Note

Effects of Several Variable Factors on the Isotope Ratio by HRGC-MS

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In the isotope ratio (Ir) analysis using GC-MS, several variable factors in sampling incidental to any food analysis were investigated for yuzu fruit. The Ir values of ten monoterpenyl hydrocarbons in yuzu essential oils from each of six fruiting positions of three trees were measured. The sign test following t-test of all the Ir values demonstrated that there was no significant difference between both sampling years of 2001 and 2002. There was also no significant variation in the Ir values among the three trees and six fruiting positions in the individual two years.

Key words: isotope ratio analysis; sampling variations; HRGC/MS; monoterpenyl hydrocarbon

A determination of the isotope value of constituents is gaining increasing importance, especially in view of the increased demand in the fragrance and food industries for the authenticity control and origin determination of the production of essential oils and food.1-3) Recently, many sensational articles have been reported on dishonest designations of raw foods, since the JAS (Japanese Agricultural Standards) were revised in the increased demand in the fragrance and food industries for the authenticity control and origin determination of the production of essential oils and food.1-3) Recently, the JAS (Japanese Agricultural Standards) were revised on July 2000. The authors4) have developed a specific and convenient method for characterization of yuzu (Citrus Junos Tanaka) fruits produced in different districts by the isotope ratio of monoterpenyl hydrocarbons with HRGC-MS. This method will be applicable to several cases such as differentiation of the quality, evaluation of genuineness of vegetable products from various producing districts, and classification of the species. It is noticed that this analytical method will be useful for chemical evaluation of agricultural products, in addition to DNA analysis.5,6) The authors7,8) have also reported successful results on the discrimination of commercial citrus essential oils of different origins, and the accuracy of the Ir method using 13C-labeled compounds. There remain, however, some questions about samplings from the fundamental scope. In order to normalize this method, the variations or individual differences which each material substantially owns should be further investigated. This paper concerns the effects of variable factors involved in fruit samples regarding the isotope ratio analysis.

Three yuzu trees (13- to 15-years age) were selected at random among a number of yuzu trees under the usual cultivation in the field of Kochi Fruit Tree Experimental Station, Kochi. The experiments were done from the following three viewpoints: (1) annual variations between 2001 and 2002; (2) variations among trees; (3) variations between fruiting positions on the trees. All the samplings were done from these three trees in November 2001 and 2002. The six fruiting positions on each tree were as follows: top, inside, and the four sides of East, West, South, and North around equatorial canopy of the tree. The cold-pressed peel oil (CPO) of about 2 kg of each sample was prepared according to the usual method.9) The yield of CPO was 0.10–0.16 g/100 g of fresh fruit, and 0.34–0.59 g/100 g of flavedo. Each CPO sample was kept at −25°C until needed for analysis. Gas chromatography-mass spectrometry (GC-MS) was done with a Shimadzu GC-17A linked to a Shimadzu QP-5000. The GC column was a DB-Wax fused silica capillary (60 m × 0.25 mm i.d., 0.25 μm film thickness; J & W Scientific, Folsom, CA, U.S.A.) The analytical conditions of GC-MS and column temperature were the same as those of the previous paper.10) The isotope ratio (Ir) of the selected monoterpenyl hydrocarbons in CPO, m/z 137/136, was measured by selected-ion monitoring (SIM). Those compounds analyzed were α-pinene, β-pinene, sabinene, myrcene, α-phellandrene, α-terpinene, limonene, β-phellandrene, γ-terpinene, and terpinolene. All the measurements were triplicated. The statistical treatments, t-test and sign test, were done as well as described before.5,10)

A sign test was applied to the following evaluation. The Ir values of ten monoterpenyl hydrocarbons in the CPOs from each of the six fruiting positions of three trees were measured in triplicate in an individual year. All the data obtained on each monoterpenyl hydrocarbon were first normalized. The resulting data were analyzed by a sign test.4) The

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The results are shown in Table 1. In the case of ten parameters (n = 10), i.e., ten monoterpene hydrocarbons, the significant level (P < 0.05) is below 1.0107. It was demonstrated that there was no significant difference between the two sampling years of 2001 and 2002. An annual variation factor, in other words, will not significantly affect the analysis based on the Ir ratio, even if fruits are sampled from any position on a tree.

All the Ir values of ten monoterpene hydrocarbons from all the fruiting positions on each tree were used for the t-test among the three trees. The result is shown in Table 2. There was no significant difference in the Ir values between the trees (P < 0.05). Thus, it suggests that any tree in a given area is allowed to be sampled without significant variation.

The Ir values of ten monoterpene hydrocarbons from each fruiting position on the three trees in 2002 were analyzed by the least significant difference test, as shown in Table 3. No significant difference (P < 0.05) among all the fruiting positions of each compound was seen. The result for 2001 was the same as that for 2002.

In conclusion, it is suggested that the data obtained from this Ir ratio analysis, which is used for discriminating producing districts or genuineness of crops, are independent upon common variations incidental to samplings as long as the samples are collected in the same area. Many common components of foods usually fluctuate among individual samples. It is expected that this analysis will be used widely for judgment of the characteristics of agricultural products.

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