama, em especial para os de origem artificial. Entre estes, o 131I e 99mTc foram detectados na maioria das amostras de mexilhões (Mytilus galloprovincialis) e de taínha (Liza ramada). A distribuição espacial das amostras contendo esses radionuclídeos indicou contaminação continuada em Alcântara e Praça do Comércio, e, nalgumas ocasiões, também em Chelas. Ao longo do ano, em geral não se detectou contaminação em Algés, na entrada do estuário.

Keywords: Tecnéutio-99m, Iodeto-131, Césio-137, Radionuclídeos naturais, Contaminação radiactiva, Esgotamento urbano

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1. INTRODUCTION

In nuclear medicine, radioisotopes are used both for diagnostic and therapeutic purposes. Currently, about 20 radioisotopes are produced for use in nuclear medicine, such as iodine ($^{131}$I, physical half-life $T_{1/2} = 8.0$ d), technetium ($^{99m}$Tc, $T_{1/2} = 60.01$ h), chromium ($^{51}$Cr, $T_{1/2} = 27.7$ d), gallium ($^{68}$Ga, $T_{1/2} = 3.26$ d) and cobalt ($^{58}$Co, $T_{1/2} = 70.96$ d) (Farrington 1996). In Lisboa, several hospitals and other medical facilities daily use some of these radioisotopes imported from main international radiopharmaceutical suppliers. From this use, hospital solid waste and liquid effluents containing radioactivity are produced. Radioisotope measures are implemented in the medical facilities according to international standards, to prevent or reduce the irradiation and contamination of the staff and facilities. Those measures include procedures for solid waste segregation and safe disposal, and procedures for liquid waste management including special bathrooms for patients under treatment with radiopharmaceuticals.

The radioactive liquid effluents produced at the medical facilities (from patient bathrooms and laboratory sinks) may contain relatively high levels of radioactivity depending on the clinical specialties in the facilities, amount of radioisotopes applied, and number of patients treated. The discharge of radioactive liquid effluents from medical facilities to the environment has been a matter of some concern and investigation in several large European cities leading to enforcement of regulations and inspections to control better radioactive discharges (Mundschunk, 1996; Barquero et al., 2008; Bergman et al., 2008; Fischer et al., 2009; Krawczyk et al., 2013). In the case of coastal discharges through marine outfalls the environmental radioactivity enhancement has also been assessed in some areas with a view to protection of fisheries and human consumers against contamination and to the protection of non-human biota against radiation (Carolan et al., 2011). Often, environmental radioactivity surveys of such areas are carried out to check compliance with radiation dose limits to members of the public (EU, 1996).

In Lisboa, some hospitals are equipped with tanks for temporary storage of radioactive liquid effluents in order to allow for radioactive decay before release into the city sewage system. Nonetheless, not every medical facility performs radioactivity analysis of the effluents and keeps records of radionuclide composition and amounts of radioactivity released into the Lisboa sewage system. Besides, patients can go home and use the ordinary sewage collection and treatment works from there. After collection in the sewers system, urban wastewaters are mostly treated. Treated wastewaters plus still untreated urban wastewater from some city areas were discharged through several sewer outlets into the Tejo estuary in the years of the sampling (Figueira and Monteiro 2001; Carvalho et al. 2002).

To assess the radioactivity levels in the Tejo estuary resulting from the sewage discharges from Lisboa, an assessment experiment was established in collaboration with the Lisboa municipal administration. Sampling took place in 2004-2005, and was based upon the use of estuarine aquatic compartments (biotic and abiotic) as indicators of contamination along the north bank.

2. MATERIALS AND METHODS

Sampling

Monthly sampling was carried out during a 19-month period, from June 2004 to December 2005, with the help of the Lisboa Fire Brigade. Sampling stations were selected along the shore, from Algés near the maritime traffic control tower at the sea entrance of the estuary, to the mid estuary near Sacavém in front of the Nuclear and Technological Institute (ITN). Most stations were located between the 25th April and the Vasco da Gama bridges. Locations were selected near the main urban wastewater discharges (sewer outlets) and in between them in places with walls or piers with intertidal mussel beds (Figure 1).

Water, bottom sediment, mussels (Mytilus galloprovincialis), and mullets (Liza ramada) were sampled at low tide. Both species were selected for monitoring because they are common biota of the Tejo estuary. Mussels are sessile bivalves eating upon suspended particulate matter that they filter from water, thus integrating and averaging contamination levels at the same place, while mullets are pelagic fish swimming up and down the estuary and are opportunistic feeders. Surface water samples were collected at about 3-5 m from the shoreline, where water depth
may vary from 2 to 5 meters, directly into polyethylene cans, using a nylon net filter (300 µm mesh) to avoid large particles. Sub tidal sediment samples were collected at the same locations, using a bulk sediment sampler operated from the boat and collecting the top 10 cm sediment layer. Mussels were collected always at the same beds. Mullets were collected from the boat and collecting the top 10 cm sediment layer. Typically, size of samples collected was 2-4 litres of sediment, 10 litres of water, 3 kg of mussels and 12 specimens of adult mullets. Samples were conditioned in plastic containers, identified, and transported in ice chests to the laboratory for immediate processing and radiometric analysis. Not every month all desired sample types could be collected due to occasional adverse weather and tide conditions in the estuary, but the rate of successful sampling was high (>90%).

Radioactivity analysis

Analyses of two types were performed. One, was immediately performed by gamma spectrometry in order to allow for measuring the short-lived gamma emitting radionuclides used in nuclear medicine; the other was performed on oven-dry (60°C) and homogenized sample materials, and included alpha and gamma spectrometry for determining naturally occurring radionuclides with longer physical half-lives (T1/2). Immediate gamma radiation measurements were performed on fresh material (in toto) placed in 1.0 L capacity Marinelli beakers on HpGe solid state detectors (Canberra). Gamma spectra were acquired for 1 hour and radionuclides identified and quantified using the software package Genie2000 (Canberra). These measurements allowed the determination of radionuclides such as 131I and 99mTc.

For deferred analyses, animal samples were treated by dissecting and separating fresh tissues from shells in the case of mussels (Mytilus galloprovincialis), and separating muscle, liver, and gonads in the case of mullets (Liza ramada). Following oven drying and homogenization, sample aliquots were taken for gamma ray measurement using long counting times (24 h) for quantification of 40K and 137Cs. Separate aliquots were used for complete dissolution with HCl and HNO3 acids followed by radiochemical separation and analysis of 210Po and 210Pb by alpha spectrometry. Briefly, 210Po was analyzed after addition of an internal isotopic tracer (209Po) to the sample, and 210Pb was determined after in-growth of 210Po from the parent radionuclide 210Pb and second 210Po plating on a silver disc (Carvalho 1995; Carvalho et al., 2010). Alpha radioactivity measurements of the alpha radiation emitted by the silver disc were performed with ion implanted silicon detectors and alpha spectrometers OctetePlus (ORTEC EG&G) for determination of polonium-210 and lead-210.

A rigorous analytical quality control was maintained through the analysis of certified reference materials using the same methods and through participation in interlaboratory comparison exercises (Pham et al. 2006; Povinec et al., 2007). Results are expressed in Bq kg-1 wet weight (w.w.) for mussels and fish, Bq kg-1 dry weight (d.w.) for sediment, and in Bq L-1 for water, unless stated otherwise.

The global assessment of radionuclide concentrations in several environmental compartments was made averaging all results per radionuclide and per sample type (Table 1). For computation of the mean concentration for each radionuclide, samples with radionuclide concentrations below the lower limit of detection were ascribed a value of zero and used in the mean computation.

3. RESULTS AND DISCUSSION

Gamma spectrometry analyses of fresh materials (biotic and abiotic) consistently allowed the identification and quantification of several artificial and natural radionuclides (Figure 2). The spectrograms show the gamma peaks of 131I, and 99mTc, short-lived gamma emitting radioisotopes used in nuclear medicine. 137Cs (artificial radionuclide from fallout of nuclear weapon tests and nuclear accidents) and 40K (naturally-occurring radionuclide) were detected as well. Long lived artificial gamma emitting radionuclides other than 137Cs generally were not detected in sediments.
Alcântara. These two sampling stations were located closer to urban wastewater discharges. During the years of sampling at Alcântara Mar dock, a sewer outlet discharging treated urban wastewater from Alcântara wastewater treatment plant, and at Praça do Comércio a sewer outlet discharging untreated urban wastewaters, were fully operational (Figure 1).

I-131 ($T_{1/2}= 8.0$ d) has a longer half–life than $^{99m}$Tc ($T_{1/2}= 6.01$ h), which allows longer-lasting transport by tidal currents from the discharge points a few km along the shore either upstream or downstream. This was shown by positive radioactivity results from Algés to Beiroras, although radionuclide concentrations displayed decreasing frequency and decreasing concentration values from the stations at Alcântara and Praça do Comércio to both ends of the city area (Figures 1 and 3). At station 8, near Sacavém, no biota samples were available and only water and sediments could be analysed. Tc-99m was the only radionuclide of artificial origin detected there, and in low concentrations.

Concentrations of $^{131}$I in sediments were in average lower than those of technecium, which is in line with known Kd sediment-water partitioning coefficients (Kd= Bq kg$^{-1}$ in sediment dry weight/ Bq L$^{-1}$ in water) reported in literature: Kd values are of about 70 for iodine and $10^{2}$-$10^{3}$ for technetium in the marine environment (IAEA, 2004). Cesium-137 was rarely detected and always in low concentrations in sediments; furthermore it would very difficult to measure $^{137}$Cs in small volume water samples due to high sorption onto sediments of this radionuclide, with Kd values of $10^{4}$-$10^{5}$ (IAEA, 2004).

At the same sampling station radionuclide concentrations showed wide variations over time, reflecting also the fluctuation of radioactivity in wastewater discharges Figure 4). Mussels were the environmental compartment that showed the highest number of positive results (i.e., displaying radionuclide concentrations above detection limit and quantifiable) for artificial radioactive contamination. Practically, in most stations there were always mussels contaminated with $^{131}$I, and often with $^{99m}$Tc also. This is due to the filter-feeding activity of these molluscs that filter large water volumes to feed upon suspended organic particles. It is interesting to notice that the mullets concentrated $^{99m}$Tc from contaminated water to values sometimes much higher than those determined in mussels (Figure 4).

Radioactive iodine and technetium are present also in waste discharges from nuclear fuel reprocessing plants in UK and France into the marine environment, and have been subject to research and environmental surveys (Keogh and al., 2007; Smith et al., 2001). For example, $^{99m}$Tc from these waste discharges was consistently measured over several years along the East coast of Ireland in concentrations from 19-45 mBq/L in seawater. Seaweeds accumulate technetium to high concentrations with concentration factors relative to water of about $1.2x10^{5}$ on dry weight (d.w.) basis (CF=Bq kg$^{-1}$ d.w. seaweed/ Bq L$^{-1}$ filtered sea water), while the concentration factor on a wet weight basis for the edible part of mussels was from about 500 to near 1000, and for fish muscle was around 12 (Smith et al., 2001).While these isotopes measured in the Irish sea are inorganic forms of Tc and I, the radioisotopes in samples from the Tejo estuary are most likely still bound to the radiopharmaceutical organic molecules. This organic

Figure 2. Examples of gamma spectra of estuarine samples: A, Water; B, Mussels; C, Sediment. The peaks of main radionuclides are identified. Radionuclides of anthropogenic origin are $^{131}$I, $^{99m}$Tc and $^{137}$Cs. All other gamma-ray peaks correspond to naturally-occurring radionuclides.

Figura 2. Exemplo de espectros de radiação gama de amostras do estuário do Tejo: A, água; B mexilhões; C, sedimento. São identificados os picos principais de radionuclídeos emissores de radiação gama. Os radionuclídeos de origem antropogénica são o $^{131}$I, $^{99m}$Tc e $^{137}$Cs. Os restantes picos correspondem a radionuclídeos de origem natural.
Figure 3. Frequency of occurrence (%) of artificial radioisotopes in estuarine samples for each sampling station in the Tejo estuary during the 19-month survey period (2004-2005).

binding might contribute to a much higher accumulation in mullet than in mussels, i.e., the reverse than observations made in Irish Sea on the bioaccumulation of inorganic technetium and iodine. This observation indicates that bioaccumulation of radiopharmaceutical residues by aquatic biota and food chain transfer in the estuarine environment might be different than bioaccumulation of inorganic forms of the same radioelements. Most likely, current knowledge obtained with studies on the environmental behaviour of inorganic I and Tc does not apply to residues of radiopharmaceuticals labelled with these radionuclides. These aspects need further investigation.

Po-210 concentrations determined in mussels’ soft tissues and in tissues of the mullets *Liza ramada* were not identical (Figure 5). Instead, they varied with the tissue and between species, as expected. The fish muscle displayed the lowest concentrations measured in fish tissues, which for fish filet consumers reduces the transfer of radionuclides, although consumption of other organs such as gonad would transfer much higher activity. It must be said that concentrations determined for naturally-occurring radionuclides, such as 210Po, 210Pb and 40K in biota from the north bank of the Tejo estuary are comparable to values reported in similar species from non-contaminated areas (Cherry 1974; Carvalho 1995; Dahlgaard 1996). This indicates that in the Lisboa area there is no significant enhancement of these radionuclide concentrations above normal background values, as opposed to observations made at the south bank around Barreiro peninsula due to discharge of industrial wastes (Carvalho et al., 2013).

The mean and range of radionuclide concentrations measured in estuarine samples throughout the sampling period, including artificial and main naturally-occurring radionuclides, is shown in Table 1. These radionuclides have different sources. The naturally-occurring radionuclides in mussels and fish tissues did not originate particularly in the urban wastewater discharges. Actually, 210Po originates in the radioactive decay of uranium series radionuclides, which are ubiquitous in nature. In mussels and fish the concentrations of natural 210Po and 210Pb varied over the months, most likely under the effect of the organism physiological conditions. Indeed, it was recently shown in mussels that animal size and accumulation/expenditure of fat are parameters that explain the fluctuation of 210Po concentrations in mussels throughout the seasons of the year (Carvalho et al., 2010). The artificial radionuclides 131I and 99mTc displayed average concentrations...
lower than those of the naturally occurring radionuclides for the entire data set. Notwithstanding, one should keep in mind that concentrations of natural radionuclides are nearly the same in most areas of the estuary while concentrations of artificial radionuclides range from zero in some areas to high values near the sewers discharges (Carvalho et al., 2013).

Absorbed radiation doses due to these natural and artificial radionuclides in mussels and in whole body mullets were computed for the maximum concentrations measured, as reported in Table 1 (Pröhl 2003). It may be noticed that in spite of the absorbed radiation dose imparted to aquatic biota by artificial radionuclides, the main contribution to the overall absorbed dose is still originated by the naturally-occurring radionuclides. Amongst these, $^{210}$Po gave the main contribution (95.5%) to the total absorbed dose in mussels while $^{40}$K gave the main contribution (73%) in fish (Figure 6). Therefore, despite adding radiation dose to the aquatic organisms and to eventual human consumers of aquatic biota, the contribution of radioisotopes from hospital releases did not represent much in comparison with the radiation dose from naturally occurring radionuclides present in the estuarine environment. Notwithstanding, as discharges fluctuated widely throughout the year the contribution of artificial radionuclides to the radiation dose may also vary over time.

### 4. CONCLUSIONS

In the Lisboa area, the Tejo estuary received urban wastewater discharges that carried artificial radioisotopes originated in medical facilities. Besides the risk of radiation exposure of workers of the city sewerage system and wastewater treatment plants, there is also the potential for exposure of members of the public to radioisotopes present in discharges into the estuary. Those radioisotopes were short lived and the temporary storage of contaminated effluents in tanks prior to the discharge should be sufficient to allow for radioactive decay. Nevertheless, there are either facilities without interim storage radioactive effluents or this interim storage was too short and, once released, a rapid transit from the medical facilities to the estuary (a few hours at the maximum) occurred in a systematic manner and allowed the round-the-year detection of $^{131}$I and $^{99m}$Tc, in estuarine water, sediment, mussels, and fish.

The highest radionuclide concentrations were consistently measured near Alcântara and Praça do Comércio where two major sewers drained the urban wastewater discharges into the estuary. Discharges at these sites were of treated and untreated wastewaters, respectively. Results for Alcântara indicate that radionuclides from hospital effluents were not removed in the Alcântara wastewater treatment plant, as could be expected because waste water treatment plants are designed to treat nutrient organic loads and pathogens. Dispersal of sewage discharges by the river flow and tidal currents allowed for transport of radioisotopes up and down along the North bank and accumulation of $^{131}$I by mussels in all sampling stations from Algés (estuary mouth) to Beirolas (mid estuary). Fish accumulated radionuclides, $^{99m}$Tc in particular, but fish mobility renders less obvious than the sessile mussels the exact place of radionuclide uptake. Notwithstanding, $^{99m}$Tc was consistently detected in water, mussels and fish in the stations of Alcântara, Praça.
Table 1. Concentrations of artificial radionuclides and main naturally-occurring radionuclides (210Po, 210Pb and 40K) measured in samples of the Tejo estuary. Mean of all data per radionuclide (n values) and range observed during the 19-month survey period.

<table>
<thead>
<tr>
<th>Radionuclide</th>
<th>Sediments (Bq kg⁻¹ww)</th>
<th>Unfiltered water (Bq L⁻¹)</th>
<th>Mussels (in toto) (Bq kg⁻¹ww)</th>
<th>Fish (in toto) (Bq kg⁻¹ww)</th>
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<tbody>
<tr>
<td>131I</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>88</td>
<td>120</td>
<td>110</td>
<td>47</td>
</tr>
<tr>
<td>mean</td>
<td>0.47</td>
<td>0.56</td>
<td>1.79</td>
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</tr>
<tr>
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<td>10±2</td>
<td>12±2</td>
<td>2.9±0.4</td>
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<td>&lt; 0.31</td>
<td>&lt; 0.27</td>
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<td>99mTc</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>n</td>
<td>88</td>
<td>120</td>
<td>110</td>
<td>47</td>
</tr>
<tr>
<td>mean</td>
<td>0.76</td>
<td>0.62</td>
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<tr>
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<td>&lt; 0.28</td>
<td>&lt; 0.31</td>
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<td>137Cs</td>
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<td>mean</td>
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<td>n</td>
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<td>6</td>
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<table>
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<tr>
<th>Radionuclide</th>
<th>Mussels (soft tissues) (Bq kg⁻¹ww)</th>
<th>Fish (Bq kg⁻¹ww)</th>
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<tr>
<td></td>
<td>Liver</td>
<td>Gonads</td>
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<tr>
<td>210Po</td>
<td>16</td>
<td>14</td>
</tr>
<tr>
<td>mean</td>
<td>-</td>
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<td>-</td>
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<tr>
<td>minimum</td>
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</tbody>
</table>

*x, lower than the detection limit x, which value is indicated in each case.
do Comércio and Chelas. Therefore, suitable measures to better control radioactivity in hospital effluents and in urban wastewater discharge are desirable in order to abate contaminants in the estuarine environment.

Recent improvements in the sewage system of Lisboa and in the Alcântara wastewater treatment plant aiming at the full treatment of urban wastewaters may have modified the pattern of waste discharges and contaminant distribution in the estuary described herein, and improved the quality of wastewater discharges. Nevertheless, this potential improvement was not checked for radioactivity as yet. Furthermore, it is likely that other discharges of radioisotopes from medical facilities exist around the estuary and the assessment of radioactive contamination of the estuary presented here probably is not complete. In addition, the use of radioisotopes in medicine and the number of treatments dispensed increase every year and new hospitals have been recently built in the Lisboa area and in other cities around the Tejo estuary and in its catchment area.

As professional fisheries along with sport fishing and leisure activities take place in the Tejo estuary, the implementation of a regular programme for radioactivity monitoring in the estuary seems much needed to keep under review the radioactivity levels and to assess and update the radiological risk assessment to members of the public. Results from radioactivity monitoring may provide also useful feedback to better control effluent discharges and to improve radioactive waste management.

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REFERENCES


Carolan, J.V.; Hughes, C.E.; Hoffmann, E.L. (2011) – Dose assessment for marine biota and humans from discharges...


