Chlorophyll a fluorescence as indicative of the salt stress on *Brassica napus* L.

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**ABSTRACT**

The effects of salinity stress on chlorophyll fluorescence and the growth of *Brassica napus* L were investigated. The chlorophyll a fluorescence transient were recorded and analyzed according to the JIP-test which can quantify PSII performance. Salt stress resulted in decreased leaf area and dry matter compared with the control treatment (0 mM NaCl). The most pronounced effects of salt stress were observed with 200 mM NaCl, and the hybrids displayed different levels of sensitivity to stress. The Performance Index (PIABS) was the most sensitive parameter to salt stress, which suggests that this parameter can be used to screen genotypes for salt tolerance.

**Keywords:** Canola; growth; JIP-Test, salt stress; spring rapessed oil.

**Abbreviations:** ABS, absorption of light energy; Chl, chlorophyll; ET, conversion of excitation energy to electron transport (ET); ET/ABS, electron transport activity; F0, Fv, and FM – minimal, variable and maximum Chl fluorescence of PSII in the dark adapted state; Fv/Fm = φp = TR/ABS, maximum efficiency of PSII photochemistry; Fv′, Fv′ and Fm′ – minimal, variable and maximum Chl fluorescence in the light adapted state; Fv/Fm′, efficiency of excitation capture by open PSII reaction centers; FW, fresh weight; PIABS, Performance Index; PSII, photosystem II; QA, electron acceptor of PSII; qP, photochemical quenching coefficient; RC/ABS, ratio of reaction centers and the absorbance; TR, trapping of excitation energy; φPSII, actual PSII efficiency; φET = ET/ABS, quantum yield of electron transport; φD, quantum yield of dissipation; Ψo = ET/TR, yield of electron transport per trapped exciton; PAR, photosynthetic active radiation.

**INTRODUCTION**

Spring rapessed oil (canola) is an important crop in many countries. Currently, the dominant species is *Brassica napus* L. The seed typically has an oil content ranging from 35 to 45% and a minimum protein content of 35%. Canola is a form of rapseed that has less than 2% erucic acid in its oil and less than 30 μmoles of total glucosinolates in its seed meal (Kris-Etherton et al., 2000).

Stress factors, such as high or low temperatures, drought, high light and salinity can alter many physiological processes and suppress the photosynthetic activity of plants. Soil salinity is a major constraint limiting agriculture productivity (Zhu, 2001). Salt stress has been reported to cause an inhibition of growth and development and reductions in photosynthesis, respiration and protein synthesis in sensitive species (Panda et al., 2006). Salt ion toxicity causes numerous deleterious effects in plants by promoting the denaturation of cytosolic...
enzymes and facilitating the formation of reactive oxygen species that can damage membranes and proteins (Zhu, 2001). The decrease in photosynthetic activity frequently observed under salt stress may be due to limitations in photosynthetic electron transport and partial stomatal closure (Flexas et al., 2004). Although the effects of salt stress on photosynthesis have been studied intensively, the mechanisms of inhibition of photosynthesis by salt stress remain unclear (Munns and Tester, 2008). Many studies have investigated the effect of salt stress in different plant species and cultivars, including maize (Azevedo Neto et al., 2004), dwarf-cashew (Abreu et al., 2008), cowpea (Maia et al., 2010), faba bean (Tavakkoli et al., 2010), Silver buffaloberry (Qin et al., 2010), among others. But the effects on chlorophyll fluorescence parameters in plants are not clear. In both salt-sensitive and salt-tolerant of Chickpea seedling the salt stress affect Fv/Fm, ϕPSII, qP, F0, and Fm (Eyidogan and Tufan, 2007). However, others authors reported that salinity does not affect chlorophyll fluorescence (Redondo-Gómez et al., 2007). Measurement of chlorophyll a fluorescence, a non-invasive method, is widely used for monitoring and screening different species and genotypes for stress resistance or tolerance (Percival et al., 2003). The objective of the present study was evaluated the effect of salt stress in five hybrids of canola by determination the growth of the plant and the fluorescence of chlorophyll a.

**MATERIALS AND METHODS**

**Growth conditions and treatments:** The seeds of *Brassica napus L.* var. *oleifera* (hybrids Hyola 43, Hyola 61, Hyola 401, Hyola 420 and Hyola 432 from Celena Alimentos S/A – some characteristics were presented in Table 1) were germinated directly in sand, and the plants were grown in 5-liter pots. Plants were grown in a greenhouse during September-January in 2008-2009. The environmental conditions during the growth were: air temperature 25–30°C during the day and 15°C roughly in the night. The PAR at 9 a.m. was 450-500 µmol m² s⁻¹. Immediately after sowing, all pots were randomly divided into four groups, and we started the application of the salt solution; one group was grown in the nutrient solution Hoagland (Hoagland and Arnon, 1950) as the non-salt stressed (control), while the others were grown in complete nutrient solution plus 50, 100 and 200 mM NaCl as the salt stress treatments.

**Table 1. The most important characteristics of the hybrids used in experiment**

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Hyola 43</th>
<th>Hyola 61</th>
<th>Hyola 401</th>
<th>Hyola 420</th>
<th>Hyola 432</th>
</tr>
</thead>
<tbody>
<tr>
<td>Emergence to beginning of flowering (days)</td>
<td>53 to 77</td>
<td>60 to 77</td>
<td>44 to 74</td>
<td>64 to 70</td>
<td>47 to 73</td>
</tr>
<tr>
<td>Plant cycle – number of days from plant emergence to harvest</td>
<td>123 to 155</td>
<td>127 to 139</td>
<td>107 to 135</td>
<td>116 to 150</td>
<td>119 to 134</td>
</tr>
<tr>
<td>Plant height (cm)</td>
<td>78 to 129</td>
<td>105 to 134</td>
<td>86 to 126</td>
<td>116 to 130</td>
<td>89 to 124</td>
</tr>
</tbody>
</table>

**Chlorophyll content, leaf area and dry weight:** Chlorophyll content was measured 48 DAS, and we used a chlorophyll meter (CL-01, Hansatech Instruments, Kings Lynn, UK) to measure the middle region of the youngest mature leaf. The values were expressed as the “chlorophyll index” (Cassol et al., 2008). Leaf area, using a area meter (LI-3100, Licor, Lincoln, NE 68504 USA), and total dry weight were measured 48 DAS. The seeds yield was estimated when they had uniformly matured (100 DAS), and express as fresh weight per plant.

**Polyphasic Chl a fluorescence transients:** Polyphasic Chl a fluorescence transients were measured with a direct fluorometer (Handy PEA, Hansatech Instruments, Kings Lynn, UK) at 48 DAS. All fluorescence measurements were taken in the middle part of the plant, in fully expanded leaves, with 10 replicates for each treatment (salinity/hybrids). Leaf samples were illuminated with continuous red light (650 nm peak wavelength) after 30 min in dark-adaptation. All measurements were taken using a saturating pulse of 3.000 µmol m⁻² s⁻¹ for 1 s. Upon irradiation, dark-adapted photosynthetic samples exhibit a fast fluorescence rise from the initial fluorescence intensity F₀ to a maximal intensity Fm (Lázár, 2006). The fluorescence intensity at 50 µs was considered to be F₁, initial (minimal) fluorescence. The JIP test (Strasser and Strasser, 1995; Strasser and Tsimilli-Michael, 2000; Strasser et al., 2000, 2004) was used to analyze each Chl fluorescence transient. The Performance Index (PIₜ₆₈₅) was one of the Chl fluorescence parameters that provided useful and quantitative information about the state of plants and their vitality (Strasser et al., 2000).

The PIₜ₆₈₅ is a combined measure of three partial performances, namely those related to the amount of
photosynthetic reaction centers (RC/ABS), the maximal energy flux that reaches the PSII reaction center (TR₀), and the electron transport at the onset of illumination (ET₀).

\[ P_I_{ABS} = \frac{RC_{ABS}}{ABS} \cdot \frac{\phi_{Po} \cdot \psi_o}{1 - \phi_{Po} \cdot \psi_o} = \frac{RC}{ABS} \cdot \frac{TR_0}{DI_0} \cdot \frac{ET_0}{TR_0 - ET_0} \]

where RC/ABS is the ratio of reaction centers and the absorbance (the concentration of reaction centers per chlorophyll), \( \phi_{Po}/(1 - \phi_{Po}) \) is an expression related to primary photochemistry and \( \psi_o/(1 - \psi_o) \) is an expression related to electron transport (Oukarroum et al., 2007).

Modulated Chl a fluorescence: Modulated Chl a fluorescence was measured 48 DAS by a fluorometer (FMS2, Hansatech Instruments, Kings Lynn, UK), and the fluorescence parameters were determined in light-adapted leaves. Calculations were made from the following: (1) the photochemical quenching coefficient \( [\rho P = (F_m - F_o)/(F_m - F'_0)] \), (2) the efficiency of excitation capture by open PSII reaction centers \( (F_v/F'_v) \), and (3) the actual PSII efficiency \( (\phi_{PSII}) \). Fluorescence nomenclature was used according to van Kooten and Snel (1990). The pulse amplitude modulation (PAM) fluorescence technique obtains both qualitative and quantitative information on the organization and functioning of a plant photosynthetic apparatus by analysis of fast and slow Chl fluorescence induction kinetics.

Statistical analysis: Statistical analysis was conducted with the software package Statistica for Windows Version 6 (StatSoft Inc., 2300 East 14th Street, Tulsa, OK, USA). The data were subjected to an analysis of variance (ANOVA). The differences in mean values between canola hybrids and salt treatment were compared by a Student’s t-test.

RESULTS

Effects of salt stress on Chl content and growth:
The hybrid type was the only factor that resulted in significant differences in Chl content. Hyola 401 and 43 were the hybrids with the highest and lowest Chl contents, respectively. However, there were no statistically significant differences in the Chl content caused by the salinity treatment stress within a given hybrid line (Figure 1A). The Chl content predominantly increased in Hyola 43 at 200 mM NaCl, whereas the Chl content declined in Hyola 401 compared to control plants. However, these differences were not significant. The impact of salt stress on the leaf area (Figure 1B) also differed between hybrids. In Hyola 43 the leaf area declined at 50 mM, but the decrease was not significant. However, Hyola 61 had a significant decline at 200 mM compared to 50 and 100 mM. In Hyola 401 there was no observed difference after salt treatment. In Hyola 420 there was a 47% reduction in leaf area in 200 mM NaCl, and in Hyola 432 no difference was observed. In Hyola 420 there was a 47% reduction in leaf area in 200 mM NaCl, and in Hyola 432 no difference was observed. Plant dry weight was reduced with increased NaCl concentration in all hybrids, but the reductions were more pronounced at 200 mM (Figure 1C), especially in Hyola 61 (65%), Hyola 420 (77%) and Hyola 432 (63%). When we compared the seed yield (Figure 1D) for hybrids without salinity (control plants), we observed that the hybrid Hyola 432 had the lowest yield (0.44 g FW seed per plant), and Hyola 401 had the highest yield (1.47 g FW seed per plant). The seed yields decreased as the salinity level increased in all hybrids, except Hyola 432, which showed increases in seed yields at 50 and 100 mM NaCl. The highest reductions in comparison with the control plants were observed in Hyola 43 (87%) and Hyola 61 (73%) at a salinity of 200 mM NaCl.
**Effect of salt stress on Chl fluorescence:** When plotted on a logarithmic time scale, the kinetics of fluorescence rise exhibited similar profiles, with distinct O-J-I-P steps in all cases. The OJIP transient represents the successive reduction of the electron acceptor pools of PSII (Strasser et al., 2000). Phase J reflects an accumulation of Q_{A}^{−} Q_{B} (Lazar, 2006). Phase I reflects an accumulation of Q_{A}^{−} Q_{B}^{−} Q_{B}^{−} (Lazar, 2006). In Figure 2 we present the curves of chlorophyll fluorescence measured at 200 mM of NaCl and control for different hybrids after 48 days of treatment, where we observed the typical curve OJIP.

The maximal efficiency of PSII photochemistry (φ_{Po}), quantum yield of electron transport (φ_{Eo}), quantum yield of dissipation of energy (φ_{Do}) and yield of electron transport per trapped exciton (Ψ_{o}) were altered by hybrids and salinity, but there were no significant differences in their interaction. The lowest values of φ_{Po}, Ψ_{o}, and φ_{Do} were observed only at 200 mM for all hybrids, although φ_{Do} increased at this salt concentration.
The effect of salinity stress on some JIP-test parameters for each hybrid is shown in Figure 3. There was no large difference for hybrid and level NaCl to specific fluxes towards the RC: a) the ABS/RC – the absorption flux per RC, which indicates the apparent size of the antenna system of PSII, e.g. completely active quantity of absorbing...
chlorophyll per RC; b) TR\textsubscript{T}/RC – the flux of trapping energy per RC in \( t = 0 \); c) ET\textsubscript{T}/RC - flux of electron transport from \( Q_A \) to PQ per RC in \( t = 0 \). Only we observed that DI\textsubscript{T}/RC ratio (flux of dissipated energy per RC in \( t = 0 \)) increased at 200 mM for Hyola 401 (Figure 3).

**Figure 3.** Derivation of the JIP-test parameters from Chl fluorescence induction curves for different canola hybrids (a) Hyola 43, (b) Hyola 61, (c) Hyola 401, (d) Hyola 420, (e) Hyola 432, subject to salt treatment relative to control (0 mM).

Using \( \text{PI}_{\text{ABS-rel}} \) \( \left[ \text{PI}_{\text{ABS-stress}/\text{PI}_{\text{ABS-control}}} \right] \) as a measure of plant performance, differences in the response to salinity between the studied hybrids were revealed. The Performance Index relative \( \left( \text{PI}_{\text{ABS-rel}} \right) \) decreased in Hyola 61 (8%), Hyola 401 (27%) and Hyola 420 (15%), principally at 200 mM. The decline in \( \text{PI}_{\text{ABS-rel}} \) observed in Hyola 401 and Hyola 420 was coupled with reductions in RC/ABS, \( \psi_0/(1 - \psi_0) \) and \( \psi/(1 - \psi) \) and increases in DI\textsubscript{T}/RC. The \( \text{PI}_{\text{ABS-rel}} \) increased in Hyola 43 at 100 and 200 mM and in Hyola 432 at 50 and 100 mM.

The Figure 4 shows the correlation of the log relative Performance Index [\log \text{PI}_{\text{ABS-rel}}]: (A) log of the relative reaction centers per chlorophyll [\log (RC/ABS)\textsubscript{rel}], (B) log of the relative efficiency of primary photochemical [\log (\phi_{\text{P0}}/1 - \phi_{\text{P0}})\textsubscript{rel}], (C) log of the relative efficiency of the conversion
of excitation energy to electron transport \[ \log \left( \Psi_o / (1 - \Psi_o) \right) \text{rel} \] and (D) log of the relative electron transport activity \[ \log (\phi_{Eo}) \text{rel} \]. All log functions were linear with correlation coefficients of higher than 0.86; however the relationship between \[ \log (P_{ABS}) \text{rel} \] and \[ \log (\phi_{Eo}) \text{rel} \] had the highest correlation coefficient (0.948).

**Figure 4.** Correlation for all treatments between \[ \log (P_{ABS}) \text{rel} \] and (a) \[ \log (R_{C/ABS}) \text{rel} \], (b) \[ \log (\phi_{Po} / (1 - \phi_{Po})) \text{rel} \], (c) \[ \log (\Psi_o / (1 - \Psi_o)) \text{rel} \], and (d) \[ \log (\phi_{Eo}) \text{rel} \]; \( r \) = Pearson correlation coefficient.

**Table 2.** The efficiency of excitation capture by open PSII reaction centers \( F_v / F_m ' \), the actual PSII efficiency \( \phi_{PSII} \) and photochemical extinction coefficient \( qP \) in different hybrids of canola cultivated with salt treatments.

<table>
<thead>
<tr>
<th>Hybrids</th>
<th>NaCl (mM)</th>
<th>0</th>
<th>50</th>
<th>100</th>
<th>200</th>
<th>Means</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>0.743* ( b^* )</td>
<td>0.746</td>
<td>0.780</td>
<td>0.772</td>
<td>0.760 ( b \</td>
</tr>
<tr>
<td>Hyola 43</td>
<td></td>
<td>0.711 ( b )</td>
<td>0.780</td>
<td>0.764</td>
<td>0.763 ( b )</td>
<td>0.763 ( b )</td>
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<tr>
<td>Hyola 61</td>
<td></td>
<td>0.701 ( b )</td>
<td>0.758</td>
<td>0.758</td>
<td>0.786</td>
<td>0.750 ( b )</td>
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<tr>
<td>Hyola 401</td>
<td></td>
<td>0.722 ( b )</td>
<td>0.691</td>
<td>0.769</td>
<td>0.756</td>
<td>0.756 ( b )</td>
</tr>
<tr>
<td>Hyola 420</td>
<td></td>
<td>0.700 ( b )</td>
<td>0.799</td>
<td>0.802</td>
<td>0.794</td>
<td>0.793 ( a )</td>
</tr>
<tr>
<td>Mean</td>
<td></td>
<td>0.743 ( b)</td>
<td>0.746 ( b )</td>
<td>0.777 ( a )</td>
<td>0.774 ( a )</td>
<td>0.774 ( a )</td>
</tr>
<tr>
<td></td>
<td>0.684 ( b )</td>
<td>0.713</td>
<td>0.754</td>
<td>0.744</td>
<td>0.724</td>
<td></td>
</tr>
<tr>
<td>Hyola 43</td>
<td></td>
<td>0.729 ( b )</td>
<td>0.711</td>
<td>0.732</td>
<td>0.714</td>
<td>0.714 ( b )</td>
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<tr>
<td>Hyola 61</td>
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<td>0.638 ( b )</td>
<td>0.692</td>
<td>0.729</td>
<td>0.757</td>
<td>0.757 ( b )</td>
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<tr>
<td>Hyola 401</td>
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<td>0.672 ( b )</td>
<td>0.621</td>
<td>0.748</td>
<td>0.730</td>
<td>0.730 ( b )</td>
</tr>
<tr>
<td>Hyola 420</td>
<td></td>
<td>0.754 ( b )</td>
<td>0.776</td>
<td>0.776</td>
<td>0.769</td>
<td>0.769</td>
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<tr>
<td>Mean</td>
<td></td>
<td>0.695 ( b )</td>
<td>0.702 ( b )</td>
<td>0.747 ( a )</td>
<td>0.742 ( a )</td>
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<td>0.913 ( b )</td>
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<td>Hyola 43</td>
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<td>0.946 ( b )</td>
<td>0.957</td>
<td>0.939</td>
<td>0.932</td>
<td>0.932 ( b )</td>
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<tr>
<td>Hyola 61</td>
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<td>0.901 ( b )</td>
<td>0.911</td>
<td>0.961</td>
<td>0.962</td>
<td>0.962 ( b )</td>
</tr>
<tr>
<td>Hyola 401</td>
<td></td>
<td>0.929 ( b )</td>
<td>0.898</td>
<td>0.973</td>
<td>0.966</td>
<td>0.966 ( b )</td>
</tr>
<tr>
<td>Hyola 420</td>
<td></td>
<td>0.967 ( b )</td>
<td>0.971</td>
<td>0.968</td>
<td>0.968</td>
<td>0.968</td>
</tr>
<tr>
<td>Mean</td>
<td></td>
<td>0.931</td>
<td>0.938</td>
<td>0.961</td>
<td>0.961</td>
<td>0.958</td>
</tr>
</tbody>
</table>

*means of ten measurements;

**Different letters indicate significant differences among mean values within each Hybrid (Student t-test, \( p \leq 0.05 \)).
The effect of salt stress treatments on the F$_{V'}/$F$_{M'}$, $\phi_{PSII}$ and qP in fully expanded leaves is shown in Table 2. The Hyola 432 hybrid showed the highest values for the F$_{V'}/$F$_{M'}$ ratio and $\phi_{PSII}$, and Hyola 420 had the lowest F$_{V'}/$F$_{M'}$ ratio. The highest values of F$_{V'}/$F$_{M'}$ and $\phi_{PSII}$ were observed with 100 and 200 mM NaCl. There were no significant differences in the quenching photochemical (qP).

DISCUSSION

The leaf area, total dry matter and seed yields were affected by salinity stress in canola hybrids, principally at 200 mM; however there were differences in sensitivity between the hybrids (Figure 1). The hybrids Hyola 43 and Hyola 432 were less affected by salinity. Studies on the effects of salinity on carbon assimilation in plants could enable a more comprehensive understanding of their productivity and growth (Chaves et al., 2009). However, the studies show that photosynthetic responses to salinity are highly complex and have some limitations when measuring at different sites on the cell or leaf and at different developmental stages. It is well established that the main effect of salt stress is to decrease photosynthesis, which reduces CO$_2$ fixation rates and plant growth. Maricle et al. (2007) concluded that salinity decreased CO$_2$ fixation, but the processes responsible for harvesting solar energy are largely unaffected by increasing salinity in estuarine grasses. While the use of this energy changes with increasing irradiance and salinity, they suggest that there is no strong relationship between fluorescence parameters and salt sensitivity in this species. Our results show that different B. napus genotypes have different levels of susceptibility to salt stress and that the photochemical process can be altered by high salt.

In the present study, we did not find a simple relationship between the growth and fluorescence parameters because the growth was affected by all salt levels, and the fluorescence parameters were only altered in high salinity. Our results showed a slight difference in the maximum yield of primary photochemistry of PSII ($\phi_{PSII}$) observed only at 200 mM for all hybrids. Similar results were obtained in the yield of electron transport per trapped exciton (\$\psi_E\$) and for the quantum yield of electron transport ($\phi_{et}$) as well as an increase for the quantum yield of dissipation ($\phi_{d}$). The salinity showed little effect in the light-adapted state fluorescence parameters, F$_{V'}/$F$_{M'}$ and $\phi_{PSII}$, and no affect on the qP (Table 2).

To better evaluate the effects of salt stress on the primary photochemistry of PSII, we examined the absorption flux (ABS), trapping flux (TR) and electron transport flux (ET) per PSII reaction center (RC), i.e. ABS/RC, TR/RC, ET/RC. Our results show that these parameters did not change significantly at 0 – 200 mM. Our results show that there was no significant change in the trapping flux per PSII reaction center (TR/RC) in salt-stressed plants. TR/RC represents the maximum rate of reduction of $Q_a$ (Strasser et al., 2000). No change in TR/RC in salt-stressed plants proposes that the high conversion efficiency of excitation energy was preserved. According to the results in this study, it is suggested that through a down-regulation of PSII reaction centers and a shift of excitation energy transfer to PSI, the PSII apparatus was thus protected from further excess excitation energy by increased in DI$_0$/RC.

One of the parameters used to analyze the response of the plant to stress is the PI$_{ABS}$. In previous studies, PI$_{ABS}$ was used to estimate plant vitality in response to different environmental stresses to create vitality groups (Strasser et al., 2000). The decrease of the PI$_{ABS-re}$ in response to salinity stress in Hyola 61, Hyola 401 and Hyola 420 was mainly due to a decrease of the photochemical efficiency of photosynthetic electron transport and an increase of DI$_0$/RC in high salt. The increase of the PI$_{ABS-re}$ in response to salinity stress in Hyola 432 was principally due to an increase in the efficiency of primary photochemistry and photochemical efficiency of photosynthetic electron transport associated with a decreased DI$_0$/RC.

Linear correlations were observed between the log functions of the Performance Index, the log of the three independent components of PI$_{ABS}$ and the log of the relative electron transport activity. The linear correlation observed between the two log functions of the Performance Index and the electron transport activity indicates that changes in $\phi_{et}$ were responsible for most of the change in the multi-parametric expression of PI$_{ABS}$ in the case of salt stress, but additional alterations in the components of PI$_{ABS}$ cannot be ruled out. When Oukarroum et al. (2007) studied drought and re-watering on 10 varieties of barley (Hordeum vulgare L.), they described the same occurrence. Misra et al. (2001) studied salt and ion effects on the photosynthetic system that showed the same linearity between these two log functions, and they proposed that this analysis allowed them
to determine the susceptibility of the different genotypes to salt or ion stress. The same correlation was used to estimate the relative health of *Platanus acerfolia* L. trees in a quality assessment of urban trees (Hermans et al., 2003). Thus, we conclude that the decrease in growth cannot be attributed only the variation in the fluorescence parameters, but the salt effect on fluorescence parameters is mainly observed in high salt. The response to salt stress is dependent on the genotype, and the Performance Index can be used to distinguish the level of tolerance to stress.

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