The Effect of Growth Medium Temperature on Corn Salad [Valerianella locusta (L.) Laterr] Baby Leaf Yield and Quality

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Abstract. Soil temperature has a crucial impact on physiological processes and growth of plants with important consequences for plant productivity and food safety including nitrate accumulation in leaf blades of leaf vegetables. Consumer demand for high-quality, fresh-cut vegetables has increased rapidly in the last decades, and temperature modulation can help control nitrate concentration in fresh vegetables, an important trait of product safety. Corn salad plants [Valerianella locusta (L.) Laterr., cultivar Gala] were grown at three root temperatures (15, 20, and 25 °C) in a floating system. This experimental setup allowed to directly evaluate the effect of root temperature on yield and plant quality excluding the effect on soil processes and properties. Nutrient solution was renewed weekly and kept aerated while air temperature was maintained constant at 20 °C for all treatments during the entire time of experiments. At harvest, plants were collected, the shelf life evaluated, and the nutrient uptake [NO₃⁻, iron (Fe) from ⁶⁰Fe-O,0EDDHA, and ³²⁵⁰SO₄²⁻] and mineral content were determined. Results showed that growing conditions at 20 °C of the nutrient solution led to the best plant performance in terms of yield, nitrate content at leaf level, root biomass, leaf area, and greenness with positive effects on postharvest quality, i.e., less rapid leaf loss of greenness and leaf fresh weight (FW) loss during conservation at 4 °C. At this temperature condition of the nutrient solution, it has also been observed an enhanced functionality of mechanisms involved in the acquisition of nutrients like NO₃⁻, Fe, and SO₄²⁻, which are known to play an important role in nitrate level in leaf tissues of crops. Plants grown at 15 °C showed minor growth, whereas the nutrient solution at 25 °C caused stress for the plants affecting negatively the quality and yield. Overall, the results obtained showed that root temperature plays a fundamental role in several plant processes that affect yield and its quality; for hydroponic system cultivations, a level of growing-medium temperature close to that of the surrounding air seems suitable.
changing of developmental and environmental demands on C and nitrogen (N) resources, have evolved highly sophisticated and complex sensory systems that “crosstalk” to regulate C or N assimilation, metabolism, and transport (Coruzzi and Bush, 2001; Coruzzi and Zhou, 2001). For this reason, a small change in the soil temperature can have a profound impact on physiological processes and growth of plants (Clarkson et al., 1992; Moorby and Nye, 1984; Pregitzer and King 2005; Zhang and Deng, 2007) with serious consequences on food productivity and safety such as nitrate accumulation. So, it appears evident that the possibility of controlling the production process from the temperature point of view is extremely important and relevant considering also that consumer demand for high-quality, fresh-cut vegetables is increasing in the last decades (Brech et al., 2004). However, for the processes occurring in the soil component, responses of roots to changes in soil temperature cannot be discriminated between those directly caused by biochemical and physiological processes and those related to simultaneous changes in soil properties like nutrient availability.

Information on the soil-temperature effects on root functionality appears of importance also for the soilless cultures in which the temperature of the nutrient solution can easily be modified by environmental conditions particularly in summer (heating) and in winter or early spring (cooling) periods. As for soil-grown plants, this can consistently affect several physiological functions (Nxawe et al., 2010) with undesired consequences on yield. The more this water-saving cultivation systems is being used (as happens in arid or semiarid regions such as the Mediterranean area where water, owing to its scarcity, costs, and quality is becoming an economically valuable resource), the more the knowledge of the effects of the growth-medium temperature on cultivated plants becomes relevant, particularly when yield could be hindered and/or its quality worsened.

For these reasons, the present work studied the effect of different temperatures of the growing medium on the production and quality of corn salad [Valerianella locusta (L.) Laterr.; cultivar Gala] plants grown in soilless culture to exclude the effect of nutrient availability on the process. Functionality of mechanisms involved in NO$_3^-$, SO$_4^{2-}$, and Fe$^{2+}$ acquisition has also been considered in relation to the levels of nitrate accumulated in the edible tissues.

**Material and Methods**

A flow chart summarizing the different treatments and analyses performed in this research is presented as Figure 1. **Plant material and growth conditions.** Plants were grown hydroponically in a growth chamber as described by Iacuzzo et al. (2011) with the following controlled climatic conditions: day/night photoperiod, 16/8; radiation, 220 µE·m$^{-2}$·s$^{-1}$; air temperature (day/night) 20 ± 0.9 °C; relative humidity 70% to 80% (Cesco et al., 2006). Briefly, corn salad seeds [Valerianella locusta (L.) Laterr.; cultivar Gala from DOTTO SpA, Italy] were sown onto expanded polystyrene boards. After emergence, the boards were transferred into rectangular pots containing the aerated nutrient solution described by Manzocco et al. (2011). The nutrient solutions were renewed every week and aerated by bubbling to prevent anoxia and to guarantee the constant mixing of the solution.

Three sets of pots were used at three different temperatures of nutrient solution, 15, 20, and 25 °C, and were named T15, T20, and T25. These values of temperature were chosen considering the results of previous experiments performed in a greenhouse with plants grown both in soil and hydroponically, where the temperature of air, soil, and nutrient solution were monitored. These measurements showed significant daily temperature fluctuation in sunny weather (Fig. 2) with average values of temperature for soil and water in the range of 15 to 25 °C depending on climatic conditions. Thus, in our experiments, the temperature of air in the growth chamber was set at the mentioned 20 °C values for all treatments during all time of the experiments and that of the nutrient solutions at ≈15, 20, and 25 °C. Because the temperature of the solution at the middle temperature (20 °C) corresponded with that of ambient conditions, it was not controlled. The pots with the solution at 15 and 25 °C were supplied with a pipe at the bottom, where cold (14 °C) or hot (26 °C) water circulated to maintain the solution at the correct temperature without any contamination. Cold and hot water were supplied by independent cooling and heating systems appropriately designed and whose activation was controlled by the temperature of the solution. The temperature of the nutrient solution in all the pots as well as of the air in the growth chamber was measured by several

![Fig. 2. Temperature profiles of air, nutrient solution, and soil in a typical week of May during the growing period of corn salad plants in the greenhouse.](image-url)
thermocouples "T" and a data acquisition unit (Agilent Technologies Italia S.p.A., Cernusco, Italy) at time intervals of 5 min during the entire experiment. The temperature values of the nutrient solution as recorded during the experiment were 15.2 ± 0.2 °C, 19.8 ± 0.6 °C, and 24.7 ± 0.4 °C.

**Sampling.** At the end of the growing period in hydroponics (45 d), the plants, with the exclusion of those at the board margins, were sampled, divided in root and shoot, and used for the analytical determinations and for the uptake assays.

**SPAD and inductively coupled plasma measurements.** Fresh biomass of plant tissues and leaf area, determined by using a LI-3100 Area Meter (LI-COR Inc., Lincoln, NE), were presented per square meter of floating board. SPAD index values of fully expanded young leaves were determined using a portable SPAD-502 m (Minolta, Osaka, Japan). Nutrient [phosphorus (P), S, potassium (K), calcium (Ca), magnesium (Mg), Fe, manganese (Mn), zinc (Zn), and copper (Cu)] contents in leaf tissues were determined, after their digestion with H2O2, as follows: inorganic P was quantified spectrophotometrically at 705 nm as described for Forbush (1983); K, Ca, Mg, Fe, Mn, Zn, and Cu contents were analyzed by inductively coupled plasma atomic emission spectrometry (VISTA MPX, Varian, Torino, Italy) as described by Zucchini et al. (2009). To determine total S concentration, according to Astolfi et al. (2006), dried leaf samples were ashed in a muffle furnace at 600 °C; the ashes were dissolved in 10 mL of 3 M HCl and filtered through Whatman No. 42 paper. In contact with BaCl2, a BaSO4 precipitate is formed, which is determined turbidimetrically.

**Washing, packaging, and storage.** At harvest, shoot samples were collected and quickly washed with distilled water at 8 °C for 3 min. The salin/water ratio during washing was 1:18 w/w. Aliquots of 50 g of corn salad were packed under air in 30 × 40-cm plastic bags. Samples were stored in the dark at 4 °C for up to 26 d. After 5, 8, 14, 19, and 26 d of storage, SPAD index value and weight loss (%) of packed corn salads were determined. Weight loss was assessed by weighing the content of the packages before and after the storage period. Weight loss was expressed as the percentage of weight loss with respect to the initial weight.

**Measurement of net NO3 uptake by roots and determination of nitrogen and NO3 contents in leaf tissues.** Nitrate uptake by root was measured as described by Nikolic et al. (2007) using excised roots (≈0.8 g of FW) of corn salad plants (45 days old) and an uptake solution (20 mL) containing 1 mM KNO3. To evaluate the functionality of the mechanisms involved in the acquisition process of the nutrient at the growing conditions, uptake medium was maintained at the temperature of 15, 20, or 25 °C for the root tissues of plants grown at 15, 20, or 25 °C, respectively. The NO3 depletion from the uptake solution was measured over 10 min by removing every 2 min 0.2-mL aliquots and the concentration of NO3 determined spectrophotometrically. The net NO3 uptake rate was calculated by linear regression analysis and expressed as μmol N03/g dry weight (DW) of root per hour. Cell juices of leaf samples were prepared by thawing the leaf tissues followed by centrifugation at 10,000 g for 15 min. The concentration of NO3 in leaf samples was determined spectrophotometrically. Total N has been also measured in these tissues using a CHN analyser.

**Measurements of SO4²⁻/Fe²⁺ uptake.** The capability of the root to acquire 35SO4²⁻ was assessed as described by Astolfi et al. (2006) with slight modifications. Briefly, excised root sampled as previously described were washed with water. After 30 min, roots were transferred to beakers containing 20 mL of a freshly prepared micronutrient- and SO4²⁻-free uptake solution. Sulphate (35SO4²⁻, specific activity 2.1 KBq μmol⁻¹ 35SO4²⁻) was added at a 0.6 mM concentration and the uptake period was 30 min. To evaluate the functionality at the growing conditions (temperature of 15, 20, or 25 °C of growing medium) of the mechanisms involved in the acquisition process of the nutrient, uptake medium was maintained at the temperature of 15, 20, or 25 °C. Thereafter, the root tissues were transferred to an ice-cold desorption solution containing 0.6 mM CaSO4 and 10 mM MES [2-(N-morpholino)ethanesulfonic acid]–KOH (pH 6.0) for 30 min. Roots were oven-dried at 60 °C, weighed, mineralized with 10% H2O2 at 60 °C, and suspended in 1 M HCl for 35SO4²⁻ determination by liquid scintillation counting. The 35SO4²⁻ uptake rate is presented in μmol 35SO4²⁻/g DW of root per hour.

**Measurements of 59Fe uptake from Fe–oEDDHA.** The capability of the root to acquire 59Fe from 59Fe(III)-oEDDHA was assessed as described by Cesco et al. (2002) with slight modifications. To achieve this, excised roots were washed with CaSO4 0.5 mM for 30 min and then transferred to beakers containing 20 mL of a freshly prepared micronutrient-free uptake solution having the following composition (mM): K2SO4 0.7, KCl 0.1, Ca(NO3)2 2.0, MgSO4 0.5, K2HPO4 0.1, MES–KOH 10 (pH 6.0). Iron(59Fe)-oEDDHA, prepared by mixing 59FeCl3 with EDDHA with a molar ratio of 1:1.1 (specific activity of 114 KBq μmol⁻¹ Fe; Rodriguez-Lucena et al., 2009) was added to give a final Fe concentration of 100 μM. To limit photochemical reduction of the micronutrient in the uptake solution added by the Fe source (Hernández-Apaolaza and Lucena, 2011; Zancan et al., 2006), beakers were covered with black plastic foils during the entire experiment. The uptake solution was buffered at pH 6.0 with 10 mM MES–KOH and the uptake period was 30 min. To evaluate the functionality of the mechanisms involved in the acquisition process of the nutrient at the growing conditions (temperature of 15, 20, or 25 °C of the growing medium), uptake medium was maintained at a temperature of 15, 20, or 25 °C. Thereafter, plants were transferred to a freshly prepared 59Fe-free nutrient solution for 10 min to remove the excess of 59Fe at the root surface and then harvested. Root apoplastic 59Fe pools were removed by 1.2 g L⁻¹ sodium dithionite and 1.5 mM 2,2′-bipyridyl in 1 mM Ca(NO3)2 under bubbling N2. Root tissues were oven-dried at 80 °C, weighed, ashed at 550 °C, and suspended in 1 M HCl for 59Fe determination by liquid scintillation counting. The 59Fe uptake rate, measured as μmol 59Fe, refers to the whole root tissues and is presented per g/DW of roots per hour.

**Statistical analysis.** Each experiment was repeated three times in climate chamber over a period of 2 years and the significance of the difference between the means was calculated using the analysis of variance analyses (Fisher's least significant difference) with SigmaPlot (Version 11; Systat Software, Evanston, IL).

**Results.** Plants were able to accumulate the highest values of fresh leaf biomass (1560 ± 85 g FW/m²) when grown at the constant temperature of 20 °C (T20) of the nutrient solution. A 5 °C decrease in the nutrient solution temperature (T15) led to a significant decrease of yield (fresh biomass) and a warming of the solution by 5 °C (T25) severely limited the yield (Table 1). The difference in productivity among the three temperatures was not attributable to leaf thickness, because leaf specific weight, indicated by leaf area per gram, did not show any significant difference among the three treatments. Number and size of leaf components were responsible for the overall difference in yield. Focusing on the

<table>
<thead>
<tr>
<th>Nutrient solution temp. (°C)</th>
<th>Leaf yield (g FW/m²)</th>
<th>Leaf area (m² m⁻¹)</th>
<th>Number of leaves (no. per plant)</th>
<th>SPAD index</th>
<th>Dry weight percentage</th>
<th>NO3 (g kg⁻¹ leaf FW)</th>
</tr>
</thead>
<tbody>
<tr>
<td>15</td>
<td>1260 ± 91 B</td>
<td>4.11 ± 0.19 A</td>
<td>6.6 ± 0.4 B</td>
<td>37.5 ± 1.0</td>
<td>6.7 ± 0.2</td>
<td>4.46 ± 0.21 A</td>
</tr>
<tr>
<td>20</td>
<td>1560 ± 185 A</td>
<td>3.84 ± 0.44 A</td>
<td>7.7 ± 0.6 A</td>
<td>38.9 ± 0.9</td>
<td>7.8 ± 0.2</td>
<td>3.94 ± 0.08 B</td>
</tr>
<tr>
<td>25</td>
<td>897 ± 129 C</td>
<td>3.43 ± 0.38 A</td>
<td>5.9 ± 0.8 B</td>
<td>34.9 ± 1.2</td>
<td>8.2 ± 0.1</td>
<td>4.11 ± 0.09 B</td>
</tr>
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</table>

*Air temperature has been maintained constant at a value of 20 °C for the entire experiment. Data of number of leaves per plant, SPAD index values, DW/FW ratios, and nitrate contents are also reported. Data are means ± SD of three independent experiments; capital letters refer to statistically significant differences among the samples (analysis of variance; Fisher’s least significant difference, P < 0.05). FW = fresh weight; DW = dry weight.*
number of leaves, plants grown T20 had significantly higher numbers of leaves than those obtained in T15 and T25; plants grown at lower temperature (T15) exhibited yield restriction with respect to those grown at 20 °C with a restrain value smaller than that observed in T25. As shown in Table 1, leaves were progressively less hydrated as temperature increased. In fact, the DW percentage of leaves was significantly higher in leaf blades of plants grown in T25, reporting the lowest water content. Significantly higher hydration values were observed in plants of T20 and T15. Corn salad leaves grown in T15 and T20 presented similar chlorophyll contents as measured with the SPAD index value. However, this parameter was significantly lower when salad was grown in T25, indicating lower greenness of the leaves. Concerning NO₃ concentration in leaf tissues, no significant differences could be observed between T25 and T20, whereas a significant increase in NO₃ concentration was detected in plants grown in T15 (Table 1). Figure 3 shows the amount of N accumulated by leaves of plant; data are expressed as grams per square meter of floating boards. When plants were grown at 20 °C, the highest accumulation of N has been recorded; on the contrary, the lowest level of N was measured in plants exposed to 25 °C. Nitrate forms represent approximately one-fourth of the total amount, exhibiting the lowest ratio in plant grown at 20 °C. Leaf nutrient concentrations at harvest are reported in Table 2. Higher concentration values of P, S, Zn, and Cu were observed in leaf tissues of T25 plants with respect to those measured when the nutrient solution was maintained at 20 °C; conversely, T15 plants exhibited a higher accumulation of K and Mg (Table 2).

![Fig. 3. Effect of the nutrient solution temperature on nitrogen (N) accumulation in leaf blades of corn salad (Valerianella locusta (L.) Laterr., cultivar Gala) grown for 45 d in nutrient solution maintained at a constant temperature of 15, 20, or 25 °C. Air temperature has been maintained constant at a value of 20 °C for the entire experiment. Data, expressed as per square meter of cultivation board, are referred to total N accumulation (black box) and N accumulated as nitrate (shaded box); ratios (%) between N as nitrate and total N are also reported. Data are means ± sd of three independent experiments; capital letters refer to statistically significant differences among the samples (analysis of variance, Fisher’s least significant difference, P < 0.05).](Image 62x203 to 182x309)

Figure 4A points out the effects of the three levels of temperature on root biomass. When corn salad plants were grown in nutrient solution maintained at 20 °C, they were able to develop a root system of 89 g FW/m². A cooling of the nutrient solution to reach the constant value of 15 °C led the plants to accumulate a greater, but not significant, root biomass, whereas warming of the hydroponic solution at 25 °C significantly limited the root yield. Considering the leaf yield at harvest, when plants were grown at T15, a ratio between root and leaf fresh biomass greater than those measured for plants treated at the other two temperatures was recorded (Fig. 4B). To evaluate the functionality of NO₃⁻, Fe(III)-chelate, and SO₄²⁻ acquisition mechanisms operating at the root level of corn salad plants grown at three levels of temperature of the nutrient solution, uptakes of NO₃⁻, Fe from ⁵⁷Fe(III)-o,oEDDHA, and ³⁵SO₄²⁻ (Table 3) were measured using excised roots from plants at the harvested stage and maintaining the temperature values of uptake media corresponding to those of the nutrient solutions. As shown in Table 3, the rise in temperature from 15 to 20 °C induced a clear increase in nutrient acquisition capability of NO₃⁻ (+81%), ³⁵SO₄²⁻ (+16%), and Fe (+15%). Conversely, a further warming of root temperature (T25) caused a drastic limitation of NO₃⁻ and SO₄²⁻ acquisition, halving their uptake rates. Similarly, a remarkable reduction was recorded for Fe uptake when the temperature of the nutrient solution was maintained at 25 °C.

The effect of the growth medium temperature was also evaluated in the post-harvest period measuring both the SPAD index and the weight loss of corn salad packed under air and stored up to 25 d at 4 °C. Temperature of the nutrient solution influenced SPAD index during post-harvest storage; bleaching of the leaves was found to proceed slower as the temperature of the nutrient solution was increased (Fig. 5A). The loss of weight from leaf blade dehydration appeared particularly fast during storage of leaves collected from T15 plants (Fig. 5B).

**Discussion**

Among all the environmental parameters, temperature is considered the most able to affect plant growth and some authors (Gavito et al., 2001) have suggested to consider air and soil temperatures in studies of plant growth with the same relevance of soil fertility. Concerning its effects, it is well known that soil temperature can strongly influence plant growth acting indirectly on soil properties (such as nutrient availability) or directly on the biochemical and physiological processes of the whole plant (such as nutrient acquisition and metabolism). Following this, plant yield may be affected considerably by this parameter with, in some cases, consequences on safety such as nitrate content in edible leaf tissues (Lawlor et al., 1987a, 1987b; Miller et al., 2001; Santamaría, 2006). However, when this aspect is studied using soil-grown plants, discriminating the relative contribution between these two components appears very difficult; furthermore, the identification of possible solutions to the problem is not so easy, particularly when addressing the critical aspects of root components. For these reasons, in the present work, the effect of temperature of the growing medium on yield and quality of ready-to-eat corn salad plants has been evaluated using a soilless system, in which the availability of nutrients was maintained at its optimal values; furthermore, *in silico* analyses with a geochemical model excluded any effect of the three values of temperature (15, 20, or 25 °C) on the chemical equilibria and the availability of the dissolved nutrients (data not shown). Then, with this approach, it was possible to evaluate the direct effects of hydroponic solution temperature on yield and quality of the edible tissues taking into account physiological processes related to these aspects such as nutrient acquisition but excluding the effect of soil processes and properties.

Results of this study showed that when corn salad plants were grown in a nutrient solution maintained at the temperature of 20 °C (T20), edible yield (leaf biomass; Table 1) was higher than that observed in plants exposed to 15 °C (T15) for the entire time of the experiment. Furthermore, the greenness of these leaves (T20) was more intense; this is an interesting aspect because consumers mainly judge the acceptability of ready-to-eat vegetables based on their appearance. Similarly, higher yield and more evident greening with temperature rise have also been described.
FeIII or SO4\textsuperscript{2-} uptake media were maintained at the same levels of those used in the growing conditions. Data are means ± SD of three independent experiments; capital letters refer to statistically significant differences among the samples (analysis of variance, Fisher’s least significant difference, \( P < 0.05 \)). FW = fresh weight.

Table 3. Effect of the nutrient solution temperature on NO\textsubscript{3}\textsuperscript{-}, SO\textsubscript{4}\textsuperscript{2-}, and Fe\textsuperscript{III} uptake by roots of corn salad [\textit{Valerianella locusta} (L.) Laterr., cultivar Gala] grown as described in Table 1.\textsuperscript{a}

<table>
<thead>
<tr>
<th>Nutrient solution temp. (\degree\textsuperscript{C})</th>
<th>NO\textsubscript{3}\textsuperscript{-} uptake (\textmu\text{mol} \text{g}^{-1} root DW/h)</th>
<th>SO\textsubscript{4}\textsuperscript{2-} uptake (\textmu\text{mol} \text{g}^{-1} root DW/h)</th>
<th>Fe\textsuperscript{III} uptake (\textmu\text{mol} \text{g}^{-1} root DW/h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>15</td>
<td>173 ± 31 B</td>
<td>70.1 ± 5.4 B</td>
<td>0.89 ± 0.08 B</td>
</tr>
<tr>
<td>20</td>
<td>313 ± 15 A</td>
<td>81.2 ± 2.9 A</td>
<td>1.02 ± 0.06 A</td>
</tr>
<tr>
<td>25</td>
<td>115 ± 23 C</td>
<td>35.4 ± 5.8 C</td>
<td>0.56 ± 0.03 C</td>
</tr>
</tbody>
</table>

\( ^{a} \text{Net nitrate uptake was measured spectrophotometrically as depletion from a solution containing 1 m}\text{M NO}\textsubscript{3}^{-}; \text{Fe}^{\text{III}} \text{or} \text{SO}_4^{\text{2-}} \text{uptake were determined at pH 6.0 by using} \text{^{59}Fe} \text{(Fe-o,oEDDHA, final iron concentration of 100 \textmu M) or} \text{^{35}S(SO}_4^{\text{2-}}, \text{final sulfur concentration of 600 \textmu M L}^{-1} \text{tracers, respectively. Temperatures of the uptake media were maintained at the same levels as those used in the growing conditions. Data are means ±} \text{SD of} \text{three independent experiments; capital letters refer to statistically significant differences among the samples (analysis of variance, Fisher’s least significant difference, } \text{P < 0.05).} \text{DW = dry weight.}

Fig. 4. Effect of the nutrient solution temperature on root biomass of corn salad [\textit{Valerianella locusta} (L.) Laterr., cultivar Gala] grown as described in Figure 3. Data of root biomass (A) and root FW/leaf FW ratios (B) are reported. Data are means ± SD of three independent experiments; capital letters refer to statistically significant differences among the samples (analysis of variance, Fisher’s least significant difference, \( P < 0.05 \)).

Fig. 5. Effect of the nutrient solution temperature on SPAD index values and weight loss (%) during postharvest storage at 4 \degree\textsuperscript{C} of corn salad leaves [\textit{Valerianella locusta} (L.) Laterr., cultivar Gala] grown as described in Figure 3. Data are means ± SD of three independent experiments.

for other soilless production systems (Nxawe et al., 2009), particularly in the early spring period when plant growth is strongly dependent on temperature (Westwood, 1988). Changing temperature of the nutrient solution from 15 to 20 \degree\textsuperscript{C} increased also the nutrient uptake rates of NO\textsubscript{3}\textsuperscript{-}, Fe, and SO\textsubscript{4}\textsuperscript{2-} (Table 2), which in turn might have speeded up metabolic processes and hence shoot growth (Dong et al., 2001; Ndakidemi and Semoka, 2006). Concerning nitrate uptake, its increased rate at T20 is concomitant with a smaller nitrate accumulation at the leaf level (Table 1); although it seems contradictory, it is well demonstrated that under abiotic stress, like nutrient deficiencies, plants accumulate nitrate in leaves even if the uptake rate of the nitrate anion is depressed (Iacuzzo et al., 2011; Nikolic et al., 2007; Prosser et al., 2001). As argued by these authors, the nitrate accumulation is mainly the result of slower growth and a comparatively higher repression of the assimilation process than the uptake one. In fact, when the ratios between N as nitrate and total N accumulated at the leaf level are considered (Fig. 3), it appears evident that an enhanced nitrate assimilation process occurred in plants maintained in a growth medium and air temperature of 20 \degree\textsuperscript{C}. Furthermore, it is interesting to note that the uptake rates of all the three nutrients were enhanced with the temperature rise indicating how an equilibrate growth of plants requires a balanced acquisition of the three elements (Iacuzzo et al., 2011). The observed changes in nutrient uptake capability, as also observed in previous work on ammonium, P, and K (see the review of Bassirirad, 2000), can be at least partially explained by the effect of temperature on root respiration, a temperature-sensitive process that mediates ion movement across the root. However, because ion uptake is not the only root function that requires energy from respiration, despite the tight linkage between root nutrient uptake and respiration, it should not be assumed that increased soil temperature would affect root nutrient uptake as a linear function of root respiration (Bassirirad, 2000). Dong et al. (2001) argued that nutrient uptake and root functions are simultaneously regulated by soil temperature. Furthermore, as reported by Buckley (2008), one general characteristic of plant growth is the strong relationship between structural development and physiological functioning of roots and shoots. For these reasons, the enhanced nutrient fluxes recorded at the root level could influence processes such as photosynthesis at the shoot level (Calatayud et al., 2004, 2008; Santarius, 2004; Yamori et al., 2006, 2008) exerting in turn pronounced effects on shoot growth and consequently on yield (Bowen, 1991). In addition, experimental evidence shows that, when plants are exposed to cold temperature, a following rising of temperature might modify the carbon allocation among roots and shoots (Lambers and Poorter, 1992) favoring the lower levels of biomass accumulation at the root level similar to that observed in this work by increasing the hydromonic solution temperature from 15 to 20 \degree\textsuperscript{C} (Fig. 4). Clarkson et al. (1988) suggested that the response of nutrient uptake to temperature might represent changes in plant nutrient demand per unit root and not an acclimatory change in root transport properties. Nonetheless, with the exception of Fe and Mn, the considerably enhanced biomass production resulting from the warming of the nutrient solution from 15 to 20 \degree\textsuperscript{C} might have caused a dilution of nutrient concentrations of leaf tissues (Table 2). In particular, with respect to nitrate contents, which are an important aspect of food safety (Santamaria, 2006), the temperature rise could also either favor NO\textsubscript{3} assimilation processes or limit the large storage capacity of plant tissues for NO\textsubscript{3} in cells, a phenomenon observed when metabolism is restricted by low temperature (Lawlor, 2002). This aspect is also evident in leaf tissues of T20 plants as compared with those of T15 plants (Fig. 3), in which the ratio of nitrate-N on the total N is significantly lower. In addition to the low nitrate contents, it is interesting to note that the ready-to-eat corn salad from T20 plants showed a lower tendency to bleaching and dehydration during storage (Fig. 5) as compared with that obtained from T15 plants. Based on these
results, it can be inferred that setting the temperature of the nutrient solution at 20 °C could be particularly effective in obtaining fresh-cut products with higher physical stability and hence longer shelf life.

When the temperature of the nutrient solution was raised to 25 °C, levels of edible yield (leaf biomass; Table 1) dropped to the lowest values of those recorded in the three treatments. Furthermore, these plants (T25) developed foliage with fewer leaves and with the lowest greenness intensity. These effects are in agreement with that observed when plants are exposed to high soil temperatures clearly indicating that the temperature of 25 °C for a nutrient solution represents too high a value for balanced and unstressed growth of corn salad plant. In fact, the characteristic limitation by high soil temperature of root growth (Xu and Huang, 2000), water content (Graves et al., 1991; Huang et al., 1991), nutrient uptake (Huang and Xu, 2000), and oxidative damage in leaves (Huang et al., 2001) was also evident in our T25 corn salad plants (Tables 1 and 3) indicating a sort of high-temperature stress for these plants. The leaves of these plants showed slower bleaching and weight loss (Fig. 5) during post-harvest storage in comparison with the values observed in T15 plants. At harvest, the worst leaf quality of leaves in T25 plants (lower greenness and higher dry weight percentage; Table 1) could reasonably contribute to the slowed tendency to bleaching and dehydration of the leaves during storage (Fig. 5). Data reported in Table 1 show that, although root NO3 uptake was more than halved in T25, nitrate contents in theses leaves were slightly increased compared with T20. Lawlor (2002) argued that the rate of protein synthesis, central in N metabolism and plant growth, when all the other factors are not limiting, strongly depends on temperature; it stops at very low temperatures, increasing to a maximum as temperature rises before decreasing with further increase. This affects all plant processes thereby influencing plant growth and changes the demand of nitrate as well as of the other nutrients. Results here reported show a similar pattern for corn salad plants when the temperature of the growing medium has been increased from 15 °C up to 25 °C. In addition, experimental evidence shows that as the requirements for protein synthesis decrease, so amino acids accumulate and the demand for NO3 falls, but not its uptake, so NO3 accumulates (Lawlor et al., 1987a, 1987b, 1988; Miller et al., 2001).

In summary, results of this study showed that, independently from the effect on nutrient availability, the temperature of the growing medium plays an important role in the functionality of root apparatus of corn salad plants, which in turn can affect their growth and food quality and safety. This aspect is particularly relevant for soilless cultures using hydroponics in which, during summer or early spring, the temperature of the nutrient solution could be too different from that required to optimize plant growth. Because the increase or decrease in temperature above or below the optimum level can alter several physiological functions such as photosynthesis, chlorophyll formation and pigmentation, nutrient uptake, synthesis of secondary metabolites in plants (Nxaue et al., 2010), and, as a consequence, the yield and the establishment of the optimum temperatures in the growing medium to meet demands of specific plant species appear very important particularly in hydroponics in which heating or cooling can be easily provided (Calatayud et al., 2008). In the environmental conditions used in the present work for corn salad plants grown in hydroponics with a floating system, maintaining the growing medium temperature at a level close to that of air appears to be the most appropriate to maximize the yield, guaranteeing the safety of the edible tissues and their properties fundamental for storing post-harvest. Moreover, the experimental evidence proves that root temperature levels higher than those of shoot (air temperature) for a long span of time are deleterious for root functionality and plant growth.

Literature Cited


