Energetic flux and performance index in copaiba (Copaifera multijuga Hayne) and mahogany (Swietenia macrophylla King) seedlings grown under two irradiance environments

José F.C. Gonçalves*, Ulysses M. Santos Jr., Adamir R. Nina Jr. and Larissa R. Chevreuil

Instituto Nacional de Pesquisas da Amazônia, Laboratório de Fisiologia e Bioquímica Vegetal, Caixa Postal 478, 69011-970 Manaus, AM, Brasil. *Corresponding author: jfc@inpa.gov.br. Phone: +55 92 3643-1938, Fax: +55 92 3643 1838

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Plant adaptation under conditions of limited irradiance can be investigated by their efficiency while using the available energy with a minimal thermodynamic energy loss. For this reason, we compared the energetic fluxes and the performance index (PI$_{ABS}$) in copaiba (Copaifera multijuga) and mahogany (Swietenia macrophylla) seedlings under both sunlight and shade. The experiment was carried out in Manaus (3º8’S, 59º52’W), Amazonas State, Brazil. The fluorescence transients were obtained in intact and healthy leaves using a Plant Efficiency Analyser (PEA) between 1200 h and 1300 h. Using the O-J-I-P test, the following parameters were calculated: 1) density of reaction centers per cross section (RC/CS), 2) maximum quantum yield for primary photochemistry of photosystem II ($\phi_{Po}$), and 3) probability of energy excitation ($\psi_o$) or that of an absorbed photon ($\phi_{Eo}$) to move an electron further than quinone A, and 4) the performance index (PI$_{ABS}$). The highest PI$_{ABS}$ was found in seedlings under shade for both species. Mahogany seedlings exhibited values of PI$_{ABS}$ about three-fold and 14-fold higher than copaiba seedlings under shade and sunlight, respectively. For RC/CS, reaction centers were inactive under sunlight, 58% for mahogany and 78% for copaiba. The high irradiance induced a decrease of $\phi_{Po}$ in mahogany (36%) and in copaiba (69%) when compared to shaded seedlings. This likely was a result of need for excess energy dissipation under sunlight. In addition, it was verified that seedlings under sunlight presented a severe decrease in $\phi_{Eo}$, particularly in copaiba (23-fold), when compared to mahogany (9-fold). Therefore, mahogany seedlings showed the highest performance in energy use under low and high irradiance, thus it may be proposed that mahogany presented lower entropy than copaiba seedlings, as demonstrated by the higher plasticity in the use of energy.

**Key words**: chlorophyll a fluorescence, energy fluxes in photosystem II, light acclimation, O-J-I-P transients, tropical tree species

Fluxo energético e índice de desempenho em plantas jovens de copaíba (Copaífera multijuga) e de mogno (Swietenia macrophylla) sob dois ambientes de irradiação: Adaptação de plantas sob condições limitantes de irradiação pode ser investigada pela eficiência do uso da energia disponível, com uma perda mínima da energia termodinâmica. Por essa razão, foram comparados o fluxo energético e o índice de desempenho (PI$_{ABS}$) em plantas jovens de copaíba (Copaífera multijuga) e de mogno (Swietenia macrophylla) sob condições de sol e sombra. O experimento foi conduzido em Manaus (3º8’S, 59º52’W). Os transientes de fluorescência foram obtidos em folhas saudáveis e intactas, utilizando-se de um fluorômetro portátil, no período entre 1200 h e 1300 h. A partir do teste O-J-I-P foram calculados os seguintes parâmetros: 1) densidade de centros de reação por seção transversal (RC/CS), 2) eficiência máxima do fotossistema II ($\phi_{Po}$), 3) probabilidade da energia de excitação ($\psi_o$) ou de um fóton absorvido ($\phi_{Eo}$) mover um elétron após a quinoma A,
INTRODUCTION

Forest environment presents high heterogeneity in terms of light availability, both spatially (forest layers, forest understory, sub-canopy environments and high gaps) and temporally (seasonal periods). Besides such heterogeneity, human activities (extensive deforestation) and/or natural events can promote frequent and significant changes in balanced environments, thus the establishment of a species and/or its continuity in the system are functions of its capacity to adapt its metabolism to the new environmental conditions (Kruger et al., 1997; Gonçalves and Santos Jr., 2005a). Irradiance availability is the basic factor for the energy flux in biological systems, largely determining the physiological processes of plants. Depending on the capture and use capacity of irradiance, plants present different and decisive responses for their survival, growth and adaptation under different light conditions. For this reason, shaded or full sunlight environments can inhibit the photosynthetic processes either because of shortage or of excess energy input (Zhang et al., 2003). In this way, the capacity of plants in adjusting their metabolism to optimize the use of primary resources such as light will determine the success or failure of their adaptation and establishment under given conditions of irradiance.

The chlorophyll a fluorescence technique is of particular significance in natural environments to study how the irradiance availability affects the tree photosynthetic apparatus and its efficiency under different light conditions (Gonçalves et al., 2001, 2005; Gonçalves and Santos Jr., 2005b; Santos Jr. et al., 2006). The analysis of chlorophyll a fluorescence emitted by PSII is a fast, easy, and non-invasive tool to evaluate the plant’s photosynthetic performance, providing precise and objective data with regard to photochemical efficiency and the processes of non-photochemical de-excitation involved in the conversion of light energy under different conditions (Clark et al., 2000; Misra et al., 2001; Baker and Rosenqvist, 2004; De Ronde et al., 2004; Gonçalves and Santos Jr., 2005b; Panda et al., 2006). The PSII unit catalyses the light-induced electron transport from water to the plastoquinone (PQ) pool molecule and consists of up to 25 different types of protein subunits, which are organized into two structurally and functionally different parts (Hankamer et al., 1997; Stirbet et al., 1998). The first part is the core complex, a well-defined structure responsible for all electron transfer reactions in PSII and organized as a dimer in the stacked, appressed regions of the thylakoid membrane. The second part is the peripheral antenna; in plants it consists of a collection of light-harvesting complex II (LHCII) proteins which absorb most of the light for PSII (Dekker and Grondelle, 2000). Under conditions of high irradiance, PSII is recognizable the first protein complex to exhibit damage. For this reason, a study of the structural and functional behavior of PSII can allow the examination of the primary effects of stress caused by excess irradiance on plants.

In this study, two species found in the primary Amazonian forest, mahogany (Meliaceae) and copaiba (Leguminosae: Caesalpinioideae), were compared to investigate interspecific variations in chlorophyll a fluorescence. Mahogany is one of the most valuable neotropical timber species in tropical forests (Whitman et
al., 1997) whereas copaiba has been traditionally and widely used as oil or capsules that have been exported from Brazil to other countries such as France, Germany and United States (Veiga and Pinto, 2001; Veiga et al., 2001). The main goal of this study was to evaluate the efficiency in capture and use of light energy and the performance index in environments of low and high irradiance through the analysis of changes in the shape of O-J-I-P transients, the parameters of chlorophyll a fluorescence and the performance index (PI$_{ABS}$) in those species.

**MATERIAL AND METHODS**

**Study area and species investigated:** This study was conducted in the experimental area of the V-8 Campus at the Instituto Nacional de Pesquisas da Amazônia, Manaus (3º8’S, 59º58’W), Amazonas State, Brazil. The experiment was carried out using eight-month-old mahogany (*Swietenia macrophylla* King) and copaiba (*Copaifera multijuga* Hayne) seedlings. Seedlings were obtained from seeds collected in the Adolfo Ducke Forest Reservation, located 10 km from the city of Manaus. Seeds were selected according to their uniformity, size and healthy state. After disinfections in sodium hypochlorite solution 1% chlorine (v/v) for 10 min, seeds were washed in running tap water and then in distilled water. Afterwards, they were placed to germinate in plastic boxes filled with washed sand as substrate. After reaching a height of 10 cm, the seedlings were transferred to black polyethylene bags containing 5 kg of an organic soil and sand mixture (2:1, w/w), where they remained for 30 d for acclimatization. The seedlings were grown under mid-intensity irradiance (maximum irradiance = 250-500 µmol m$^{-2}$ s$^{-1}$) until reaching eight months old. Afterwards they were assigned to two light treatments: shade, maximum irradiance = 10-30 µmol m$^{-2}$ s$^{-1}$, and sunlight, maximum irradiance = 1800-2300 µmol m$^{-2}$ s$^{-1}$, under natural day/night cycles of about 12-13/12-11 h. During the experiment, plants were watered daily. The data were collected after 120 d of the experiment when the young leaves growing in the new environment were fully expanded.

**Measurements of Chlorophyll a fluorescence:** The chlorophyll a fluorescence was measured in healthy and completely expanded leaves using a hand fluorometer (Plant Efficiency Analyser-MK2–9600, Hansatech, Norfolk, UK). Measurements were taken between 1200 and 1300 h using 10 plants per species per irradiance level. The selected leaves were subjected to a 20 min dark-adaptation period, a period of time sufficient for all reaction centers of PSII to become open (Rhee et al., 1998; Zouni et al., 2001). Immediately after the dark adaptation, the leaves were exposed to a pulse of saturating light at an intensity of 2700 µmol m$^{-2}$ s$^{-1}$, with a wavelength of 650 nm for 5 s. The fluorescence transients were recorded from 10 µs to 5 s at 12 bit resolution (Strasser et al., 2000). The Biolyzer software (Maldonado-Rodriguez, 2000) was used to load the full fluorescence transients and to calculate the O-J-I-P parameters from variable fluorescence values at $F_{500\mu s}$, $F_{100\mu s}$, $F_{500\mu s}$, $F_{2\mu s}$ and $F_{300\mu s}$, according to the equations of the O-J-I-P-test (Strasser and Govindjee, 1992; Strasser and Tsimilli-Michael, 2001) (see Table 1 for further details). The specific fluxes expressed per reaction centers (ABS/RC; TR$_O$/RC; DI$_O$/RC; ET$_O$/RC) and phenomenological fluxes per cross section (ABS/CS; TR$_O$/CS; DI$_O$/CS; ET$_O$/CS) were derived from the theory of energy flux from biomembranes (Sironval et al., 1981) and were calculated using the O-J-I-P test (Strasser and Tsimilli-Michael, 2001). Absorbance (ABS) refers to the absorption of photons by the chlorophyll molecules in the antenna complex. Part of the absorbed energy was trapped (TR$_O$) by the reaction centre of PSII (P$_{400}$) while the remainder was dissipated (DI$_O$) in the form of heat and fluorescence. Of the trapped energy, part of it was converted to redox energy by electron transport (ET$_O$) through $Q_A$ and $Q_B$ (Strasser et al., 2000). Furthermore, the following variables were determined: the density of reaction centers (RC) per cross section (RC/CS), the maximum quantum efficiency of PSII primary photochemistry ($\phi_{po}$ = TR$_O$/ABS), the maximum efficiency of non-photochemical de-excitation ($\phi_{po} = DI_O$/ABS) and, the probability that a trapped exciton ($\psi = ET_O/TR_O$) or that an absorbed photon ($\phi_{es} = ET_O/ABS$) moves an electron further than quinone A ($Q_A$). The performance index (PI) has been defined as the ratio of two structure-function indexes (SFI). The first, SFI$_{1}$ (%ABS$)$ ((Chl$_{RC}$/Chl$_{tot}$) x $\phi_{po}$ x $\psi$), responds to structural and functional PSII events leading to electron transport within photosynthesis (Tsimilli-Michael et al., 1998). The second, SFI$_{2}$ (%ABS$)$ [(1-Chl$_{RC}$/Chl$_{tot}$) (1 - $\phi_{po}$) x (1 - $\psi$)], refers to the energy that is dissipated or lost from photosynthetic electron...
Table 1. The O-J-I-P-test parameters formulae and definitions. See Strasser et al. (2004) for further details.

| Parameters | O-J-I-P test formulae | Definitions |
|------------|------------------------|-------------|
| **Extracted fluorescence parameters** | | |
| FO | \( \text{O}(F_{50\mu s}) \) | Initial fluorescence |
| O | \( F_{(F_{50\mu s})} \) | Fluorescence intensity at 50 \(\mu\)s |
| J | \( F_{(2ms)} \) | Fluorescence intensity at 2 ms |
| I | \( F_{(30ms)} \) | Fluorescence intensity at 30 ms |
| P or Fm | Maximum fluorescence |
| Fm | \( F_{100\mu s} \) | Fluorescence intensity at 100 \(\mu\)s |
| | \( F_{300\mu s} \) | Fluorescence intensity at 300 \(\mu\)s |
| | \( t_{Fm} \) | Time to reach \(F_m\) (ms) |
| **Calculated parameters** | | |
| Fv | \( (F_{m} - F_{0}) \) | Variable fluorescence |
| VJ | \( \frac{[F_{2ms} - F_{0}]}{[F_{m} - F_{0}]} \) | Relative variable fluorescence at J-step |
| MO | \( \frac{4(F_{300\mu s} - F_{50\mu s})}{(F_{m} - F_{50\mu s})} \) | Net rate of PSII closure |
| Area | \( \frac{\text{Area}}{F_{m}} \) | Area between the fluorescence curve and \(F_{m}\) |
| SM | Normalised area |
| **Specific fluxes (RC)** | | |
| ABS/RC | \( \frac{[\text{TR}_{\gamma}/\text{RC}]}{[\text{TR}_{\gamma}/\text{ABS}]} \) | Effective antenna size of an active RC |
| TR\(_{\gamma}/\text{RC} \) | \( \frac{(M_{\gamma}/V_{\gamma})}{(M_{\gamma}/F_{m})} \) | Maximal trapping rate per RC |
| DI\(_{\gamma}/\text{RC} \) | \( \frac{[\text{ABS}/\text{RC} - \text{TR}_{\gamma}/\text{RC}]}{[\text{ABS}/\text{RC} - \text{TR}_{\gamma}/\text{RC}]} \) | Dissipation of an active RC |
| ET\(_{\gamma}/\text{RC} \) | \( \frac{[\text{TR}_{\gamma}/\text{RC}]}{[\text{ET}_{\gamma}/\text{TR}_{\gamma}]} \) | Electron transport of an active RC |
| **Phenomenological fluxes (CS)** | | |
| ABS/CS | Approximately proportional to \(F_{0}\) | Number of photons absorbed per CS |
| TR\(_{\gamma}/\text{CS} \) | \( \frac{\text{ABS/CS} \times [\text{TR}_{\gamma}/\text{ABS}]}{\text{ABS/CS} \times [\text{TR}_{\gamma}/\text{ABS}]} \) | Energy flux for trapping per CS |
| DI\(_{\gamma}/\text{CS} \) | \( \frac{\text{ABS/CS} \times [\text{TR}_{\gamma}/\text{CS}]}{\text{ABS/CS} \times [\text{TR}_{\gamma}/\text{CS}]} \) | Energy dissipation per CS |
| ET\(_{\gamma}/\text{CS} \) | \( \frac{[\text{ET}_{\gamma}/\text{TR}_{\gamma}]}{[\text{RC}/\text{CS}]} \) | Electron transport per CS |
| RC/CS | \( \frac{[\text{ABS/CS} \times \text{RC/ABS}]}{[\text{ABS/CS} \times \text{RC/ABS}]} \) | Density of reaction centres per CS |
| **Yields (or flux ratios)** | | |
| \( \phi_{Po} \) | \( F_{m}/F_{o} = 1-(F_{o}/F_{m}) \) | Maximum quantum yield of primary photochemistry |
| \( \phi_{Do} \) | \( \frac{\text{DI}_{\gamma}/\text{ABS}}{\text{ABS}} \) | Maximum quantum yield of non-photochemical de-excitation |
| \( \psi_{a} \) | \( \frac{1}{\text{ET}_{\gamma}/\text{TR}_{\gamma}} \) | Probability that a trapped exciton moves an electron further than \(Q_{A}^{-}\) |
| \( \phi_{Eo} \) | \( \frac{\text{ET}_{\gamma}/\text{ABS}}{\text{ABS}} \) | Probability that an absorbed photon moves an electron further than \(Q_{A}^{-}\) |
| **Performance index** | | |
| PI\(_{\text{ABS}} \) | \( \frac{\text{SFI}_{\text{po}}(\text{ABS})}{\text{SFI}_{\text{No}}(\text{ABS})} = \frac{[\text{Chl}_{\text{RC}}/\text{Chl}_{\text{tot}}][1-(\text{Chl}_{\text{RC}}/\text{Chl}_{\text{tot}})]}{[\text{Chl}_{\text{RC}}/\text{Chl}_{\text{antenna}}]} \) | Performance index |

Experimental design and statistical analysis: The experiment was arranged in a completely randomized design in a 2x2 factorial scheme with two species (co-transport, in which Chl\(_{\text{tot}}\) is the total Chl \(a\) concentration, and Chl\(_{\text{antenna}}\) = Chl\(_{\text{RC}}\) + Chl\(_{\text{antenna}}\) (Strasser et al., 1999).

The combination of both structure-function indexes leads to the expression performance index (PI) and when based on absorption of antenna Chls of PSII (PI\(_{\text{ABS}}\)) can be represented as:

\[
\text{PI}_{\text{ABS}} = \text{SFI}_{\text{po}}(\text{ABS})/\text{SFI}_{\text{No}}(\text{ABS}) = \frac{[\text{Chl}_{\text{RC}}/\text{Chl}_{\text{tot}}][1-(\text{Chl}_{\text{RC}}/\text{Chl}_{\text{tot}})]}{[\text{Chl}_{\text{RC}}/\text{Chl}_{\text{antenna}}]} \left[ \frac{\phi_{Po}}{1-\phi_{Po}} \right] \left[ \psi_{a}/(1-\psi_{a}) \right]
\]

or, in terms of the expression used in the O-J-I-P-test (Srivastava et al., 1999):

\[
\text{PI}_{\text{ABS}} = \left( \frac{\text{RC/CS}}{[\text{ABS/CS}]} \right) \left[ \frac{\phi_{Po}}{1-\phi_{Po}} \right] \left[ \psi_{a}/(1-\psi_{a}) \right].
\]
paiba and mahogany) and two irradiance environments (shade and full sunlight). For each treatment 10 replicates (seedlings) were used. The results were submitted to variance analysis and the means were compared by the Tukey test at 5% probability. The statistical analyses were conducted using the software SYSTAT (version 8.0).

RESULTS

Analysis of Chl a fluorescence transients: The leaves of the copaiba and mahogany plants under shade exhibited typical polyphasic Chl a fluorescence O-J-I-P transients, rising from an initial fluorescence level (F₀) to the maximum fluorescence level (Fm) (Figure 1A-B). Thus, Chl a fluorescence transients rose sharply from O to P steps under shade compared to sunlight. Furthermore, leaves under full sunlight failed to show typical Chl a fluorescence O-J-I-P transients. The times (t₁,o) to reach maximum fluorescence in both species under shade were higher than under sunlight (Table 2). The area beneath the fluorescence curve between F₀ and Fm were lower in sunlight than in shade, suggesting a decrease in the electron pool size of PSII, including QA, QB and PQ (Figure 1A-B, Table 2) (Joliot and Joliot, 2002).

Under shade, copaiba presented a higher increase in fluorescence intensity at 2 ms (J phase) than mahogany, suggesting an accumulation of the fraction of reduced QA pool (Figure 1C-D). In addition, mahogany under shade presented higher values than copaiba for the area beneath the fluorescence curve between F₀ and Fm, suggesting that mahogany exhibited the highest electron pool size of PSII (Table 2). The resulting difference of normalized transients ΔWₜ₀ between sunlight and shade allowed the visualization of the K-band in copaiba (Figure 1E) and mahogany (Figure 1F). This indicates that the k-band was more pronounced after high irradiance compared to shade.

Chl a fluorescence parameters (specific and phenomenological fluxes): Examination of the specific flux (per RC) showed an increase in the functional “antenna size” i.e. (ABS/RC), as given by light absorbed per active RC in seedlings under high irradiance. The increases of ABS/RC were close to 4.7-fold and 2.3-fold in copaiba and mahogany under full sunlight compared to shade, respectively (Figure 2A). This increase in ABS/RC was due to an inactivation of active RC units per cross section (RC/CS) under high irradiance (Table 3). The trapping rate of PSII (TR₁/RC) was 42% and 47% higher in copaiba and mahogany, respectively, under full sunlight compared to shade (Figure 2B). In addition, due to the excess of energy and the decrease of RC/CS under high irradiance, an increase of 17.6-fold and 6.4-fold was observed in the effective dissipation per active RC (DI₀/RC) in copaiba and mahogany, respectively (Figure 2C). As a result of high DI₀/RC, the values of ET₀/RC were 75% and 80% in mahogany and copaiba seedlings, respectively, under full sunlight compared to shade (Figure 2D).

Between the studied species it was observed that in both irradiance environments copaiba seedlings presented the highest values of ABS/RC (135% in sunlight against 16% in shade), TR₁/RC (8% in sunlight and 12% in shade) and DI₀/RC (239% in sunlight and 38% in shade) than mahogany (Figure 2A-C). However, due to the high values of DI₀/RC in copaiba, the electron transport rates in active RC were 24% and 56% higher in mahogany than in copaiba under shade and sunlight, respectively (Figure 2D).

Considering the phenomenological fluxes, it was found that copaiba and mahogany presented higher values of TR₀/CS and ET₀/CS under shade than sunlight (Figure 3B, 3D). In addition, the values of DI₀/CS were 155% and 253% higher in mahogany and copaiba, respectively, under full sun than under shade (Figure 3C). Copaiba seedlings showed higher values of ABS/CS (15%), TR₀/CS (10%) and DI₀/CS (37%) than mahogany under shade (Figure 3A-C). Under high irradiance, copaiba presented the lowest value of TR₀/CS and the highest value of DI₀/CS (Figure 3B-C). For ET₀/CS, in both irradiance environments, mahogany exhibited the highest values (Figure 3D).

Yields and performance index: The results of flux ratio (yields) demonstrated that both copaiba and mahogany presented a decrease in TR₀/ABS (ϕₒ), ET₀/TR₀ (ψₒ) and, consequently, in ET₀/ABS (ϕₑₒ) under high irradiance (Table 3). The decrease of ϕₒ in mahogany (36%) and copaiba (69%) under high irradiance indicates a loss of maximum quantum efficiency of primary
Figure 1: (A, B) – Polyphasic Chl a fluorescence transients recorded in copaiba and mahogany under shade and sunlight. Each transient represents the mean of 10 transients. (C, D) – Transients normalized between O and P. (E, F) – Change in the shape of the Chl a fluorescence transient curves normalized between O and J (WOJ) showing k-band. \[ \Delta W_{OJ} = (W_{OJ(\text{sunlight})} - W_{OJ(\text{shade})}) \].

Photochemistry due to photoinhibition promoted by excess energy. In addition, such excess induced an inactivation of 58 and 78% of active RC per cross section in mahogany and copaiba under sunlight, respectively, inducing an increased energy dissipation as well as lower values of \( \phi_p \) (Table 3). Further, it was verified that seedlings under sunlight presented a decrease in \( \phi_e \), especially for copaiba (23-fold) when compared to mahogany (nine-fold).

The performance index (PI\text{ABS}) combines three
The shape of the O-J-I-P transients is very sensitive to different light conditions. Under high irradiance the shape of the O-J-I-P transient changed in copaiba and mahogany with a severe decrease in F_m, resulting in lowered F_v (Figure 1.A-B, Table 2). The fast rise from O to J is the photochemical phase and J to P is the thermal phase (Strasser et al., 1995; Lazar, 2003). The rise in transients from O to J and from J to P are due to the reduction of QA to QA\(^{-}\) and the reduction of the PQ pool (Strasser and Govindjee, 1992). Thus the high irradiance severely affected both donor and acceptor sides of PSII due to the disconnected oxygen-evolution complex with impairment of the electron transport chain (Figure 1.E-F). Comparing both species, stronger photoinhibition induced by excess of light energy, promoting more damage at both sides of PSII, is to be expected for copaiba than in mahogany.

The time to reach the complete reduction of PSII acceptor QA was higher in shade than in sunlight. The shorter time under high irradiance is caused by the faster rate of QA reduction due to high light availability in the environment. Since F_o remained unchanged regardless of species and treatments (Table 2), a regulatory or protective mechanism under high irradiance is to be expected (Björkman and Demmig Adams, 1987; Osmond et al., 1993). In addition, the decline in both F_m and F_v/F_m

### Table 2.

| Variables | COPAÍBA | MAHOGANY |
|-----------|---------|----------|
| FO        | 662 ± 144 Aa | 681 ± 59 Aa |
| F_50µs    | 666 ± 147 Ba | 823 ± 66 Aa |
| F_100µs   | 730 ± 178 Ba | 971 ± 81 Aa |
| F_300µs   | 822 ± 223 Ba | 1494 ± 132 Aa |
| F_2ms     | 880 ± 247 Ba | 2265 ± 193 Aa |
| F_30ms    | 861 ± 237 Bb | 3058 ± 200 Aa |
| F_m       | 230 ± 110 Bb | 2635 ± 171 Ab |
| F_v       | 893 ± 251 Bb | 3317 ± 219 Aa |
| Area      | 212 ± 26 Ba | 304 ± 18 Ab |
| Area      | 3438 ± 1575 Ba | 32856 ± 3505 Ab |

The relationship between log (PI\(_{ABS}\))\(_{rel}\) = \[\log\left(\frac{PI\(_{ABS}(Copaíba\ or\ Mahogany)}{PI\(_{ABS}(Copaíba\ or\ Mahogany\ average)}\right)\] and the relative yield of electron transport (ET/ABS)\(_{rel}\) = \[\frac{ET/ABS\(_{Copaíba\ or\ Mahogany}\)}{ET/ABS\(_{Copaíba\ or\ Mahogany\ average}\)}\] in Figure 4 shows that shaded seedlings exhibited better capacity to use irradiance than seedlings under full sunlight. As expected from the above data, mahogany presented higher capacity to convert light energy into chemical energy than copaiba in both irradiance environments (Figure 5A-B).

**DISCUSSION**

The shape of the O-J-I-P transients is very sensitive to different light conditions. Under high irradiance the shape of the O-J-I-P transient changed in copaiba and mahogany with a severe decrease in F_m, resulting in lowered F_v (Figure 1.A-B, Table 2). The fast rise from O to J is the photochemical phase and J to P is the thermal phase (Strasser et al., 1995; Lazar, 2003). The rise in transients from O to J and from J to P are due to the reduction of QA to QA\(^{-}\) and the reduction of the PQ pool (Strasser and Govindjee, 1992). Thus the high irradiance severely affected both donor and acceptor sides of PSII due to the disconnected oxygen-evolution complex with impairment of the electron transport chain (Figure 1.E-F). Comparing both species, stronger photoinhibition induced by excess of light energy, promoting more damage at both sides of PSII, is to be expected for copaiba than in mahogany.

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Figure 2. Specific fluxes expressed per reaction center (RC) in copaiba and mahogany under sunlight and shade environments. (A) – Flux of photons absorbed by chlorophyll molecules per active RC (ABS/RC); (B) – Energy flux for trapping that is conserved in chemical components per active RC (TRo/RC); (C) – Flux of energy that enters into the electron transport chain further than Q_A per active RC (ETO/RC); (D) – Flux of energy that is dissipated in the form of heat and fluorescence, or transferred to other systems per active RC (DIO/RC). Statistics as in Table 2.

Table 3. Density of active reaction centers per cross section (RC/CS), maximum efficiency of PSII (\( \phi_p = TRo/ABS \)), maximum efficiency of non-photochemical de-excitation (\( \phi_D = DIo/ABS \)), probability that a trapped exciton (\( \psi_o = ETo/TRo \)) or that an absorbed photon (\( \psi_Eo = ET_o/ABS \)) can move an electron further from Q_A, and performance index (PI\_ABS) in copaiba and mahogany under two irradiance environments. Statistics as in Table 2.

| Variables             | COPAIBA       | MAHOGANY     |
|-----------------------|---------------|--------------|
|                       | Sunlight      | Shade        | Sunlight      | Shade        |
| RC/CS                 | 58 ± 23 Bb    | 264 ± 21 Aa  | 113 ± 29 Ba   | 265 ± 4 Aa   |
| TRo/ABS (\( \phi_p \)) | 0.25 ± 0.05 Bb | 0.80 ± 0.01 Ab | 0.53 ± 0.08 Ba | 0.83 ± 0.003 Aa |
| DIo/ABS (\( \phi_D \))  | 0.75 ± 0.05 Aa | 0.20 ± 0.01 Ba | 0.47 ± 0.08 Ab | 0.17 ± 0.003 Bb |
| ETo/TRo (\( \psi_o \))  | 0.057 ± 0.024 Bb | 0.399 ± 0.041 Ab | 0.095 ± 0.033 Ba | 0.556 ± 0.018 Aa |
| ETo/ABS (\( \psi_Eo \)) | 0.014 ± 0.006 Bb | 0.317 ± 0.030 Ab | 0.052 ± 0.022 Ba | 0.460 ± 0.015 Aa |
| PI\_ABS               | 0.002 ± 0.001 Bb | 1.013 ± 0.186 Ab | 0.028 ± 0.019 Ba | 2.725 ± 0.286 Aa |
indicates an adaptation to a physiological state which requires, under high-irradiance conditions, a much lower quantum yield of PSII to sufficiently reduce the primary acceptor Q_a.

The high irradiance induced a severe decrease in the area above the fluorescence between $F_o$ and $F_m$ in seedlings under sunlight, indicating a reduction in the pool size of PSII (Strasser et al., 1995; Joliot and Joliot, 2002).

The $\phi_{po}$ ($F_o/F_m = TR_o/ABS$) is a parameter that expresses the maximum efficiency of PSII which is controlled by the primary photochemistry of PSII (charge separation, recombination and stabilization), non-radiative loss of excited states in light-harvesting antennae and by excited states quenched by oxidized PQ molecules from the PQ pool (Lazar, 2003). The low values of $\phi_{po}$ in seedlings under high irradiance could have resulted from the inactivity of RC, which might favor higher energy dissipation as heat and fluorescence, as deduced from the high values for $\phi_{Do}$. This might be associated with increased heat sinks (heat-sink centers or silent centers). These heat sinks may absorb light in a similar manner to active RC, but are unable to store the excitation energy as redox energy, dissipating their total energy as heat (Hermans et al., 2003). In addition, due to excess of irradiance, transfer of energy to other systems

Figure 3. Phenomenological fluxes expressed per cross section of leaf tissue (CS) in copaiba and mahogany under sunlight and shade environments. (A) – Flux of photons absorbed by chlorophyll molecules per CS (ABS/CS); (B) – Energy flux for trapping that is conserved in chemical components per CS (TR_o/CS); (C) – Flux of energy that enters into the electron transport chain further than Q_A per CS (ET_o/CS); (D) – Flux of energy that is dissipated in the form of heat and fluorescence, or transferred to other systems per CS (DI_o/CS). Statistics as in Table 2.
such as energy-dependent formation of reactive oxygen species could also took place (Strasser et al., 2000). Analyzing $\psi_0$, the lowest values of $\varphi_{Po}$ were found in seedlings under high irradiance. The values of $\psi_0$ decreased to a remarkably greater extent in both mahogany and copaiba under sunlight than in shade. This result reflects a reduction in the pool of plastoquinone in an oxidized state (PQ) and reoxidation inhibition of QA$.^-.$ and demonstrates that besides the loss of energy to QA, the loss of excitation energy further from QA was significant (Force et al., 2003). Results of $\varphi_{Po}$, $\psi_0$ and $\varphi_{Eo}$ allow us to deduce that mahogany may use light energy more efficiently than copaiba under both shade and full sunlight conditions (Table 3).

The performance index (PI$_{ABS}$) is a consistent parameter to evaluate the plant performance in light energy absorption, excitation energy trapping and conversion of excitation energy to electron transport by photosynthesis under stress conditions of irradiance, drought, heat, dark chilling, ozone and salt (Clark et al., 2000; Misra et al., 2001; De Ronde et al., 2004; Gonçalves and Santos Junior, 2005b; Strauss et al., 2006). As observed in this study, PI$_{ABS}$ was much lower in full-sun-grown seedlings. Among the factors that have likely

Figure 4. (A) – Relationship between the log function of the relative Performance Index (Log (PI$_{ABS}$)$_{rel}$) [$=$Log (PI$_{ABS(copaiba or Mahogany)}$/PI$_{ABS(copaiba or mahogany average)}$)] and the relative yield of electron transport (ET$_o$/ABS)$_{rel}$ [(ET$_o$/ABS$_{(copaiba or mahogany)}$/ET_o$/ABS$_{(copaiba or mahogany average)}$)]. (B) – Details of second quadrant; (C) – Details of third quadrant. The filled and empty squares represent copaiba under shade and sunlight, respectively. The empty triangles and empty circles represent mahogany under sunlight and shade, respectively.
Figure 5. (A) – Relationship between the log function of the relative Performance Index (Log (PI_{ABS})_{rel} [Log (PI_{ABS(sunlight)}/PI_{ABS(sunlight average)})] and the relative yield of electron transport (ET_{o}/ABS)_{rel} [(ET_{o}/ABS\_{(sunlight average)}/ET_{o}/ABS\_{(sunlight) average})]. (B) – Relationship between the log function of the relative Performance Index (Log (PI_{ABS})_{rel} [Log (PI_{ABS(shade)}/PI_{ABS(shade average)})] and the relative yield of electron transport (ET_{o}/ABS)_{rel} [(ET_{o}/ABS\_{(shade average)}/ET_{o}/ABS\_{(shade) average})]. The filled and empty squares represent copaiba under shade and sunlight, respectively. The empty triangles and empty circles represent mahogany under sunlight and shade, respectively.

The PI_{ABS} expresses both a function of the fluorescence extreme F_{o} and F_{m}, as well as the intermediate J-step and the slope at the origin of the fluorescence rise, whereas \(\varphi_{Po}\) expresses a function of only F_{o} and F_{m} independently of how the trajectory of the fluorescence intensity reaches its maximal value (Strauss et al., 2006). Furthermore, PI_{ABS} allows broader analysis of photosynthetic performance such as the relationship between photon absorption efficiency and capture of excited energy in PSII, as well as analysis of density of active RC and the probability that excited energy moves an electron further than Q_{A}^{-} (Gonçalves and Santos Junior, 2005a). Therefore, PI_{ABS} is a better parameter to evaluate responses of PSII to stressful conditions than \(\varphi_{Po}\) alone.

The correlation between log (PI_{ABS})_{rel} and (ET_{o}/ABS)_{rel} can be considered a typical property of a plant’s capacity to transform light energy into chemical energy, which is directed into metabolic reactions in the biochemical process of photosynthesis (Hermans et al., 2003). The low values of (ET_{o}/ABS)_{rel} and log (PI_{ABS})_{rel} in copaiba and mahogany under the stress condition, caused by high free energy availability in the system, suggest that adaptation to full sunlight of these species at the seedling stage is thermodynamically unfavorable due to the difficulty of using the excess energy in the photosynthetic process. On the other hand, under low irradiance, the thermodynamically optimal state is more easily achieved due to minimal entropy production, as a result of low free energy in the system.

The similar performance of mahogany seedlings under both sun and shade when compared to the contrasting performance of copaiba seedlings also under both sun and shade conditions (Figure 4A; see details in Figure 4B-C), and the superior performance of mahogany in both light environments (Figure 5A-B), suggest that this species is more plastic than copaiba for using the available energy at seedling stages.

**CONCLUSIONS**

We conclude that the investigation of the shape of O-J-I-P transients and the parameters obtained using Chl a fluorescence such as the performance index were effective to evaluate the efficiency in capture and use of low and high light under shade and sunlight, respectively, in copaiba and mahogany seedlings.
Copaiba and mahogany were not very efficient in their use of excess energy under high irradiance, suffering from photo inhibition. In contrast, seedlings use light energy more efficiently under low irradiance due to minimal entropy production. Mahogany presented higher plasticity for using the available energy than copaiba in both irradiance treatments. This suggests that mahogany has a greater capacity to become established under contrasting light environments than does copaiba in the juvenile phase. Finally, these indexes could also possibly be of great value as screening tools for high-irradiance tolerance of tree species in highly degraded areas.

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REFERENCES

Baker NR, Rosenqvist E (2004) Application of chlorophyll fluorescence can improve crop production strategies: an examination of future possibilities. J. Exp. Bot. 55:1607-162121.

Bjorkman O, Demmig-Adams B (1987) Photon yield of O2 evolution and chlorophyll fluorescence characteristics at 77 K among vascular plants of diverse origins. Planta 170:489-504.

Clark AJ, Landolt W, Bucher JB, Strasser RJ (2000) Beech (Fagus sylvatica) response to ozone exposure assessed with a chlorophyll a fluorescence performance index. Environ. Pollut. 109:501-507.

De Ronde JA, Cress WA, Kruger GHJ, Strasser RJ, Van Staden J (2004) Photosynthetic response of transgenic soybean plants, containing an Arabidopsis P5CR gene, during heat and drought stress. J. Plant Physiol. 161:1211-1224.

Dekker JP, Van Grondelle R (2000) Primary charge separation in photosystem II. Photosynth. Res. 63:195-208.

Force L, Critchley C, Rensen JJS (2003) New fluorescence parameters for monitoring photosynthesis in plants. Photosynth. Res. 78:17-33.

Gonçalves JFC, Marenco RA, Vieira G (2001) Concentration of photosynthetic pigments and chlorophyll fluorescence of mahogany and tonka bean under two light environments. Braz. J. Plant Physiol. 13:149-157.

Gonçalves JFC, Barreto DCS, Santos Junior UM, Fernandes AV, Sampaio PTB, Buckeridge MS (2005) Growth, photosynthesis and stress indicators in young rosewood plants (Aniba rosaeodora Ducke) under different light intensities. Braz. J. Plant Physiol. 17:325-334.

Gonçalves JFC, Santos Junior UM (2005a) Assimilação de carbono e indicadores de estresse da Amazônia. In: Nogueira RJMC, Araújo EL, Willadino LG, Cavalcante UMT (eds), Estresses Ambientais: Danos e Benefícios em Plantas, pp.165-181. UFRPE, Imprensa Universitária, Recife.

Gonçalves JFC, Santos Junior UM (2005b) Utilization of the chlorophyll a fluorescence technique as a tool for selecting tolerant species to environments of high irradiance. Braz. J. Plant Physiol. 17:307-313.

Hankamer B, Barber J, Boekema EJ (1997) Structure and membrane organization of photosystem II from green plants. Annu. Rev. Plant Physiol. Plant Mol. Biol. 48:641-672.

Hermans C, Smeyers M, Rodriguez RM, Eyletters M, Strasser RJ, Delhaye JP (2003) Quality assessment of urban trees: A comparative study of physiological characterization, airborne imaging and on site fluorescence monitoring by the O-J-I-P-test. J. Plant Physiol. 160:81-90.

Joliot P, Joliot A (2002). Cyclic electron transport in plant leaf. Proc. Natl. Acad. Sci. USA 99:10209-10214.

Kruger GHJ, Tsimilli-Michael M, Strasser RJ (1997) Light stress provokes plastic and elastic modifications in structure and function of photosystem II in camellia leaves. Physiol. Plant. 101:265-277.

Lazar D (2003) Chlorophyll a fluorescence rise induced by high light illumination of dark-adapted plant tissue studied by means of photosystem II and considering photosystem II heterogeneity. J. Theor. Biol. 220:469-503.

Maldonado-Rodriguez (2000) Biolizer: program developed in the Bioenergetics Laboratory of Geneva and available as free ware.
Screening the vitality and photosynthetic activity of plants by fluorescence transient. In: Behl RK, Punia MS, Lather BPS (eds), Crop Improvement for Food Security, pp.79-126. SSARM, Hisar.

Strasser RJ, Srivastava A, Tsimilli-Michael M (2000) The fluorescence transient as a tool to characterize and screen photosynthetic samples. In: Yunus M, Pathre U, Mohanty P (eds), Probing Photosynthesis: Mechanisms, Regulation and Adaptation, pp.445-483. Taylor & Francis, London.

Strasser RJ, Srivastava A, Tsimilli-Michael M (2004) Analysis of chlorophyll a fluorescence transient. In: Papageorgiou G, Govindjee (eds), Advances in Photosynthesis and Respiration: Chlorophyll Fluorescence a Signature of Photosynthesis, pp.321-362. Kluwer Academic Publishers, Netherlands.

Strasser RJ, Tsimilli-Michael M (2001) Stress in plants from daily rhythm to global changes, detected and quantified by the JIP test. Chim. Nouvelle (SRC) 75:3321-6.

Strauss AJ, Kruger GHJ, Strasser RJ, Van Heerden PDR (2006) Ranking of dark chilling tolerance in soybean genotypes probed by chlorophyll a fluorescence transient O-J-I-P. Environ. Exp. Bot. 56:147-157.

Tsimilli-Michael M, Pêcheux M, Strasser RJ (1998) Vitality and stress adaptation of the symbionts of coral reef and temperate foraminifers probed in hospite by the fluorescence kinetics O-J-I-P. Arch. Sci. Genéve 51:205-240.

Tsimilli-Michael M, Eggenberg P, Biro B, Koves-Péchy K, Voros I, Strasser RJ (2000) Synergistic and antagonistic effects of arbuscular mycorrhizal fungi and Azospirillum and Rhizobium nitrogen-fixers on the photosynthetic activity of alfalfa, probed by polyphasic chlorophyll a fluorescence transient O-J-I-P. Appl. Soil Ecol. 15:169-182.

Veiga Jr. VF, Pinto AC (2001) The Copaifera L. genus. Quimica Nova 25:273-286.

Veiga Jr. VF, Zunino L, Calixto JB, Patitucci ML, Pinto AC (2001) Phytochemical and antioedematogenic studies of commercial Copaiba oils available in Brazil. Phytother. Res. 15:476-480.

Whitman AA, Brokaw NVL, Hagan JM (1997) Forest damage caused by selective logging of mahogany (Swietenia macrophylla) in northern Belize. For. Ecol. Manage. 92:87-96.
Zhang S, Ma K, Chen L (2003) Response of photosynthetic plasticity of *Paeonia suffruticosa* to changed light environments. Environ. Exp. Bot. 49:121-133.

Zouni A, Witt HT, Kern J, Fromme P, Kraub N, Saenger W, Orth P (2001) Crystal structure of photosystem II from *Synechococcus elongatus* at 3.8 Å resolution. Nature 409:739-43.