

DETERMINATION OF POLONIUM-210 IN SEAFOOD WITH LARGE AREA GRID IONIZATION CHAMBERS ALPHA SPECTROMETRY

Yanqin Ji¹, Liangliang Yin², Xianzhang Shao³

China CDC Key Laboratory of Radiological Protection and Nuclear Emergency, National Institute for Radiological Protection, China CDC, 2 Xinkang Street, Deshengmenwai, 100088 Beijing, China
¹jiyanqin@nirp.cn, ²yinliangliang2010@163.com, ³shaoxzh@nirp.cn

A rapid and reliable method for determination of ²¹⁰Po using large-area grid ionization chamber α spectrometry was established. Samples were digested using microwave digestion system. After preparation of sample source with ultrasonic disperse and vacuum evaporation, the radioactivity concentration of ²¹⁰Po in clam was detected by large-area grid ionization chamber α spectrometry (ϕ 25cm). ²⁰⁹Po tracer was used to obtain the recovery.

I. INTRODUCTION

Polonium-210 (²¹⁰Po) occurs in natural environment at trace levels as part of the ²³⁸U decay chain. As a pure α emitter, ²¹⁰Po is one of the most radiotoxic nuclides. With wide dispersion in the environment, ²¹⁰Po enters human body via food chains, thus being a serious threat to human health. ²¹⁰Po was known as one of the most important sources of internal dose received by humans from foods.¹ The maximum permissible human body-burden of ingested ²¹⁰Po was 1.1×10^3 Bq.² Therefore, the capability to rapidly analyze large numbers of samples in variety of matrices is necessary for risk assessment and post incident decision making. The purpose of this study is to develop a rapid and reliable method for determination of ²¹⁰Po using large-area grid ionization chamber α spectrometry.

II. METHODS

II.A. Sample preparation

In the experiment, the soft tissues of clam sample were taken out, dried at 105 °C, and grinded into powder. 0.1-0.5 g of powder was digested with HNO₃-H₂O₂ in microwave system, ²⁰⁹Po standard solution was added as the tracer. Followed eliminating acid, adding ultrapure water to disperse sample solution under ultrasonic. Then the sample solution was transferred directly into dish (ϕ 25cm), and the sample dish were then put in the vacuum oven, evaporated to dryness. The counting source prepared in this way can be put in large-area grid ionization chamber directly.

II.B. Spectrometry

The sample source were counted using large-area grid ionization chamber. The chamber filled with commercially available mixed gas of 90% Ar and 10% CH₄ to about 48.26 kPa³. Standard source (²³⁷Np-²³⁹Pu-²⁴¹Am mixed source, ϕ 25cm) was used for energy calibration and efficiency determination.

III. RESULTS AND DISCUSSION

The ϕ 25cm blank dish was counted for 24 h, while mixed standard source for 30 min, to obtain detection efficiency 33% of the large-area grid ionization chamber α spectrometry. The minimum detectable activity (MDA) was 10×10^4 Bq. ²⁰⁹Po standard solution was spiked in clam powder and the present method was used to prepare source. The recovery was 98.5%.

The dry clam samples were analyzed using two methods. The result of 0.057 Bq/g activity in clam samples is good agreement with the commonly used method that the spontaneous deposition source (ϕ 2cm) measured using multichannel α spectrometer.

The method developed for measurement of seafood could achieve higher recovery (above 98%) and lower MDA, using small quantity of sample (0.1 - 0.5 g of dry sample).

IV CONCLUSIONS

Compared with the traditional method, the developed method can avoid separation process, using less quantity of sample and simplify the measurement process. The method is useful for measurement of other biological or environmental sample.

ACKNOWLEDGMENTS

The authors acknowledge financial support by the Chinese Ministry of Science and Technology (Grant No. 2014FY211000 and 2013BAK03B00).

REFERENCES

1. P. Planinšek, L. Benedik, et al. Comparison of various dissolution techniques for determination of ²¹⁰Po in biological sample. *Appl Radiat Isot*, **81**(11): 53-56 (2013).
2. H. Lee, J.Wang, et al. Annual dose of Taiwanese from the ingestion of ²¹⁰Po in oysters. *Appl Radiat Isot*, **73**: 9-11 (2013).
3. H.H. ötzl, R. Winkler, et al. Experiences with large-area Frisch grid chambers in low-level alpha spectrometry. *Nucl Instrum Methods Phys Res*, **223**(84): 290-294 (1984).