Monitoring of tritium in surface waters in PC, Nuclear facilities of Serbia

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Abstract: The isotope of hydrogen, tritium, is a radioactive isotope. In most cases, it is found bonded to molecules that replace hydrogen. It can be found in nature as a product of interactions of cosmic radiation with air or atmospheric molecules nitrogen. Such tritium is natural. There is also anthropogenic tritium that contributes to increased levels of natural tritium. Artificially produced tritium is produced in the testing of nuclear weapons, in the operation of nuclear power plants, and processing of nuclear fuel. This paper will describe the methods of tritium detection liquid scintillation detector Quantulus 1220 by the direct method in 300 minutes at the Department of Nuclear Physics. Surface water samples with tritium were taken from the location of PC Nuclear facilities of Serbia from the Mlaka stream. The monitoring recorded increased tritium content at two locations, indicating failures of nuclear waste storage.

Keywords: tritium; radioactivity; detector; quantulus1220

1. Introduction

The concentration of tritium on Earth is very low. The atmosphere only has quantities of traces created by interacting its gases with cosmic rays. The atomic nucleus is unstable and decays with a half-life of 12.32 years per emission of one of the electrons in 3He (beta decay) [1–9]. The general properties of tritium are given in the table.
Table 1. Tritium basic properties [1,10]

<table>
<thead>
<tr>
<th>Property</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spin</td>
<td>$\frac{1}{2}^+$</td>
</tr>
<tr>
<td>Half-Life period</td>
<td>12.3 years</td>
</tr>
<tr>
<td>Atomic mass</td>
<td>3.01605u</td>
</tr>
<tr>
<td>Melting point</td>
<td>-252.5 °C</td>
</tr>
<tr>
<td>Boiling point</td>
<td>-248.1 °C</td>
</tr>
</tbody>
</table>

2. Tritium detection technique

Since tritium significantly impacts the environment, analysis and testing have found a significant role in monitoring the condition of nuclear facilities, public protection, dosimetry, and hydrogeology [11–12]. The tested tritium used today for testing and analysis is not entirely of cosmic origin. A large amount of this isotope was created due to man, after 40 of those years, nuclear weapons tests. As we stated, tritium is a weak beta emitter with an energy of $E_{\text{max}} = 18.6\,\text{keV}$ [13–16]. There are several techniques used for tritium detection, including:

- Liquid scintillation counting involves mixing the tritium sample with a scintillation cocktail, emitting light when the tritium decays. The emitted light is then detected and counted using a photomultiplier tube.

- Gas proportional counting: In this technique, the tritium sample is mixed with a proportional gas, and the emitted ionization from tritium decay is measured using a gas-filled detector.

- Ionization chamber: An ionization chamber measures the ionization created by tritium decay in the sample. The chamber consists of a gas-filled container with two electrodes, and the amount of ionization is proportional to the amount of tritium in the sample.

- Electrochemical methods involve using an electrode to detect tritium in a sample. The tritium atoms are ionized at the electrode surface, and the resulting current is measured.
- Scintillation spectrometry: This technique uses a scintillation detector to measure the energy of the emitted light from tritium decay, allowing for the identification and quantification of tritium in a sample [17,18].

In this paper, we used the LSC method so we will base it on it.

2.1. LSC method

First and foremost, in the LSC method, the radioactive sample (for example, tritium water after combustion) should be combined with the scintillation cocktail so that it is in direct contact with a substance that tends to fluoresce when the atoms are excited, thus creating scintillations, i.e., photons. A substance commonly labeled as “fluorine” proved to be appropriate. Fluorine is an appropriate name because of the substance’s ability to scintillate. It lights up when the atoms are excited, so it does not directly refer to the element fluorine [18-24].

The tritium sample is typically dissolved or suspended in a solvent compatible with the liquid scintillator. Mixing with the scintillator: The sample is then mixed with the liquid scintillator, typically a cocktail consisting of a solvent, a scintillant, and other additives to enhance the counting efficiency and reduce background noise. Counting the emitted light: The sample/scintillator mixture is placed in a counting vial and inserted into the LSC instrument. The instrument uses a PMT to detect and count the emitted light from tritium decay in the sample. Data analysis: The counting data is analyzed to determine the concentration of tritium in the sample, typically reported in units of becquerels per liter (Bq·L⁻¹) or disintegrations per minute (DPM) (Fig. 1).

**Figure 1.** Preparation of the sample.
There are several methods of testing tritium using liquid scintillation detectors, and they are:

- **ASTM** - a standard method for testing tritium
- Electrolytic enrichment
- Direct method

We used the direct method in this work, so more will be said.

### 2.1.1. Direct method

The direct LSC method is a relatively simple and fast technique for tritium detection. It is often used in environmental monitoring, nuclear power plants, and other applications where tritium may exist.

The direct LSC method offers high sensitivity and can detect tritium at very low levels, typically down to tens of tritium atoms per liter of sample. However, care must be taken to minimize background radiation and to ensure that the liquid scintillator is properly calibrated and free of impurities that could interfere with the counting efficiency.

This method is called direct because it does not require electrolytic enrichment. It is direct determines the tritium values. The method belongs to the rapid tests. It is used as the method for examining limit values of radionuclides in drinking water. It is also applied when checking water radiation at nuclear reactors, which serves for moderation or cooling [20]. The method uses a scintillation detector that can detect small energy values in which the emission of beta particles that accompanies the decay of tritium falls.

The following equipment requires:

- Liquid scintillation detector (Quantulus1220);
- Bottles of 20 ml (plastic);
- Apparatus for filtration;
- Apparatus for distillation.
3. Results

Before measuring, we define locations and reference values (Fig. 2). Locations:

- Mlaka 1 - Place without any impact of nuclear facilities on the environment.
- Mlaka 2 - Sampling site at the recipient.
- Mlaka 3 - Sampling site downstream from the recipient.

![Figure 2. Selected locations.](image)

For the detection of tritium in precipitation, sampling with location Zeleno brdo, which is located 7 km from the Nuclear facility, while for the detection of tritium in surface water was taken from Mlaka 1.

3.1. Detector optimization

The detection efficiency is determined by equation 1:

\[
\varepsilon_t = \frac{s}{8A_r}
\]

where is \( A_r \) [Bq/l]-activity concentration of \(^3\)H solution (equation 2).
\[ A = \frac{r-(b+r_q)}{60 \varepsilon t V} \]  

Detection efficiency for Ultima Gold LLT and Optiphase, Ultima Gold LLT is shown in Fig. 3.

![Detection efficiency for Ultima Gold LLT and Optiphase](image)

**Figure 3.** Detection efficiency for Ultima Gold LLT and Optiphase, Ultima Gold LLT used in measurements.

The minimum detectable tritium MDA activity achieved during the measurement of 300 min is calculated using equation 3:

\[ MDA = \frac{2.71+4.65 \sqrt{(b+r_q)T}}{60 \varepsilon t V} \]  

A display of minimum detectable activity depending on measurement times is shown in Fig. 4.

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**Figure 4.** Display of minimum detectable activity depending on measurement times for Ultima Gold LLT.

Fig. 5 presents the dependency of specific activity in one year.

**Figure 5.** Presentation of specific activity in one year.

Mean values of tritium concentration for Mlaka 2 and Mlaka 3 in the period from December 2017 to June 2019 are calculated using equation 4:
\[ A_{sr}(3H) = \frac{A_1 + A_2 + A_3 \ldots + A_{19}}{19} \]  

Values obtained for Mlaka 2: \( A_{sr}(3H)_{Mlaka2} = 7.84 \text{ Bq}^{-1} \)

Values obtained for Mlaka 3: \( A_{sr}(3H)_{Mlaka3} = 6.81 \text{ Bq}^{-1} \)

Statistical presentation of maximum, minimum, and mean values of tritium activity in 19 months is presented in Fig. 6.

\[ \text{Figure 6. Statistical presentation of maximum, minimum, and mean values of tritium activity in 19 months.} \]

4. Conclusion

The isotope of hydrogen, tritium, is a radioactive isotope of hydrogen. In most cases, it is found bound to molecules that replace hydrogen. It can be found in nature as a product of the interaction of cosmic radiation with air molecules, i.e., atmospheric nitrogen. Such tritium is natural. There is also anthropogenic tritium that contributes to increased levels of natural tritium. Artificially produced tritium is produced in nuclear weapons tests, in the operation of nuclear power plants, and in the processing of nuclear fuel, as stated in the paper.
It was determined that the most effective method for tritium detection is the direct method due to its advantages over the other two mentioned methods.

Liquid scintillation detector Quantulus 1220 is an "ultra-low-level" counter and, as such is suitable for measuring extremely low concentrations of alpha and beta emitter activity, and as such provides a wide range of possibilities for work and measurement of not only tritium but also other radionuclides. The calibration of the Quantulus 1220 detector system using the direct method for 300 min using the LLT cocktail is 2.2 Bq\textper litre, which meets the detection limits. A value of 31.2% was taken for the detection efficiency.

Through monitoring, it was established that increased tritium values are found in the recipient’s surroundings.

During the processing of the results, it was shown that the maximum concentrations of tritium in the tepid stream are as high as 19 Bq\textper litre, and the minimum concentrations are almost at the limit of the minimum detectable activity of 2.3 Bq\textper litre. Also, these values do not exceed the parametric values in drinking water defined by the "Official Gazette of the RS" of 100 Bq\textper litre. All measured values do not exceed the legally prescribed values and are far below the critical values.

References

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