Iodine uptake and distribution in horticultural and fruit tree species

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Abstract

Iodine is an essential microelement for mammals (Welch and Graham, 1999) with a unique role in organisms. In fact, it is an indispensable component of the thyroid hormones, thyroxine (T₄) and triiodothyronine (T₃), which are involved in the regulation of various enzymes and metabolic processes. Inadequate dietary iodine intake can cause insufficient thyroxin production. This results in iodine deficiency disorder (IDD). The clinical manifestations of IDD include hypothyroidism, goiter, mental retardation, reproductive impairment, deaf-mutism, and lower child survival rates (Hetzel, 1983; Stanbury et al., 1996). The recommended daily nutrient intake of iodine is 50 mg for infants (aged 0-12 months), 90 mg for children (aged 2-6 years), 120 mg for schoolchildren (aged 7-12 years), 150 mg for adolescents (aged over 12 years) and adults, and 200 mg for pregnant and lactating women (WHO/UNICEF/ICCIDD, 1996). These amounts are thought to allow normal hormone production without stressing the thyroid iodide trapping mechanism. Excess iodine intake is more difficult to define. Many people habitually ingest huge amounts of iodine (approx. 10-200 mg day⁻¹) with no evidence of negative effects (FAO/WHO, 2004).

In the human diet, seaweed, fish and shellfish, all of which absorb iodine from the water, are the best sources of iodine with amounts ranging from 163 to 3180 μg kg⁻¹ fresh weight (FW). The iodine content of terrestrial foods is generally much lower, and concentrations vary widely according to geographical location: from 10 to a maximum 200 μg kg⁻¹ FW (FAO/WHO, 2004).

Iodine is an essential microelement for humans and iodine deficiency disorder (IDD) is one of the most widespread nutrient-deficiency diseases in the world. Iodine biofortification of plants provides an attractive opportunity to increase iodine intake in humans and to prevent and control IDD. This study was conducted to investigate the iodine uptake and accumulation in the edible portion of two fruit trees (plum and nectarine) and two horticultural crops (tomato and potato). We tested two types of iodine treatment (soil and foliar spray application) and, for fresh market tomato, two production systems (open field and greenhouse hydroponic culture). We investigated the distribution of iodine in potato stem and leaves, and in plum tree fruit, leaves and branches. Iodine content of potato tubers after postharvest storage and processing (peeling) were also determined. Differences in iodine accumulation were observed among the four crops, between applications, and between production systems. In the open field, the maximum iodine content ranged from 9.5 and 14.3 μg 100 g⁻¹ for plum and nectarine fruit, to 89.4 and 144.0 μg 100 g⁻¹ for potato tuber and tomato fruit, respectively. These results showed that nectarine and plum trees accumulated significantly smaller amounts of iodine in their edible tissues compared with potato and tomato. Results also showed hydroponic culture to be the most efficient system for iodine uptake in tomato, since its fresh fruit accumulated up to 2423 μg 100 g⁻¹ of iodine. In all species investigated, iodine was mainly stored in the leaves. Only a small portion of iodine was transported to plum tree branches and fruit, and to potato stems and tubers. No differences in iodine content were observed after peeling. A significant increase in iodine content of potato was observed after baking, whereas a significant decrease was observed after boiling. We concluded that iodine biofortified fresh market tomato salad, both from field and hydroponic cultivation, and baked potatoes can be considered potential functional foods for IDD prevention.

Introduction

Iodine is an essential microelement for mammals (Welch and Graham, 1999) with a unique role in organisms. In fact, it is an indispensable component of the thyroid hormones, thyroxine (T₄) and triiodothyronine (T₃), which are involved in the regulation of various enzymes and metabolic processes. Inadequate dietary iodine intake can cause insufficient thyroxin production. This results in iodine deficiency disorder (IDD). The clinical manifestations of IDD include hypothyroidism, goiter, mental retardation, reproductive impairment, deaf-mutism, and lower child survival rates (Hetzel, 1983; Stanbury et al., 1996; Delange et al., 1999). The recommended daily nutrient intake of iodine is 50 μg for infants (aged 0-12 months), 90 μg for children (aged 2-6 years), 120 μg for schoolchildren (aged 7-12 years), 150 μg for adolescents (aged over 12 years) and adults, and 200 μg for pregnant and lactating women (WHO/UNICEF/ICCIDD, 1996). These amounts are thought to allow normal hormone production without stressing the thyroid iodide trapping mechanism. Excess iodine intake is more difficult to define. Many people habitually ingest huge amounts of iodine (approx. 10-200 mg day⁻¹) with no evidence of negative effects (FAO/WHO, 2004).

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In 1993, the WHO and UNICEF recommended universal salt iodization (USI) and iodine supplementation as highly effective strategies for preventing and controlling IDD (UNICEF/WHO, 1994) and USI programs are now applied in most countries. Despite this, approximately 31% (1,900.9 million) of the world population has insufficient iodine intake. The most affected regions are Europe and South-East Asia (WHO/UNICEF/RCCD, 2007). In 2005, Italy introduced legislation requiring retailers to sell only iodized salt; 30 ppm as either potassium iodide (KI) or potassium iodate (KIO₃) unless consumers specifically request otherwise (Moleti et al., 2008). Nevertheless, in Italy there is still a low dietary supply with mild to moderate iodine deficiency (Andersson et al., 2007). Since the effectiveness of USI largely depends on the economic, social, and cultural context of the population in question, an alternative approach of biofortification could also be considered. Biofortification can be defined as the supplementation of trace elements in the food chain through plant uptake, and it is believed to be a cost-effective strategy to reduce mineral malnutrition. Fortified food can be defined as functional foods, i.e., foods that provide a health benefit beyond basic nutritional functions (Henry, 2010).

Iodine is a non-essential element for higher plants, although all plants can assimilate it from the soil. The capacity of the soil to absorb iodine is crucial for iodine fertilization. In humus and clay soils, iodine is fixed by organic matter and aluminium oxides and iron oxides. Therefore, the plants grown in these soils have a low iodine content (Jopke et al., 1996). Iodine fertilization of these types of soils is less efficient in terms of iodine uptake when compared to sandy soils and water culture (Jopke et al., 1996; Zhu et al., 2003; Blasco et al., 2008; Voogt et al., 2010). In recent years, iodine biofortification has drawn the attention of researchers who have studied the capacity of edible crops to accumulate iodine. It has been demonstrated that iodine cannot be easily stored in the grains of cereals. In fact, Mackowiak and Grossl (1999) showed that even with high concentrations of iodide or iodate in growing solutions the iodine concentration in rice grains was still not sufficient to meet the recommended dietary allowance (RDA). Horticultural crops can store iodine, and its uptake increases with the quantity of iodine used, as demonstrated for potato (Caffagni et al., 2011), Chinese cabbage (Weng et al., 2003; Hong et al., 2008), radish (Weng et al., 2003), tomato (Hong et al., 2008; Caffagni et al., 2011; Landini et al., 2011), carrot (Dai et al., 2004; Hong et al., 2008), lettuce (Blasco et al., 2008; Hong et al., 2008; Voogt et al., 2010) and spinach (Weng et al., 2003; Zhu et al., 2003; Dai et al., 2004). Furthermore, different crops were shown to respond differently to increasing iodine dose in soil (Weng et al., 2003; Dai et al., 2004; Hong et al., 2008). However, at higher concentrations, iodine can be toxic, leading to leaf damage (chlorosis and necrosis) and stunted growth (Mackowiak and Grossl, 1999; Weng et al., 2008; Caffagni et al., 2011).

Iodine is stored better in the vegetative organs, such as root, stem and leaves (Dai et al., 2004; Voogt et al., 2010), suggesting differences in iodine accumulation among organs (Weng et al., 2003; Hong et al., 2008). Iodine biofortification studies, conducted in greenhouse conditions, showed an adequate iodine accumulation in crops such as spinach (Zhu et al., 2003; Dai et al., 2004), lettuce (Blasco et al., 2008; Voogt et al., 2010), tomato (Hong et al., 2008; Caffagni et al., 2011; Landini et al., 2011) or potato (Caffagni et al., 2011), making them favorite crops for iodine biofortification programs. Iodine content may also be influenced by other factors such as postharvest processing (e.g. peeling and cooking) and storage. These can largely determine final quality, whether a crop is sold for fresh consumption, or used as an ingredient in a processed food product (Winger et al., 2008).

This paper reports results obtained within the frame of the Italian national project IODOPLAN, aimed at iodine fortification of horticultural and fruit tree species. In particular, it was our aim to find suitable methods of iodine fortification to be applied in normal cultivation practices in Italy. The second aim was to study the effect of different postharvest treatments (peeling, storage and cooking) in one horticultural and one fruit tree species. The third aim was to study the distribution of the applied iodine in various organs of one horticultural and one fruit tree species.

Materials and methods

Plant material and treatments

Four cultivated plant species were used, two fruit trees (plum, Prunus domestica L. cv. Angeleno; nectarine, Prunus persica L. Batsch, cv. Big Top) and two horticultural crops (potato, Solanum tuberosum L. cv. Daisy; fresh market tomato, Solanum lycopersicum L. CRA breeding line Sel. 25/08, a small-fruited breeding line, of Datterino typology).

A field experiment was carried out from 2007 to 2009 on 8-year old plum trees [Prunus domestica L. cv. Angeleno grafted onto GF 677 rootstock (Prunus persica x Prunus amygdalus)]. The experiment was performed at a field site (44°13′59.99″N and 11°30′53.60″E) located in Marzeno (Province of Ravenna, northern Italy), in different plots to avoid any residual effect from the previous year. The orchard was planted in a typical growing area, on a soil classified as a Bathic Cambisols (FAO, 1998), consisting of 19% total CaCO₃, 8% active CaCO₃, 0.7% organic matter, pH 7.8, 27% clay, 58% silt and 15% sand. For replicated treatments and control, each year each plot consisted of a single, non-contiguous, 140 x 4.5 m row of trees.

A field study was conducted from 2007 to 2009 on 8-year old nectarine trees (Prunus persica L. Batsch, cv. Big Top grafted onto GF 677 rootstock (Prunus persica x Prunus amygdalus)). The experiment was performed at a field site (44°29′50.16″N and 11°57′08.40″E) located in Fusignano (Province of Ravenna, northern Italy), in different plots to avoid any residual effect from the previous year. The orchard was planted in a typical growing area, on a soil classified as a Calcaric Gleyic Cambisols (FAO, 1998), consisting of 17% total CaCO₃, 12% active CaCO₃, 2.3% organic matter, pH 7.8, 34% clay, 62% silt and 4% sand. For replicated treatments and control, each year each plot consisted of a single, non-contiguous, 180 x 4 m row of trees.

Potato experiments were conducted in 2007 and 2008 at a field site (44°29′41.08″N and 11°57′20.73″E) located in Fusignano (Province of Ravenna, northern Italy) in two different plots to avoid any residual effect from the previous year. For replicated treatments and control, each year each plot consisted of a 200 x 9 m, non-contiguous, field area, including 10 rows of potato plants (90 cm between rows, 21 cm between plants in the row). The tubers were planted on April 17 and April 23 in 2007 and 2008, respectively. Plum, nectarine and potato experimental fields, including plots, were subject to standard cultivation techniques, under the Regional regulations for integrated pest management (Regione Emilia Romagna). Tomato experiments were conducted in the open field and in the greenhouse. The open field study was conducted in 2007 and 2008 at a field site (42°53′09.47″N and 13°47′52.83″E) located in Monsampolo del Tronto (Province of Ascoli Piceno, central Italy) in two different plots to avoid any residual effect from the previous year. For replicated treatments and control, each year plot size was 7 m², consisting of 20 tomato plants. The soil is classified as a Calcaric-Fluvic Cambisols (FAO, 1998) consisting of 19.4% total CaCO₃, 7.2% active CaCO₃, 2.2% organic matter, pH 7.6, 28.3% clay, 20% silt and 52% sand. Seeds were sown on March 27 and March 25, and were transplanted on May 4 and May 2 in 2007 and 2008, respectively. The tomato field trial was managed with standard cultivation techniques, under the Regional regulations for integrated pest management (Regione Marche). The greenhouse experiments were conducted in 2008 and 2009 in Montanasso Lombardo (Province of Lodi) Italy (45°20′31.01″N 9°26′53.97″E) and the cultivation was carried out in hydroponic culture. Tomato plants were grown in an open rockwool sys-
tem as spring crop. We used Rockwool Grodan Master slabs (100x20x7.5 cm) filled with the organic medium. In 2008, tomato seeds were sown individually in special rockwool minicubes on March 3, and after two weeks minicubes containing tomato seedlings were inserted into rockwool starter cubes (10x10 cm). Tomato transplants were placed onto rockwool production slabs on April 29. Tomato plants were grown until the 7th cluster. In 2009, all the cultural practices were repeated as in 2008, except for the dates on which sowing and transplantation on rockwool slabs took place (March 25 and May 15, respectively). For replicated treatments and control, each year each plot consisted of 8 plants.

In the open field experiments two treatment systems were used, and two pre-market preparations of fertilizer containing potassium iodide (KI) and coformulants (BMS Micro-Nutrients NV, Bornem, Belgium) were applied. The treatments were as follows: i) foliar spray application of a liquid fertilizer with 25 g L\(^{-1}\) I; and ii) soil application of a crystalline fertilizer with 90 g Kg\(^{-1}\) I. The dose of iodine called N (125 g ha\(^{-1}\)) was used as reference. The iodine dose was adjusted each year in order to increase iodine concentration in the plants. Details of all treatments and doses in each year of trials (2007, 2008 and 2009) are summarized in Table 1. For nectarine trees, both soil and foliar spray treatments were applied. For plum tree, only foliar spray treatments were applied. For potato and open field tomato, soil, foliar spray, and soil plus foliar spray treatments were applied. In all field trials, a control plant without any iodine application was included every year. Irrigation and treatments were applied by an automated trickle irrigation system, and each plum tree, nectarine and potato treatment was replicated twice according to a completely randomized design, except for 2007, when they were performed without replicates. Tomato field treatments were replicated three times, according to a randomized block design, except for 2007, when they were performed without replicates.

In hydroponic culture, only the 25 g L\(^{-1}\) I liquid fertilizer (BMS Micro-Nutrients NV) was used for experiments; three treatments, 1 mM, 2 mM and 5 mM of potassium iodide (KI), were applied to tomato plants. The control treatment consisted of applying the complete mixture on rockwool production slabs on April 29. Tomato plants were grown until the 7th cluster. In 2009, all the cultural practices were repeated as in 2008, except for the dates on which sowing and transplantation on rockwool slabs took place (March 25 and May 15, respectively). For replicated treatments and control, each year each plot consisted of 8 plants.

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<tr>
<td>2N</td>
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<td>Plum tree</td>
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<td>Potato</td>
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<td>N</td>
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<td>-</td>
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<td>2N</td>
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<td>-</td>
<td>20N+5N</td>
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<tr>
<td>40N**</td>
<td>2N</td>
<td>-</td>
<td>20N+5N</td>
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<tr>
<td>Tomato</td>
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<tr>
<td>N</td>
<td>-</td>
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<tr>
<td>2N</td>
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</tr>
<tr>
<td>40N</td>
<td>2N</td>
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<td>2N</td>
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<td>8N</td>
<td>-</td>
<td>-</td>
<td>2mM</td>
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<tr>
<td>N+N</td>
<td>20N+8N</td>
<td>1mM</td>
<td>2mM</td>
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<td></td>
<td>2N</td>
<td>2mM</td>
<td>5mM</td>
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</table>

* Treatments from which samples for post harvest, peeling and cooking experiments were collected; ** treatments from which also samples of vegetative parts were collected.

### Processing and storage treatments

Effect of postharvest storage and processing (cooking) for potato tubers, and storage and processing (peeling) for nectarine fruits were analyzed in 2008. Nectarine fruit peeling was performed by simply eliminating 2-3 mm thick peel with a clean knife. For potato tuber, two cooking methods were used, and processing times and temperatures were based on the optimum times and required temperatures to cook the average sample.

**Boiling**

Water was brought to the boil on a gas stove. Potato samples treated with doses of iodide (N) of 2 N, and 40 N were cooked for 25 min in the boiling water. After cooking, the leachate was removed and discarded. The cooked potato sample was frozen at -20°C until iodine determination.

**Bake**

A gas oven was heated to 200°C. Potato samples treated with a dose of 40 N were cooked for 45 min. Cooked samples were frozen at -20°C until iodine determination.

In order to analyze the effect of postharvest storage, samples of potato tubers, treated with doses 2 N, and 40 N, and of nectarine fruits, treated with dose N (Table 1), harvested in the periods indicated above, were placed in a cold storage room for six months at 10°C, and for two weeks at 5°C, respectively.
Iodine content analysis

Iodine content in plum and tomato fruit was determined with peel and in potato tubers without peel. In nectarine fruit, due to consumption habits, iodine content was determined both with and without peel. Iodine content in plant tissues was determined using the USEPA Method 3052 (HNO₃-H₂O 2, microwave digestion), and an inductively coupled plasma mass spectrometry (ICP-MS) analysis procedure. Chemical analyses were performed at NEOTRON S.p.a. (Santa Maria di Muggiano, Modena, Italy). The closed digestion was performed by placing the sample in a polytetrafluoroethylene (PTFE) vial (or bomb). After adding the digestion reagents (10 mL HNO₃ 65% and 2 mL H₂O 2 30%), the bomb was hermetically sealed and located in a microwave oven for irradiation. The determination of iodine by ICP-MS was performed by using isotope dilutions of I¹²⁷, and iodine concentrations in the samples were determined by means of a calibration curve obtained with the method of standard additions (Larsen and Ludwigsen, 1997).

Statistical analysis

Results obtained were expressed as means ± SEM and statistical analysis was performed according to the Systat 12 statistical package (Systat Software, Inc.). Data were analyzed using a general linear model (GLM). Duncan’s multiple range test (DMRT) was used to examine differences between groups and among group means. P≤0.05 was considered statistically significant.

Results

Iodine accumulation in crops

Tables 2, 3, 4 and 5 show the relation between iodine content in the edible portion of different crops, and the dose of exogenous iodine in soil, spray and hydroponic applications.

Table 2 shows iodine accumulation in nectarine and plum fruit over the three years of the experiment. The iodine content in nectarine fruit increased significantly after foliar spray application at dose N only in 2007, while in 2008, the increase was significant only at the highest dose of applied iodide (2.5 N). In the first year of the experiment (2007), iodine content in nectarine fruit treated with dose N applied as foliar spray was determined with and without peel (Figure 1A) and no significant change in iodine content was observed. Thus, in the following two years, iodine content in fruit was determined only with peel. The iodine content in plum fruit was determined only with peel. A significant increase in iodine content in plum fruit was observed already at the lowest dose of iodide (N) applied as a foliar spray (2008). Considering all the treatments applied (2007-2009), nectarine and plum tree accumulated only low amounts of iodine in their edible tissues: from 4.0 to 14.3 μg 100 g⁻¹ and from 5.6 to 9.5 μg 100 g⁻¹, respectively (Table 2).

Table 2. Iodine accumulation in nectarine and plum fruits during three years of experiment.°

<table>
<thead>
<tr>
<th></th>
<th>2007</th>
<th>2008</th>
<th>2009</th>
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<tbody>
<tr>
<td></td>
<td>Soil treatment</td>
<td>Spray treatment</td>
<td>Soil and spray treatment</td>
</tr>
<tr>
<td></td>
<td>Control N</td>
<td>2N</td>
<td>Control N</td>
</tr>
<tr>
<td>Nectarine</td>
<td>0±0ab</td>
<td>0±0a</td>
<td>0±0a</td>
</tr>
<tr>
<td>Plum tree</td>
<td>0.7</td>
<td>0.7</td>
<td>3.3±0.1b</td>
</tr>
</tbody>
</table>

°Iodine content (μg 100 g⁻¹, FW) is expressed as replicate means ± SEM. a,bValues followed by different letters are significantly different by Duncan’s multiple range test (P≤0.05).

Table 3. Iodine accumulation in potato tubers during two years of experiment.°

<table>
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<tr>
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<th>2007</th>
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<tbody>
<tr>
<td></td>
<td>Soil treatment</td>
<td>Spray treatment</td>
</tr>
<tr>
<td></td>
<td>Control N</td>
<td>2N</td>
</tr>
<tr>
<td>Potato</td>
<td>0.9±0.1b</td>
<td>2.0±0.2a</td>
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</table>

°Iodine content (μg 100 g⁻¹, FW) is expressed as replicate means ± SEM. a-dValues followed by different letters are significantly different by Duncan’s multiple range test (P≤0.05).

Table 4. Iodine accumulation in fruits of tomato grown in open field during two years of experiment.°

<table>
<thead>
<tr>
<th></th>
<th>2007</th>
<th>2008</th>
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<tbody>
<tr>
<td></td>
<td>Soil treatment</td>
<td>Spray treatment</td>
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<td>Control N</td>
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<tr>
<td></td>
<td>Control N</td>
<td>20N</td>
</tr>
<tr>
<td>Tomato</td>
<td>0.8</td>
<td>0.6</td>
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°Iodine content (μg 100 g⁻¹, FW) is expressed as replicate means ± SEM. a,bValues followed by different letters are significantly different by Duncan’s multiple range test (P≤0.05).

Table 5. Iodine accumulation in fruits of tomato grown in hydroponic culture during two years of experiments.°

<table>
<thead>
<tr>
<th></th>
<th>2008</th>
<th>2009</th>
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<tbody>
<tr>
<td></td>
<td>Hydroponic treatment</td>
<td>Control</td>
</tr>
<tr>
<td></td>
<td>1 mM</td>
<td>2 mM</td>
</tr>
<tr>
<td>Tomato</td>
<td>4.6±1.6a</td>
<td>454.5±151.5b</td>
</tr>
</tbody>
</table>

°Iodine content (μg 100 g⁻¹, FW) is expressed as replicate means ± SEM. a,bValues followed by different letters are significantly different by Duncan’s multiple range test (P≤0.05).
years of the experiments. The iodine content in tubers was determined only without peel. During the first year of treatment, no significant uptake was observed after soil treatments at the lowest dose N and 2 N doses; therefore, during the second year, the iodine doses were increased to 20 and 40 N. Furthermore, foliar spray treatments were also applied, alone and in combination with soil treatment at 20 N. The iodine content in potato tubers increased as a greater amount of iodine was applied. The iodine uptake was higher after soil treatments than after foliar spray treatments, and no significant increase was observed when soil applications were combined with foliar applications. There was a significant increase in iodine content compared to the control plants after soil treatments at 20 and 40 N and after foliar spray treatment at 5 N; the amounts of iodine accumulated in the tubers were 25.0, 89.4 and 5.7 μg 100 g⁻¹, respectively (Table 3).

Table 4 shows iodine accumulation in tomato fruit grown in the open field over the two years of experiments. During the first year of treatment, no significant difference in uptake was observed either after soil treatments at N and 2 N, spray treatment at N or after a combination of soil and spray treatments; therefore, during the second year, the iodine doses were increased. Significant uptake was observed after foliar spray application at 8 N (94.6 μg 100 g⁻¹); however, no significant uptake was obtained after soil treatments and no significant increase in iodine uptake was observed when foliar applications were combined with soil applications (Table 4). During 2007, the iodine uptake obtained in fresh market tomato grown in the open field was unsatisfactory; therefore, since 2008, a hydroponic system has been included.

Table 5 shows the iodine accumulation in fruit of tomato grown in hydroponic culture over the two years of experiments. In 2008, a significant iodine uptake was observed at a lower dose of iodide (1 mM) and no significant increase was observed with a further increase in the dose. Significant iodine uptake was observed with the lower dose of iodide (2 mM) in 2009 and this was higher than in 2008. Furthermore, no significant increase was observed with a further increase in the dose. Tomatoes grown in hydroponic culture successfully accumulated large amounts of iodine in their edible tissues (454 to 2423 μg 100 g⁻¹) although this varied widely among replicates.

**Effect of postharvest storage and processing on iodine uptake**

No significant change in iodine content was observed either in nectarine fruit after storage and peeling, or in potato tubers after storage (Figure 1). The effect of cooking on iodine content was investigated in potato tubers (Figure 1B). A significant increase in iodine content was observed after baking whereas a significant decrease in iodine content was observed after boiling.

**Iodine distribution in different plant organs**

Figure 2 shows iodine distribution in different organs of plum tree (Figure 2A) and potato (Figure 2B) as a function of iodine doses applied. Iodine content is expressed as dry weight (DW). Iodine was mainly accumulated in the leaves and only a small portion of iodine uptake was transported to the plum tree branches and fruit, and to potato stems and tubers. The ratio of I in leaves to I in fruit in plum tree was 35:1, 71:1 and 59:1 for foliar sprays treatments at N, 2 N and 2.5 N, respectively, and the ratio of I in leaves to I in branches was 4:1, 9:1 and 11:1 for foliar sprays treatments at N, 2 N and 2.5 N, respectively (Figure 2A). The ratio of I in leaves to I in tubers in potato was 343:1 for foliar spray treatment at 5 N whereas for soil treatment at 40 N, the ratio was 9:1, and the ratio of I in leaves to I in stems was 9:1 for foliar spray treatment at 5 N and for soil treatment at 40 N the ratio was 3:1 (Figure 2B).

Iodine content in leaves of plum tree increased with increasing amounts of iodine whereas a significant increase was already observed in branches and in fruit of plum at the lowest dose (N). Increasing the dose of application did not lead to any significant further increase in iodine content (Figure 2A). Iodine content in potato leaves was higher after foliar spray treatment than after soil treatment. A significant increase in iodine content was observed in potato stems under both

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*Article* [Italian Journal of Agronomy 2012; 7:e32] [page 233]
treatments compared to the control, but no significant change in iodine content was observed after foliar spray treatment at 5 N compared with soil treatment at 40 N. Iodine content in potato tubers was higher after soil treatment than after foliar spray treatment (Figure 2B).

Discussion

In this study, the uptake of iodine in soil, spray and hydroponic applications, and its accumulation in the edible portion of plants were studied in two fruit tree and two horticultural species. Clear differences in iodine accumulation were observed among the four crops, between applications (soil and foliar spray) and between production systems (open field and greenhouse hydroponics). As regards differences among the four crops observed in the open field system, the maximum iodine content ranged from 9.5 and 14.3 μg 100 g⁻¹ for plum and nectarine fruit, and to 89.4 and 144.0 μg 100 g⁻¹ for potato tuber and tomato fruit, respectively (Tables 2, 3 and 4). Iodine doses could not be increased in both fruit tree species to increase the iodine accumulation in edible tissues because of phytotoxic symptoms shown by foliar-sprayed plants at increased doses. The iodine content in tomato fruit and potato tubers was much higher than in nectarine and plum fruit. This suggests that these horticultural crops (potato and fresh market tomato) might be a better target for strategies to improve dietary iodine supply. As far as differences between soil and foliar applications are concerned, the iodine content in nectarine, plum and tomato fruit increased significantly only after foliar spray treatments, even though fruit trees, nectarine and plum, accumulated much lower amounts of iodine in their edible tissues than tomato (Tables 2 and 4) while the iodine uptake in potato tubers was higher after soil treatments than after foliar spray treatments (Table 3). In order to further clarify the mechanisms of iodine uptake, we investigated the distribution of iodine in potato stem and leaves, and in plum tree fruit, leaves and branches. The transpiring plant parts, i.e. the leaves, contained much more iodine than the fruit and tubers, in agreement with data reported by other authors (Weng et al., 2003; Zhu et al., 2003; Dai et al., 2004; Hong et al., 2008; Voogt et al., 2010). Indeed, the flow of mineral nutrients and water from roots to shoots through the xylem is reduced in organs with low rates of transpiration such as fruit and tubers (Herrett et al., 1962).

It has been suggested that plant root cells take up iodine as the iodide anion (Umaly and Poel, 1971; Mackowiak and Grossi, 1999; Zhu et al., 2003; Blasco et al., 2008; Caffagni et al., 2011), and that I⁻ follows the chloride (Cl⁻) transport pathway with H⁺/anion symporters catalysing I⁻ uptake and anion channels releasing I⁻ into the xylem (White and Broadley, 2001; Roberts, 2006). Other authors demonstrated that iodine was easily accumulated in roots and leaves (Mackowiak and Grossi, 1999; Zhu et al., 2003; Dai et al., 2004, 2006; Kashparov et al., 2005; Mackowiak et al., 2005; Blasco et al., 2008), and little iodine was considered to have been reallocated via the phloem to fruit or seeds (Herrett et al., 1962; Muramatsu et al., 1993, 1995). However, Landini et al. (2011) showed that iodine was also stored in tomato fruit after leaf treatment. This suggests that a moderate phloem flux of iodine occurred, in agreement with our data collected in the open field where a significant increase of iodine in tomato fruit was obtained only after foliar spray treatments. In the case of potatoes, it was suggested that xylem connections between tubers and basal roots are non-functional (Kratzke and Palta, 1985), so iodine might be directly absorbed from soil through the epidermis. This might explain the higher iodine uptake in potatoes treated with soil application.

The hydroponic system provides excellent opportunities for biofortification with iodine (Blasco et al., 2008; Voogt et al., 2010). This is the most efficient system to control plant uptake of nutrients as the iodine concentration in the root environment can be managed quite simply. Fresh market tomato is one of the crops grown commercially in hydroponic systems and was shown to be a very suitable crop for iodine uptake (Caffagni et al., 2011). In the present study, tomatoes grown in hydroponic culture successfully accumulated up to 2423 μg 100 g⁻¹ of iodine in their edible tissues, much more than tomato grown in an open field system. Considering a single serving of this vegetable as 100 g of tomato salad, the iodine content obtained in hydroponic culture was approximately 3-fold the RDA (determined for adults as 150 μg day⁻¹) for plants treated with the lowest dose of application (1 mM) and 16-fold RDA for plants treated with 5 mM. The results of the open field experiments showed successful iodine accumulation in tomato fruit treated with foliar spray application at 8 N. Considering a single serving of this vegetable as 100 g of tomato salad, the iodine contents were...
A second aim of this work was to investigate the iodine content of nectarine fruit after both postharvest storage and peeling, and the iodine content of potato tubers after both postharvest storage and cooking. No significant change in iodine content was observed between peeled and unpeeled fruit, suggesting that iodine might be stored within the flesh of fruit and therefore consumption habits do not influence the iodine intake. Food preservation methods, such as deep freezing and freeze drying, were reported to reduce the iodine content of food by as much as 20-25% (Lee et al., 1994). However, our data show no significant change in iodine content after postharvest non-freezing storage either in nectarine fruit or in potato tubers (Figure 1). These findings showed that non-freezing storage could be the most appropriate method for preserving these crops once enriched. It has been shown that cooking with water or steam causes a greater loss of iodine in comparison to, for example, roasting or frying (Goindi et al., 1995). In fact, frying and grilling were reported to reduce the iodine content of food by 25%, while boiling may result in a loss of up to 60% (Lee et al., 1994). This was confirmed by our data that showed that iodine content was reduced by 65% after boiling (Figure 1B). Finally, it has been suggested that cooking methods with high-temperature oil and baking might, therefore, be the best way to increase iodine content and to prevent IDD, since a single serving (100 g) of baked potato treated with soil application at 40 N was enough to reach full RDA.

Conclusions

Horticultural and fruit tree species can absorb and accumulate iodine when exogenous iodine is applied. However, an obvious discrepancy is observed in its absorption among crops. Based on the results of these experiments, it can be concluded that priority should be given to horticultural crops as candidates for iodine biofortification and that salad of fresh market tomato treated with foliar spray application at 8 N and baked potatoes after soil application at 40 N could be considered potential functional foods for IDD prevention. Our results suggest that iodine might be stored within the flesh of fruit and that non-freezing might be the most appropriate storage method for the crops studied in order to retain the iodine content. Furthermore, our results confirmed that hydroponic culture is the most efficient system for nutrient uptake. However, sufficiently good results of iodine accumulation in two horticultural crops were also obtained in normal Italian cultivation practice.

References


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