Preparation of Tritium Standard Gas by the Use of Tritiated Methane

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Tritium standard gas for calibrating radioactivity detectors was prepared by the use of tritiated methane synthesized by the reaction of tritiated water with aluminum carbide at ca. 470°C. A long-path proportional counter was used to standardize the radioactivity of the prepared gas. Tritiated gas of less than 10^4 Bq could be measured by this counter with the overall error of 3.5%(3σ). Finally, two types of standard were prepared in a metal cylinder and a glass ampoule. The latter type could be used as a handy calibration source for detectors.

Key Words: tritium, radioactive standard, tritiated methane

1. Introduction

A great number of studies on tritium have been made from various standpoints, especially in connection with the development of fusion reactors. For the measurement of the radioactivity of gaseous tritium, proportional counters and ionization chambers are often employed. Tritium standard gases commercially available from several suppliers are used for the calibration of those apparatuses. These standards, however, are not always satisfactory in the accuracy of radioactivity (e.g. 3 to 10% uncertainty) and are somewhat expensive in spite of the short period to be guaranteed for the use (e.g. 4 months). For periodic calibrations of detectors of tritium, Yoshida, et al. prepared secondary standard gases of tritium from primary standards purchased from a foreign supplier.

The authors have attempted to prepare a primary tritium standard gas with good accuracy and of convenience to calibrate detectors. Tritiated methane was employed as a standard gas because permeation through and adsorption on a container wall are less effective for methane than for tritium gas (HT, T2). Tritiated methane was synthesized by the reaction of tritiated water with aluminum carbide. The optimum reaction temperature for the production of radiochemically pure tritiated methane was examined. To standardize the radioactivity of tritiated gas prepared by this work, a long-path proportional counter developed by the authors was employed.

2. Experimental

2.1 Synthesis of tritiated methane

2.1.1 Materials

Aluminum carbide (Al4C3, ≥98% in purity, 325 mesh) purchased from Mitsuwa Chemicals Co., Ltd. was used without further purification. Tritiated water (ca. 0.4 MBq ml−1) was supplied from a stock solution which was generated as a waste solution of tritium production experiments in the authors’ laboratory. Tritiated water (ca. 0.4 MBq ml−1) was supplied from a stock solution which was generated as a waste solution of tritium production experiments in the authors’ laboratory. To synthesize the tritiated methane of a relatively high specific radioactivity, tritiated water with nominal specific activity of 195 GBq ml−1 was obtained from Amersham International plc, England, and was used after dilution procedures to get proper concentrations. Helium (99.9999%) and N2 (99.999%) gases, which came from Nippon Sanso Co., Ltd. and Hitachi Sanso Co., Ltd. respectively, were used as carriers without further purification.

2.1.2 Apparatus and procedures

The apparatus used for the synthesis of tritiated methane is illustrated in Fig. 1. Tritiated water was injected with a micro syringe (A) through a silicone packing into the reaction column (15 mm ∅ × 200 mm; B) of quartz glass
Fig. 1 Schematic diagram of the apparatus used for the synthesis and distribution of tritiated methane.

packed with the mixture of aluminum carbide (Al₄C₃, ca. 2g) and quartz wool. The reaction products were led with He carrier gas through a cold trap (D) cooled at -72°C, in which unreacted tritiated water was collected, to a gas-chromatograph in order to measure the amount of synthesized methane. The gas-chromatograph (Shimazu GC-6AM) was equipped with a thermal conductivity detector and a column (4 mmφ x 2 m) packed with activated carbon (30/60 mesh). Temperature of the column was kept at 100°C. The gaseous products were finally collected in a silica gel-containing cold trap (E) chilled at -196°C.

After a desired amount of tritiated methane had been collected in the trap (E), a stopcock (F) was closed and a three-way cock (G) was turned to connect the trap (E) to a rotary pump. The He carrier gas was removed from the trap (E) by evacuation for more than 10 min. Turning the three-way cock (H), the trap (E) was then warmed up to an ambient temperature. The retaining methane in the trap was evaporated by this treatment and transferred with the aid of a Toepler pump (K) to the ampoule system (L) which was evacuated in advance. The ampoule system was equipped with stopcocks for each glass ampoule (2 to 10 ml in volume). These stopcocks were closed before the ampoules filled with tritiated methane were fuse-sealed off. By this operation, scatterings of the amount of the sealed gas per unit volume of the ampoules were reduced.

In general, the reaction of aluminum carbide with water is expressed in the equation

$$\text{Al}_4\text{C}_3 + 12\text{H}_2\text{O} \rightarrow 4\text{Al(OH)}_3 + 3\text{CH}_4,$$  \hspace{1em} (1)

where half of the hydrogen atoms in the water (tritium atoms when tritiated water was used) is transformed into methane. And it takes a few hours to complete this reaction at relatively low temperatures (100 - 200°C) because of its moderate reactivity\(^5,6\). On the other hand, aluminum hydroxide produces water by thermal decomposition at higher temperatures (>300°C);

$$2\text{Al(OH)}_3 \rightarrow \text{Al}_2\text{O}_3 + 3\text{H}_2\text{O}.$$  \hspace{1em} (2)

Therefore, the reaction

$$\text{Al}_4\text{C}_3 + 6\text{H}_2\text{O} \rightarrow 2\text{Al}_2\text{O}_3 + 3\text{CH}_4,$$  \hspace{1em} (3)

which has been derived from the equations (1) and (2), can be expected at higher temperatures (>300°C). In this case, hydrogen (tritium) will be transformed into methane efficiently and the reaction time can be reduced at the same time. To examine these advantages, temperature effect of this reaction was investigated at first.

Water (H₂O, 3 μl) was injected into the reaction column at various temperatures, while the flow rate of He carrier gas was adjusted at 15 ml min⁻¹. Effect of the flow rate on the yield of methane, on the other hand, had been surveyed from 0 ml min⁻¹ (water being kept in the reaction column for a few minutes) to 30 ml min⁻¹ in advance. This effect, however, proved
to be of less importance compared with the temperature effect. Accordingly, He carrier gas was controlled to flow at 15 ml min\(^{-1}\) afterwards.

2.2 Measurement of radioactivity
Radioactivity of the tritiated gas was measured with a long-path proportional counter\(^5\). Figure 2 shows a setup of the counting system. The long-path proportional counter (A) made from stainless steel tube of 1 000 mm long and 21.3 mm i.d. was coupled with a gas handling system. As an anode of the counter, a 50 µm tungsten wire was stretched in the center of the tube. The operation voltage was 3 200 V. The aliquot of sample gas in the cylinder (B) was introduced into the sampler (D) whose volume was precisely known. Temperature and pressure of the gas were measured simultaneously. The sample gas was then transferred with counting gas (pure methane) into the counter (A). Counting was carried out while the sample gas was moving through the counter tube together with the counting gas at the flow rate of 30 ml min\(^{-1}\).

For the measurement of radioactivity of tritium in a sealed ampoule, the ampoule was broken in the ampoule breaker (C) in the atmosphere of methane counting gas and the released gas was introduced into the counter (A). Counting was performed in the same way as described above.

3. Results and Discussion

3.1 Synthesis of tritiated methane
3.1.1 Effect of reaction temperature
The temperature effect on the yield of methane was measured by a gas chromatograph, shown in Fig. 3. The amount of methane synthesized was very small at temperatures below 400°C. The maximum yield of methane was obtained at 520 to 560°C and the yield of methane decreased gradually beyond ca. 560°C.

Radio-gas chromatographic assay of tritiated methane synthesized at 520°C revealed the presence of HT in the product. Temperature dependency on the production of hydrogen was then investigated. As seen in Fig. 4, a rapid increase in hydrogen generation was observed above 500°C. The decrease in the yield of methane at temperatures above ca. 560°C could be explained by the generation of hydrogen as a by-product. The presence of HT in a tritium standard gas seemed to reduce the accuracy of the radioactivity value because of the permea-
A : Standard gas cylinder  B : Ampoule breaker
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for the preparation of cylinder-type standard
gas by compressing tritiated methane
with N₂ carrier gas.

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Sample gas started to flow toward the counter at (a), reached the inlet of the tube at (b), entered in the tube completely at (c), reached the outlet at (d), and went out of the tube entirely at (e).

Fig. 7 Response curve of tritium measurement obtained by using a long-path proportional counter.

volume during this time. The constancy of count rate indicates that the end effect of the counter was avoided, and this was confirmed by using a position sensitive detecting system\(^7\).\(^8\).
Though a small correction for wall effect should be necessary, the radioactivity of the sample gas could be determined without end effect. Tritiated gas of 1 to 10 ml of less than 10\(^4\) Bq could be measured by using this setup. The statistic error was below 1.0\(\%\) (3\(\sigma\)) when tritiated gas of 30 to 10\(^4\) Bq ml\(^{-1}\) in radioactive concentration was measured. The systematic error consisting of wall effect, sampling, threshold, and dead time was \(\pm 2.5\%\). The overall error was then estimated to be \(\pm 3.5\%\). This method of measurement will be reported in detail elsewhere\(^9\).

4. Conclusion

Radiochemically pure tritiated methane was synthesized by the reaction of tritiated water with aluminum carbide at ca. 470\(^\circ\)C. The prepared gas was filled in two types of containers; a metal cylinder and a glass ampoule, and standardized by the use of a long-path proportional counter with an overall error of 3.5\% (3\(\sigma\)). For the case of ampoule-sealed samples, a small error may arise in the course of sealing procedure, but this type of sample is easy to handle in the calibration work of radioactivity detectors.

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References

9) to be published
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トリチウム標識メタンを用いたトリチウム標準ガスの作製

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トリチウム水を炭化アルミニアムと約 470℃ で反応させることにより合成したトリチウム標識メタンを用いて、放射能検出器校正用トリチウム標準ガスを作製した。得られたガスの放射能値の絶対測定は長径管比例計数管を用いて行った。この計数管により 10^6 Bq 以下のトリチウム化ガスを ±3.5% (3σ) 以内の誤差で値付けることができる。最終的に、ボンベ型およびアンプル型の 2 種類の標準ガスを作製した。とくに後者は校正用ガスとして簡便に使用することができる。