Comparison of Proximate and Mineral Composition Between Asparagus officinalis and Momordica dioica: Iranian and Indian Vegetables

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Abstract: The proximate composition and mineral constituents of Asparagus officinalis stem and Momordica dioica fruit were evaluated in order to standard methods. The stem contained a ashes: 10.70%, crude protein: 32.69%, crude lipid: 3.44%, crude fiber: 18.50% and carbohydrates: 34.67%. Stem also have high energy value (384.27 kcal/100 g dry weight). Mineral ranges (mg/100 g dry weight, DW) were: K (10.94), Na (1.84), Ca (0.67), Fe (0.19) and Zn (2.60). The fruits contained a ashes: 9.1%, crude protein: 5.44%, crude lipid: 3.25%, crude fiber: 22.99% and carbohydrates: 59.31%. The fruits also have high energy value (288.25 kcal/100 g dry weight). Mineral ranges (mg/100 g dry weight, DW) were: K (4.63), Na (1.62), Ca (7.37), Fe (5.04) and Zn (3.83). Comparing proximate and minerals contents of the stem and the fruit, the results indicated that Asparagus officinalis stem could be a good supplement for some nutrients such as protein, lipid, potassium and Zinc, fibre and carbohydrates while Momordica dioica fruit was good source of lipid, crude fiber, carbohydrates, Fe and Zn.

Key words: Asparagus officinalis stem · Momordica dioica · Micronutrients · Proximate and Mineral composition

INTRODUCTION

In developing nations, numerous types of edible wild plants are exploited as sources of food hence provide an adequate level of nutrition to the inhabitants. Recent studies on agro pastoral societies in Africa indicate that these, plant resources play a significant role in nutrition; food security and income generation [1].

Furthermore, FAO report, at least one billion people are thought to use wild foods in their diet [2]. In Ghana alone, the leaves of over 300 species of wild plants and fruits are consumed. In Swaziland, wild plants provide a greater share of the diet than domesticated cultivars. In India, Malaysia and Thailand, about 150 wild plants species have been identified as sources of emergency food [3]. Similarly, in South Africa about 1400 edible plant species are used, in Sahel region of Africa, over 200 wild foods were identified to be used by the rural communities [4]. In most of these reports, it was emphasized that nutritionally, these unconventional plants foods could be comparable to or even sometimes superior to the introduced cultivars [5]. It is, therefore, worthwhile to note that the incorporation of edible wild and semi-cultivated plant resources could be beneficial to nutritionally marginal populations or to certain vulnerable groups within populations, especially in developing countries where poverty and climatic changes are causing havoc to the rural populace. In this context, analyses were carried out to evaluate the nutritional content of Asparagus officinalis stem and Momordica dioica fruit with hope that it would be incorporated into the food basket of the country [3,6-8]. Aim of analysis of nutrients in the plant foods is preliminary assessment of nutritional value of the plant-based diet.

Plant Material: Asparagus officinalis stem and Momordica dioica. Fruits used as experimental material were collected from farm and Agricultural lands(garden) in around Behbahan, South Iran, in October 2007. The collected plant material was placed in a polyethylene bag to prevent loss of moisture during transportation to the laboratory.

Preparation of the Plant Materials for Chemical Analyses: Asparagus officinalis stem and Momordica dioica. Fruits were washed with distilled water and dried

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at room temperature to remove residual moisture, then placed in paper envelope and oven-dried at 55°C for 24 hours [9]. The dried stem were ground into powder using pestle and mortar and sieved through 20-mesh sieve. The root tubers powder was used for the nutrients analyzes.

**Plant Foods Chemical Analysis:** The methods recommended by the Association of Official Analytical Chemists were used to determine ash (#942.05), crude lipid (#920.39), crude fibre (#962.09) and nitrogen content (#841.13) [9].

**Determination of Crude Lipid and Crude Fibre Content:** Two grams of dried root tubers or fruits were weighed in a porous thimble of a Soxhlet apparatus, with its mouthed cotton wool plugged. The thimble was placed in an extraction chamber which was suspended above a pre-weighed receiving flask containing petroleum ether (b.p. 40-60°C). The flask was heated on a heating mantle for eight hours to extract the crude lipid. After the extraction, the thimble was removed from the Soxhlet apparatus and the solvent distilled off. The flask containing the crude lipid was heated in the oven at 100°C for 30 minutes to evaporate the solvent, then cooled in a dessicator and reweighed. The difference in weight was expressed as percentage crude lipid content.

Crude fibre was estimated by acid-base digestion with 1.25% H₂SO₄ (prepared by diluting 7.2 ml of 94% conc. acid of specific gravity 1.835g ml⁻¹ per 1000 ml distilled water) and 1.25% NaOH (12.5 g per 1000 ml distilled water) solutions. The residue after crude lipid extraction was put into a 600 ml beaker and 200 ml of boiling 1.25% H₂SO₄ added. The contents were boiled for 30 minutes, cooled, filtered through a filter paper and the residue washed three times with 50 ml aliquots of boiling water. The washed residue was returned to the original beaker and further digested by boiling in 200 ml of 1.25% NaOH for 30 minutes. The digest was filtered to obtain the residue. This was washed three times with 50 ml aliquots of boiling water and finally with 25 ml ethanol.

The washed residue was dried in an oven at 130°C to constant weight and cooled in a dessicator. The residue was scraped into a pre-weighed porcelain crucible, weighed, ashed at 550°C for two hours, cooled in a dessicator and reweighed. Crude fibre content was expressed as percentage loss in weight on ignition, [9,10].

**Determination of Nitrogen Content and Estimation of Crude Protein:** Macro-Kjeldahl method was used to determine the nitrogen content of the stem 2g of dried plants were digested in a 100 ml Kjeldahl digestion flask by boiling with 10 ml of concentratin tetraoxosulphate (VI) acid and a Kjeldahl digestion tablet (a catalyst) until the mixture was clear. The digest was filtered into a 100 ml volumetric flask and the solution made up to 100 ml with distilled water. Ammonia in the digest was steam distilled from 10 ml of the digest to which had been added 20 ml of 45% sodium hydroxide solution. The ammonia liberated was collected in 50 ml of 20% boric acid solution containing a mixed indicator. Ammonia was estimated by titrating with standard 0.01 mol L⁻¹ HCl solution. Blank determination was carried out in a similar manner. Crude protein was estimated by multiplying the value obtained for percentage nitrogen content by a factor of 6.25 [9].

**Estimation of Carbohydrates and Energy Values:** Available carbohydrate was estimated by difference, by subtracting the total sum of percent crude protein, crude lipid, crude fibre and ash from 100% DW of the plant the plant calorific value (in kJ) was estimated by multiplying the percentages of crude protein, crude lipid and carbohydrate by the factors 16.7, 37.7 and 16.7 respectively [9].

**Mineral Analysis:** The mineral elements Na, K, Ca, Fe and Zn were determined on 0.3g plant powder by the methods of Funtua using Energy Dispersive X-ray Fluorescence (EDXRf) transmission emission spectrometer carrying an annual 25 mCi 109Cd isotopic excitation source that emits Ag-K X-ray (22.1 keV) and a Mo X-ray tube (50KV, 5mA) with thick foil of pure Mo used as target material for absorption correction. The system had a Canberra Si (Li) detector with a resolution of 170eV at 5.9keV line and was coupled to a computer controlled ADCCard (Trump 8K).

Measurements were carried out in duplicate. Na was analyzed after wet digestion of one gramme of the stem powder with nitric/perchloric/sulphuric acid (9:2:1 v/v/v) mixture. Sodium was analyzed with a Corning 400 flame photometer [9].

**RESULTS AND DISCUSSION**

**Proximate Analysis:** The results of proximate composition of Asparagus officinalis stem and Momordica dioica fruits are shown in Tables 1 and 2. The ash content, which is an index of mineral contents, for Asparagus officinalis stem was more than to the values reported for other edible leaves such as Momordica balsamina leaves where as for Momordica dioica fruits the value of 6.7% DW was less than the reported values [1,3,11]. It is apparent that Asparagus officinalis stem are
a good source of Potassium. The crude protein content of both edible plants was more than what is reported for some lesser known wild leafy vegetables such as Morinda balsamina (11.29 ± 0.07%), Moringa oleifera (20.72%), Lessianthera africana leaves (13.10-14.90%) and Leptadenia hastate (19.10%) [12,13]. Plant food that provide more than 12% of their calorific value from protein are a good source of protein. In that context, Asparagus officinalis stem and Morinda dioica fruits are good sources of protein. The crude lipid contents of both plant foods were less than the range (8.3-27.0% DW) reported for some vegetables consumed in Nigeria and Republic of Niger [10,14].

The estimated carbohydrate contents in Asparagus officinalis stem and in Morinda dioica fruits were stand to be higher than that for Senna obtusifolia leaves (20%) and Amaranthus inculvatus leaves (23.7%). On the other hand, Asparagus officinalis stem and M. dioica fruits contain comparable amount of carbohydrate for Morinda balsamina (39.05 ± 2.01%). The crude fibre content in both plant foods were more than the reported values (8.50-20.90%) for some Nigeria vegetables, [15]. One discussed drawback to the use of vegetables in human nutrition is their high fibre content, which may cause intestinal irritation and a decrease of nutrient bioavailability. The fibre RDA values for children, adults, pregnant and breast-feeding mothers are 19-25%, 21-38%, 28% and 29% respectively.

**Mineral Content:** Table 3 and 4 show the results of the mineral concentrations of Asparagus officinalis stem and Morinda dioica fruits. Nutritional significant of elements is compared with the standard recommended dietary allowance. When compared with standard values as showed in Table 3 and 4, Asparagus officinalis stem and M. dioica fruits less than adequate level of K, Fe, Zn, Ca and Na, but the plant stem could be good sources of K.

**Concluding Remarks:** The results of the nutritional analysis shown that Asparagus officinalis stem could be a good supplement for some nutrients such as protein, lipid, potassium and Zinc, fibre and carbohydrates while Morinda dioica fruit was good source of lipid, crude fiber, carbohydrates, Fe and Zinc.
The results suggests that the plant fruits if consumed in sufficient amount could contribute greatly towards meeting human nutritional requirement for normal growth and adequate protection against diseases arising from malnutrition. From the result, Asparagus officinalis stem and Momordica dioica fruits are recommend for continues used for nutritional purporses, considering to the amount and diversity of nutrients it contains. Chemical analysis alone however, should not be the exclusive criteria for judging the nutritional significance of a plant parts. Thus, it becomes necessary to consider order aspects such as presence antinutritional/toxicological factors and biological evaluation of nutrient content, [16].

**Abbreviations Used:** AOAC, Association of Official Analytical Chemists; FAO, Food and Agricultural Organization; RDA, recommended dietary allowances; DW, dried weight.

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