Seasonal characterization of antioxidant responses in plants of Ipomoea nil cv. Scarlet O’Hara

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Received August 31, 2011 – Accepted December 1, 2011 – Distributed November 30, 2012
(With 2 figures)

Abstract

Reactive oxygen species can be produced in leaf cells during normal aerobic metabolism or in a variety of exogenous factors, which may cause oxidative damage to plants, unless they have an efficient antioxidant defense system, consisting of enzymatic and non-enzymatic substances. This work raised the hypothesis that plants of Ipomoea nil cv. Scarlet O’Hara, a native species and ornamental vine of the tropics, might tolerate oxidative stress factors imposed by natural fluctuations in weather conditions through changes in the antioxidant profile. The objective of this study was to determine the variations in three leaf antioxidants in plants growing inside a greenhouse without air pollutants and exposed to varying meteorological conditions throughout the four seasons of the year and to observe if such variations are related to the oscillations in meteorological factors. Four experimental campaigns were carried out, one in each season of 2006. Each campaign lasted 28 days and started with 45 plants. Ascorbic acid (AA) concentrations and superoxide dismutase (SOD) and peroxidase (POD) activities were determined in leaves of five plants in nine sampling days of each campaign. The antioxidant responses oscillated throughout the year. The highest values were found during the spring. This seasonal antioxidant profile was associated to variations in temperature, relative humidity and global radiation. Plants of this cultivar may then tolerate oxidative stress naturally imposed by meteorological conditions.

Keywords: climber plant, ascorbic acid, superoxide dismutase, peroxidase, meteorological factors.

Caracterização sazonal de respostas antioxidativas em plantas de Ipomoea nil cv. Scarlet O’Hara

Resumo

As espécies reativas de oxigênio podem ser produzidas em células de folhas durante o metabolismo aeróbico normal ou o sob uma diversidade de fatores exógenos que, por sua vez, podem causar danos oxidativos às plantas, a menos que estas tenham um eficiente sistema de defesa antioxidativo, formado por substâncias enzimáticas e não enzimáticas. Neste trabalho, levantou-se a hipótese de que as plantas de Ipomoea nil cv. Scarlet O’Hara, uma espécie trepadeira ornamental e nativa dos trópicos, podem tolerar fatores de estresse oxidativo imposto por oscilações naturais nas condições meteorológicas por meio de mudanças no perfil antioxidativo. Assim, este trabalho objetivou determinar as variações em três espécies antioxidantes foliares em plantas crescidas em casa de vegetação sob ar filtrado e expostas a condições meteorológicas variáveis ao longo das quatro estações do ano de 2006, bem como verificar se tais variações estão relacionadas às oscilações de fatores meteorológicos. Para tanto, realizaram-se quatro campanhas experimentais. Cada campanha durou 28 dias e começou com 45 plantas. Concentrações de ácido ascórbico (AA) e as atividades de superóxido dismutase (SOD) e peroxidase (POD) foram determinadas em folhas de cinco plantas distintas e retiradas da casa de vegetação em nove dias de amostragem de cada campanha. As respostas antioxidativas oscilaram durante todo o ano, sendo os maiores valores encontrados durante a primavera. Este perfil sazonal de antioxidantes foi associado às variações de temperatura, umidade relativa e radiação global. As plantas desta cultivar podem, então, tolerar o estresse oxidativo naturalmente imposto pelas condições meteorológicas.

Palavras-chave: planta trepadeira, ácido ascórbico, superóxido dismutase, peroxidase, fatores meteorológicos.
1. Introduction

Some plants may tolerate oscillations in the abiotic conditions of the environment where they are growing more than others, which may subject them to an oxidative stress induced by increased contents of reactive oxygen species in plant tissues (De Gara et al., 2010; Foyer and Noctor, 2011). This occurs due to the differential efficiency of antioxidants in neutralizing the noxious effects of ROS to the cells. It is well known that antioxidants of the ascorbate-glutathione cycle play an important role in this defensive process, during both normal metabolic activities, in the photoreduction of the O$_2$ into the thylakoid or by exogenous reasons and the entrance of ozone to the plant (Burkey et al., 2006; Iriti and Faoro, 2008; De Gara et al., 2010; Foyer and Noctor, 2011).

Ascorbic acid, a non-enzymatic antioxidant, besides regulating cell division, is one of the most important antioxidants in plants. Hydrogen peroxide (H$_2$O$_2$) is decomposed while this antioxidant is oxidized to dehydroascorbic acid, a reaction that is mediated by ascorbate peroxidase. AA is also considered an important molecule against ozone due to its occurrence in the apoplas, once this pollutant enters the plant predominantly through the stomata and reacts immediately with water intensifying the ROS formation. Other non-enzymatic molecules implicated into the Halliwell Asada cycle may also be relevant to neutralize the ROS. The glutathione, for example, serves as a substrate to remove H$_2$O$_2$ by glutathione peroxidase, converting it to an oxidized form, which prevents lipid peroxidation and reacts with radicals hydroxyl and singlet oxygen. It also protects the SH groups of essential cell metabolism enzymes and has the important role of regenerating the AA. (Foyer and Noctor, 2005, 2011).

As a scavenger of radical superoxide, the enzyme superoxide dismutase (SOD) is another important tool to defend plants against oxidative stress, SOD converts O$_2^-$ to H$_2$O$_2$, and then H$_2$O$_2$ is converted to H$_2$O by ascorbate peroxidase (APX), by means of the oxidation of AA or by catalases (CAT) (Iriti and Faoro, 2008; De Gara et al., 2010; Lin et al., 2011).

The formation and level of such antioxidants may be seasonality marked in response to changes in the environmental conditions, even in the absence of anthropic interferences. This is plausible because the seasonality in solar irradiation, photoperiod, temperature and relative humidity, among other meteorological factors, as well as influencing the stomata aperture, regulates the photosynthesis and respiration processes on chloroplasts and mitochondria and thus the natural production of ROS in the cells. Variations in the levels of antioxidants may also occur during the aging of the plant, which occurs concomitantly with higher protein degradation, and loss of chlorophyll (Bulbovas et al., 2005; Ohe et al., 2005; Foyer and Noctor, 2005; 2011). Therefore, we may raise the hypothesis that variations in antioxidant responses throughout the life cycle occurs, making the plant more or less susceptible to seasonal variations in diverse environmental stressors. In theory, this hypothesis is also valid for the ornamental cultivar Scarlet O’Hara of Ipomoea nil, which is the aim of the present study, due to its climbing habit and high sensitivity to ozone, an interesting characteristic for biomonitoring purposes (Nouchi and Aoki, 1979).

Therefore, the range of antioxidant responses to naturally varying meteorological conditions should be known before its usual application as an ornamental or bioindicator plant in a determined environment.

Thus, the aims of this work were: 1) To determine the variations in three antioxidant species in leaves of Ipomoea nil cv. Scarlet O’Hara, throughout their development in the four seasons of the year; 2) To observe if such variations are related to the oscillations in temperature, relative humidity and solar radiation throughout the seasons.

2. Material and Methods

2.1. Plant cultivation and experimental campaigns

Seeds of Ipomoea nil cv. Scarlet O’Hara were commercially acquired (from CN Seeds LTD, www.cnseeds.co.uk) and derived from the same lot. They were germinated in a mixture of a commercial substrate mainly consisting of bark of Pinus (Plantimax-Eucatex) and fine vermiculite, at a ratio of 3:1, respectively. Seedlings containing the cotyledonal leaves were transplanted to plastic vases with the same substratum mixture. At this moment, a small wooden prop was placed to support the plants during their growth.

Four experimental campaigns were carried out, one in each season of 2006. The summer, autumn, winter and spring campaigns were done in February/March, May/June, August/September and November/December respectively.

Each campaign started when plants had at least seven expanded leaves including the cotyledonal ones, approximately one month after seedling transplantation, and lasted 28 days. It was initiated with 45 plants produced as described above. During the period of 28 days, in the time zero and in intervals between three or four days (totalizing nine sampling days), the concentrations of ascorbic acid (AA) and the activity of superoxide dismutase (SOD) and peroxidase (POD) were determined in the 5th, 6th and 7th older leaves of the main stem of five plants.

During cultivation and experimental periods, the plants had an adequate irrigation ensured by capillarity thorough nylon strings and received periodic fertilization by aqueous nutrient solution prepared according to Epstein (1975).

All the experimental campaigns, from the germination of the seeds to plant sampling, were carried out in a greenhouse supplied by filtered air located at the Institute of Botany. This is in the Southeastern region of the São Paulo city, exactly at 23° 38’ 28.8” S and 46° 37’ 15.8” W; 805 m above sea level (Fernandes et al., 2002).

Air conditioning regulated daily maximum temperatures of air so that it varied in a similar way to that in the external environment. Average values of temperature, humidity and radiation during each campaign indicated that the plants were exposed to an acceptable range of these meteorological parameters and to similar environmental conditions observed in São Paulo (Table 1).
2.2. Analytical procedures

AA was determined in fresh leaves (0.2 g), homogenized with 12 mL of EDTA-Na₂ (0.07%) and oxalic acid (0.5%). The mixture was centrifuged at 40000 g for 30 minutes at 2 °C. An aliquot of the supernatant was added to 2.5 mL of DCPiP (0.02%) and absorbance was measured spectrophotometrically at 520 nm (first lecture). After adding 0.05 mL of ascorbic acid (1%), the second absorbance measurement was estimated. Both absorbance measurements were used to estimate the ascorbic acid content (Keller and Schwager, 1977).

SOD activity was also determined in frozen leaves at -40 °C (0.4 g), homogenized with 12 mL of potassium phosphate buffer (50 mM pH 7.5) with EDTA-Na₂ 1 mM, NaCl 50 mM and ascorbic acid 1 mM in the presence of a pinch of polyvinyl polypyrrolidone (PVPP) 2% and centrifuged at 22000 g for 25 minutes at 2 °C. The activity of SOD was assayed by measuring the SOD inhibition of the NBT photochemical reduction (Osswald et al., 1992). Each reaction mixture contained 0.5 mL of EDTA-Na₂, 0.54 mM, 0.8 mL of potassium phosphate buffer (0.1 M, pH 7.0), 0.5 mL of methionine 0.13 mM, 0.5 mL of NBT 0.44 mM, 0.2 mL of riboflavin 1 mM and 0.2 mL of leaf extract. The samples were incubated for 20 minutes under a fluorescent lamp (80 W). The absorbance of the reaction mixture was determined at 560 nm. A similar mixture lacking the leaf extract was used as a control, and a dark control mixture served as a blank. The enzymatic activity was expressed as the amount of extractable needed to inhibit the reduction of NBT by 50%.

POD activity was determined in frozen leaves at -40 °C (0.3 g), homogenized with 12 mL potassium phosphate buffer (0.1 M, pH 7.0) in the presence of a pinch of polyvinyl polypyrrolidone (PVPP) 2%. The homogenate was centrifuged at 40000 g for 30 minutes at 2 °C. The Peroxidases were determined in a reaction mixture of plant extracts, 0.1 M potassium phosphate buffer (pH 5.5) and fenylendiamine (1%), to which was added an aliquot of H₂O₂ (0.3%), according to Klumpp et al. (1989). Unspecific POD activity was measured spectrophotometrically following the increase in absorbance (dE) at 485 nm due to the formation of an H₂O₂-POD complex at two different times in the linear reaction curve.

2.3. Statistical analyses

Differences in antioxidant responses among leaves in each sampling day were searched by means of one way analyses of variance. Two way analyses of variance with two factors were carried through in order to identify differences between the seasons (factor 1) and throughout the time in each season (factor 2). In all the cases, the post-hoc multiple comparison test (Student-Newman Keuls) was applied if the analysis of variance indicated significant differences. When necessary, the data were transformed to reach the normal distribution and/or equal variances. Principal component analysis (PCA) was carried out in order to evaluate the overall variability of antioxidant responses in leaves of I. nil cv. Scarlet O’Hara throughout the four seasons of the year, searching for evidence of seasonality. Analyses of correlation (Pearson) were carried out to determine the relations and antioxidant responses in the leaves and meteorological conditions inside the greenhouse in each experimental campaign and between the antioxidant responses in plants analyzed during the overall experimental period.

3. Results and Discussion

In most of the cases, no significant differences in the antioxidant responses were found among the leaves analyzed in each sampling day during each experimental campaign, which might characteristically indicate a leaf aging effect (data not shown). This result opposes the expectation that younger leaves should present more efficient capacity of defense than older leaves in the same plant. Moreover, the young leaf generally presents a high metabolic rhythm due to its stage of development, which should intensify the formation of reactive oxygen species and demands a higher efficiency of the antioxidative system. Ohe et al. (2005), for example, observed that older leaves of Nicotiana tabacum cv. Xanthi exposed to conditions of phototoxic stress presented a lower content of AA and lesser activity of ascorbate-peroxidase in the chloroplasts when compared with younger leaves.

As a consequence of the absence of differential responses among leaves, the antioxidants measured in each sampling day were presented in Figure 1 and Table 2 as average per plant. Such responses oscillated in their levels throughout the year. The contents of AA in the plants of I. nil cv. Scarlet O’Hara exposed during the summer campaign gradually increased over time, reaching maximum values after seven days of experiments and then gradually decreased. The leaf AA concentration remained almost constant during plant growth in the autumn campaign. The winter campaign was characterized by a significant high concentration of the AA in plants taken on the last day of sampling. The leaf AA varied throughout all 28 days of the experiment performed during the spring, reaching maximum values after 11 to 14 days (Figure 1a). On average, no significant variations in the leaf levels of AA were observed among the seasons (Table 2).
The activity of SOD in plants of *I. nil* cv. Scarlet O’Hara oscillated during the summer campaign. It was significantly higher after 04, 21 and 28 days of this campaign. During the autumn, the activity of SOD was relatively constant in the plants. A gradual decrease in its levels occurred from the 21st day on, reaching the lowest value in the last day of analysis. Similar to AA, the activity of SOD was significantly low in plants sampled in the middle of the winter campaign and reached the maximum average value in plants sampled in the last day of sampling. The activity of SOD gradually increased over time during the spring campaign, reaching maximum values in the three last days of the sampling (Figure 1b). On average, significantly higher value of SOD was observed in the plants analyzed during the spring campaign (Table 2).

The activity of POD in plants of *I. nil* cv. Scarlet O’Hara did not significantly differ in plants sampled during the autumn campaign. During the winter campaign, significantly low activity of POD was measured in the plants taken after four days of exposure and high after 18 days of experiment. POD showed a clear oscillatory profile during the spring campaign. Its activity gradually increased during the first 11 days and then decreased until the 21st day. Similar oscillation was also observed at the end of this campaign (Figure 1c). A significantly higher average value of POD was also observed in the plants analyzed during the spring campaign (Table 2).

The principal component analysis (PCA) carried out with the biological measurements (content of AA and activities of SOD and POD) explained 93.7% of the variability of the data in axis one and two (55.9% in axis one and 37.8% in axis two). The level of both enzymatic antioxidants (SOD and POD) was strongly related with axle one; contrarily AA was weakly related with it. The graph from this analysis confirmed the occurrence of an evident seasonality in the antioxidant responses in plants of *I. nil* cv. Scarlet O’Hara, mainly marked by the enzymatic antioxidants (Figure 2). The sampling units of the spring campaign were grouped on the positive side of axle 1, and were characterized by the highest values of SOD and POD. On the other hand, the sampling units of the other seasonal experiments were generally grouped at the opposite side of this axle. Plants generally presented a low activity of SOD, but higher levels of POD during the autumn and winter and a low activity of both SOD and POD during the summer.

**Figure 1.** Average profiles of antioxidants in plants of *I. nil* Scarlet O’Hara, in nine days of analysis, throughout 28 days of each experimental campaign carried out in 2006. a) Concentrations of ascorbic acid (AA). b) Activity of superoxide dismutase (SOD); c) Activity of peroxidases (POD). Distinct letters indicate significant differences in antioxidant responses among the sampling days in each season. # Data not available.

**Table 2.** Average values of ascorbic acid and activity of superoxide dismutase and peroxidases in plants of *I. nil* ‘Scarlet O’Hara, throughout 28 days of each experimental campaign carried out in 2006.

<table>
<thead>
<tr>
<th>Campaign</th>
<th>AA (µg/g dw)</th>
<th>SOD (Unid/g dw)</th>
<th>POD (dE/min/g dw)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Summer</td>
<td>7.9&lt;sup&gt;a&lt;/sup&gt;</td>
<td>332.6&lt;sup&gt;b&lt;/sup&gt;</td>
<td>#</td>
</tr>
<tr>
<td>Autumn</td>
<td>6.3&lt;sup&gt;a&lt;/sup&gt;</td>
<td>370.5&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1008.4&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Winter</td>
<td>6.9&lt;sup&gt;a&lt;/sup&gt;</td>
<td>384.0&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1300.2&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Spring</td>
<td>8.6&lt;sup&gt;a&lt;/sup&gt;</td>
<td>619.8&lt;sup&gt;b&lt;/sup&gt;</td>
<td>2153.3&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

Distinct letters indicate differences in antioxidant responses among the different seasons of the year.
Finally, the analyses of the Pearson correlation indicated that the variations in antioxidants were more strongly stimulated by daily oscillations in climatic factors such as temperature, humidity and radiation five days before plant sampling throughout the different seasons of the year (Table 3), showing that the relation between the changes in the environment and the plant antioxidant responses is time dependent. These correlations showed to be the clearest among those tested between leaf antioxidants and environmental conditions zero to ten days before the leaf sampling (data not shown). Similar results were obtained by Dafré-Martinelli et al. (2011).

No significant relations were found between the antioxidants and meteorological conditions during the summer campaign. Air temperature positively influenced the content of AA during the autumn and winter and the activity of both enzymes during spring. Relative humidity (RH) positively influenced the levels of AA in the autumn, the activity of SOD and POD in the winter and SOD again in the spring. A significant and negative relation was observed between RH and AA in the spring period. Solar radiation was negatively related to the concentrations of AA during the autumn and winter and positively related to AA and SOD in the spring. In brief, the differential intensity of the meteorological conditions seemed to affect the prooxidant/antioxidant equilibrium in the cells of *I. nil* cv. Scarlet O’Hara, as pointed out by Muggli (1993). Bulbovas et al. (2005) also observed no uniform relationship between antioxidant responses in...
plants of *Caesalpina echinata* Lam. cultivated in the same greenhouse used in the present study and oscillations in air temperature and relative humidity, which in turn regulate the stomata movement. The authors could similarly attribute the varying antioxidant responses to greater or minor rates of plant photosynthesis, transpiration and respiration.

The highest values of antioxidants found in the spring campaign performed with plants of *I. nil* may indicate two antagonist situations. They could have reflected ideal environmental conditions for growth of *I. nil* cv. Scarlet O’Hara. If so, the meteorological conditions observed inside the greenhouse during that period promoted high rates of photosynthesis in the plants. Consequently the formation of EAO should have been intensified, demanding increased antioxidant responses. In fact, this positive influence is expected according to Foyer and Noctor (2005, 2011), who affirmed that these seasonal oxidation-reduction relations may be part of how plants perceive and respond to environmental triggers.

However, the highest levels of antioxidants in plants of *I. nil* grown during the spring may also have reflected environmental stressing conditions instead of ideal conditions for growth. According to Takahashi and Badger (2011), extreme amounts of radiation and an increase of radiation absorption of UV create a situation of stress that may firstly affect the photosystem II reaction center. This fact may have occurred in the present study during spring, considering that it was characterized by the highest values of solar radiation and temperature inside the greenhouse in comparison to the meteorological conditions observed in the other seasons and that positive relationships were preponderantly found between the level of the antioxidant responses and solar radiation or temperature during this season. Additionally, Foyer and Noctor (2011) affirmed in their review that ascorbic acid fulfills several important roles in the protection of photosynthesis from the adverse effects of high sunlight. These authors add that the abundance of ascorbate in leaves is regulated both in response to the amount of light available during the photoperiod and the red/far-red ratio of the incident light. The strong positive correlation proved between the concentrations of ascorbic acid and global radiation observed in the present study during the spring points particularly to this kind of protection of the *I. nil* against the solar radiation in excess.

Other authors also observed such seasonality in antioxidants in response to the meteorological characteristics of each season of the year. Gilham and Dodge (1987) showed that the ascorbate levels, ascorbate-peroxidase and glutathione-reductase in leaves of *Pisum sativum* (L.) under control conditions presented an outstanding seasonal variation. Low activities and concentrations of antioxidants marked respectively the winter and summer times. The authors associated these results to differences in the density of light flow. During the hottest seasons of the year, the respiratory rates of the plants are higher and can also induce an increasing antioxidant response due to a consequent higher ROS production in mitochondria. This respiratory increase is associated with an increase in the NADH synthesis (Dizengremel et al., 2009), which is related to the production of enzymes such as SOD. For example, Zhang et al. (2008) found a significant copper-zinc SOD increase in *Elsholtzia haichowensis* during the enhancement of respiration rates induced by some stress factors. Therefore, plants have evolved various mechanisms to cope with the stresses imposed by naturally fluctuating environmental conditions, generally modulated by the gene expression and synthesis of compounds that may result in higher stress tolerance (Ahmad et al., 2010). Willekens et al. (1994) showed that ozone promoted an increase in the mRNA levels, resulting in an enhancement of the synthesis of glutathione peroxidase and some isoenzymes of CAT in plants of *N. plumbaginifolia*. Kwon et al. (2002) demonstrated that the simultaneous expression of Cu/Zn SOD and APX genes in tobacco chloroplasts enhanced tolerance to oxidative stress compared to the expression of either of these genes alone. Gillespie et al. (2011) also observed significant changes in the transcription of specific proteins and antioxidant responses in plants of *Glycine max* (soybean) exposed to acute doses of ozone.

The antioxidant profile observed in plants of *I. nil* cv. Scarlet O’Hara during the summer campaign seemed to mark more characteristically the rhythm of growth and aging of the whole plant than the influence of daily meteorological conditions, as commented by Musselman and Massman (1998), Ohe et al. (2005) and Foyer and Noctor (2005, 2011). Maximum leaf AA concentrations and SOD activity after seven days of experiments might have indicated maximum cell division and growth rate of plants, followed by an evident aging process.

Analyses of Pearson correlation finally showed that the antioxidant responses of *I. nil* ‘Scarlet O’Hara’ were themselves positively related throughout the four campaigns [AA × SOD (*r* = 0.37, *p* = 0.03); AA × POD (*r* = 0.44, *p* = 0.02) and SOD × POD (*r* = 0.44, *p* = 0.02)]. This fact suggests that integrated plant responses to ambient variations occurred reflecting the good capacity against oxidative stress, as found by Foyer et al. (1997). It means that the high activity of SOD led to an increase in the H$_2$O$_2$ production that was, in turn, eliminated by other antioxidative substances of the cycle ascorbate-glutathione as AA and peroxidases (more specifically ascorbate-peroxidase).

The results from this study revealed an evident seasonality in the antioxidant responses of plants of *Ipomoea nil* ‘Scarlet O’Hara’ growing throughout the four seasons of the year. These oscillating responses could be associated to a gradient of environmental conditions imposed by temperature, relative humidity and solar radiation. The action of determined ambient stimulations, such as that, can demand greater or minor efficiency of the antioxidative system.

Therefore, plants of this cultivar, growing in the greenhouse under filtered air and submitted to similar ambient conditions of São Paulo, are capable of defending themselves against oxidative stress naturally imposed by the climate.

Acknowledgements – We would like to thank FAPESP (Fundação de Amparo à Pesquisa no Estado de São Paulo) for giving financial support and to CNPq (Conselho Nacional de Pesquisa e Desenvolvimento) for offering a MSc scholarship to the first author.
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