Leaf gas exchange and biomass partitioning in *Jatropha curcas* L. young plants subjected to flooding and drought stresses

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Abstract

With the purpose of assessing the physiological responses of *J. curcas* young plants to drought and flooding, plants were subjected to three water regimes: Control, drought and flooding. In the same period, five plants were submitted to drought (irrigation suspension), five to flooding and five were maintained near the field capacity (control). Measurements of leaf gas exchange, chlorophyll content, biomass of each organ (leaf, stem and root), biomass ratios, as well as the root:shoot ratio were made. When compared to control, drought and flooding treatments led to significant reductions (P≤0.05) of gas exchange rates at 14 and 7 days after imposition treatments (DAIT), respectively. At the 16th DAIT, significant reductions of leaf, root, stem and total biomass were observed, which resulted in changes of biomass partitioning of each organ. Significant reduction of leaf dry biomass (LDB) was observed in water deficit plants. Moreover, no significant effects of water deficit were observed in root dry biomass. Such response has led to a higher root:shoot ratio, suggesting an improved ability for water and nutrients uptake. In turn, flooding has induced a significant decrease in the dry biomass of all organs. Stem dry biomass ratio of flooded plants (0.67 g g⁻¹) was significantly higher than that of control plants (0.41 g g⁻¹), which was seemingly due to leaf senescence and root system rotting of flooded plants. Altogether, present results reinforce the susceptibility of the studied genotype of *J. curcas* to flooding, as well as highlight some characteristics of drought tolerance in this species.

Keywords: Abiotic stress; biomass allocation; Euphorbiaceae; stomatal control; water use efficiency.

Abbreviations: DAIT_days after imposition treatments; DBR_dry biomass ratio; LA_leaf area; LDB_leaf dry biomass; SDB_stem dry biomass ratio; SLA_specific leaf area.

Introduction

In the current scenario of climate changes, plants are often exposed to various stresses, such as excess (flooding) or deficiency (drought) of water, which may affect their metabolic processes and, consequently, growth, development and yield (Chaves and Oliveira, 2004). Flooding-induced plant damage occur, especially by reducing the availability of O₂ and the diffusion of gases (mainly O₂ e CO₂) (Perata et al., 2011), inhibiting seed germination, vegetative and reproductive growth and, depending on the intensity, plant death (Kozlowski, 1997). The drought affects many physiological processes of the plants, generally by increasing the stomatal resistance, reducing transpiration and thus the supply of CO₂ (Nogueira et al., 2001). Despite the damage caused by this stress, the decrease in stomatal conductance, caused by the limited availability of water, may allow the plant to assimilate more molecules of CO₂ for each unit of water transpired, *i.e.*, increased water use efficiency is generally observed in water stressed plants (Blum, 2009). The use of appropriate crops that may grow in such harsh environments is justified (Chaves et al., 2003; Chaves and Oliveira, 2004). Given the climate changes, coupled with need of replacing fossil fuels with renewable fuels, the physic nut (*Jatropha curcas* L.) has drawn scientific and economic interest. Amongst the advantages of *J. curcas* over other oilseed crops, one may highlight the high oil production ranging from 30-35% of seed’s weight, the easy conversion of oil into biodiesel, and the lack of competitiveness with the food industry, unlike *Glycine max* L. and *Arachis hypogaea* L. (Fairless, 2007). In addition to these agronomic characteristics, *J. curcas* has been considered tolerant to drought, being reported consistent changes in biomass allocation in favor of root system, allowing them to survive when subjected to water deficit (Achten et al., 2010; Díaz-Lopez et al., 2012; Maes et al., 2009). The hardiness and adaptability of *J. curcas* to different agro-climatic conditions justify its broad distribution (Achten et al., 2008; Divakara et al., 2010) and allows its cultivation in degraded and/or dry climate areas (Srivastava et al., 2011). Based on the possible tolerance of this species to several stressful conditions, we workout the hypothesis that *J. curcas* can survive, for a determined period, to drought and flooding with no significant damage in terms of biomass production. Indeed, there is still the need for a better characterization of the physiological responses of this species when subjected to water stress. Taking into account the above information, this study aimed to evaluate leaf gas exchange and growth responses of *J. curcas* young plants to water stresses by drought and flooding.
Results

Changes in leaf gas exchange

Soil moisture was maintained at around 18.0% in control pots, whereas irrigation stoppage has led to a moisture decrease from 18.4% to 10.6%. Water deficiency or excess led to significant (P≤0.05) reductions in gas exchange variables. These reductions were observed from 7 and 14 days after imposition treatments (DAIT) in plants under flooding and water deficit, respectively (Fig 1).

After 16 days under flooding and drought conditions, the plants showed significant reduction in photosynthetic rate (A), with values of 8.0 and 20.7 μmol CO₂ m⁻² s⁻¹, respectively, against around 27.1 μmol CO₂ m⁻² s⁻¹ in control plants (Fig 1A). Stomatal conductance (gs) was lower in flooded plants, reaching the average value of 0.04 mol H₂O m⁻² s⁻¹ at 16 DAIT, i.e., 85% lower than in control plants (Fig 1B).

The transpiration rate (E) showed a similar trend. As compared to the results of plants exposed to water deficit and flooding showed reductions of 36% and 77%, respectively, at 16 DAIT (Fig 1C). Over the same period, the ratio of internal to atmospheric CO₂ concentration (Ci/Co) was around 46% of control in water deficit plants, whereas those subjected to flooding showed a reduction of 68% (Fig 1D).

In this period, regardless of stress, it was observed that the values of A and E had a positive relationship with gs (Fig 2), the reductions in these variables were strongly influenced by decreased gs. Despite the marked reduction in gs after irrigation suspension, no corresponding reduction in A was observed. Therefore, higher water use efficiency (WUE) was observed at 16 DAIT in stressed plants (160 and 195 CO₂ mol⁻¹ H₂O, respectively, in drought and flooded plants), as compared to irrigated plants (102 μmol CO₂ mol⁻¹ H₂O) (Fig 3).

When compared to control plants, it was observed that the instantaneous carboxylation efficiency (A/Ci) responded differently in stress conditions. While in plants under water deficit, A/Ci was higher from 14 DAIT. Flooded plants, there was an early (9 DAIT) reduction of these values, around 36% (Fig 4).

Effect of stress on photosynthetic pigment and growth

Despite the early response of J. curcas to flooding, the chloroplastic pigment content was not significantly affected (P≤0.05) throughout the period of stress, as well as in plants under drought. It must be noted that, at 16 DAIT, the contents of chlorophylls a, b and total, as well as those of carotenoids were significantly (P≤0.05) higher in plants under drought as compared to flooding and control treatments (Table 1).

The leaf area (LA) of J. curcas was negatively affected by the stresses, therefore, significant (P≤0.05) reductions of 48% and 84%, in plants under deficit and flooding, respectively, as compared to control. Unlike the effect on LA, drought and flooding led to significant (P≤0.05) increases in specific leaf area (SLA), with increases of 25% and 51%, respectively (Table 2).

Reduction in total biomass was 47 and 80%, respectively (Fig. 5A). In all treatments, dry biomass ratio (DBR) was peculiar to each organ (Fig. 5B), indicating that the stresses may alter the dry biomass partition in the plants of J. curcas. Although the control treatment has the highest biomass value for all organs, it is possible to observe that leaf DBR value was higher in plants submitted to drought (Figure 5B). This greater contribution of the root in relation to the total biomass of plants subjected to drought is confirmed by the higher root/shoot ratio (R/S) in this treatment (0.16 g g⁻¹) (Fig 5B).

Discussion

Reductions in leaf gas exchange rates are commonly observed in response to water deficit (da Silva et al., 2016; Pompelli et al. 2010; Silva et al. 2010; Tezara et al., 2003) and flooding (Santos et al., 2012; Verma et al., 2012). Stomatal closure is seen as a survival strategy, since it reduces hydraulic conductivity in plants subjected to these stresses reducing transpiration rate, and consequently, water absorption (de Santana et al., 2015; Hubbard et al., 2001; Pezeshki, 2001). As a response to soil flooding, this reduction is common in tolerant as well as non-tolerant to flooding (Kozlowski, 1997). Yet, in tolerant plants, the values are established due to development of hypertrophied lenticels. Such gs recovery was not observed in J. curcas plants, as reported for G. americana L., which showed reductions after 14 days even with the emergence of lenticels (Mielke et al., 2003). A species considered tolerant to flooding, Annona glabra L. showed a reduction in gs values from 4 days after the onset of the treatment, and a recovery to values comparable to control after 11 days (Mielke et al., 2005). The authors attributed this recovery to the development of hypertrophied lenticels, together with increased mass allocation to roots.

As opposed to water loss reduction, stomatal closure also reduces the carbon assimilation, thereby causing reduced photosynthetic rate (Chaves et al., 2002). The positive relationship of variables A and E in comparison with gs, in J. curcas specimens subjected to drought and flooding treatments indicates that stomatal closure may be the major limiting factor of these variables (Cornic, 2000; Sapeta et al., 2013; Silva et al., 2010). This interaction between gs and E, shows the importance of controlling stomatal movement (opening and closing), in plants adapted to stress condition, as occurred in J. curcas plants under water deficit, allowing the plant to maintain a high WUE (Naves-Barbiero et al., 2000; Yang et al., 2005).

Increased WUE in flooded plants results from a known physiological response in which transpiration rate is more affected than carbon assimilation (Medrano et al., 2009). However, the decrease of WUE after 14 days, suggest the occurrence of non-stomatal factor limiting the photosynthetic activity in J. curcas. That limitation may be due to lower pigments contents, with great effect on the irradiance uptake capacity (Pezeshki, 2001). Moreover, low rates A/Ci, observed in flooded plants, suggest the occurrence of biochemical limitation (Silva et al., 2011). Flooding led to reduction of leaf gas exchange variables in contrasting clones of Thespesia cacao for flooding tolerance, but non-stomatal limitation were observed only in the susceptible one (Bertolde et al., 2012). In the present work, non-stomatal limitation to photosynthesis in flooded plants seems to be related more to biochemical constraints (as demonstrated by decreased A/Ci) than to photochemical constraints (as no stress effect on foliar pigment content has been detected).

On the other hand, A/Ci was not affected in water deficit plants, which showed even an increase of this variable at the end of experiment, suggesting that photosynthetic limitation in these plants were due to stomatal closure. This characteristic would contribute for a rapid recovery of leaf gas exchange rates upon rewatering, as observed by Pompelli et al. (2010).
Table 1. Chlorophyll a (Chl a), chlorophyll b (Chl b), total chlorophyll (Chl a+b) and carotenoids (Carot) contents (μg cm$^{-2}$) in plants of *Jatropha curcas* under different water conditions, at 16 days after the imposition of treatment (DAIT).

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Chl a</th>
<th>Chl b</th>
<th>Chl (a+b)</th>
<th>Carot</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>29.41 ± 1.31b</td>
<td>7.31 ± 0.33b</td>
<td>36.72 ± 1.65b</td>
<td>8.07 ± 0.23b</td>
</tr>
<tr>
<td>Drought</td>
<td>35.31 ± 1.25a</td>
<td>8.81 ± 0.32a</td>
<td>44.12 ± 1.57a</td>
<td>9.06 ± 0.20a</td>
</tr>
<tr>
<td>Flooding</td>
<td>25.34 ± 1.66b</td>
<td>6.28 ± 0.42b</td>
<td>31.62 ± 2.08b</td>
<td>7.33 ± 0.30b</td>
</tr>
</tbody>
</table>

Different letters indicate significant differences by Tukey test (P ≤0.05). Values are means ± standard error (n=5).

Water stress, either by deficiency or excess of water, have provoked leaf dehydration, reduction in growth rate and leaf biomass yield (Else et al., 2001; Liu Stützel, 2004). This study has verified that the effect of stresses on carbon assimilation have caused a more severe reduction in leaf biomass accumulation than in the leaf area. Such response, in turn, was evidenced by a higher specific leaf area in the treatments (drought and flooding) than in the control. All plants of *J. curcas* survived to 16 days that were subjected to stress, despite decreased biomass production was observed. As described by Achten et al. (2010), the scarcity of water in the soil alters the pattern of biomass allocation of *J. curcas*, favoring the development of root and thereby elevating R/S values. The increase in R/S enhances the ability of this species in collecting water. Also, the reduction of $g_s$ explains the increase in WUE values.

The absence of damage to the root system of *J. curcas* plants subjected to water deficit is an interesting trait for breeding, since it improves the effective use of water, besides influencing the plant’s tolerance to stress and preventing reduced productivity (Blum, 2009). Furthermore, these characters influence the acclimation of this species to various locations, including areas with limited water availability (Srivastava et al., 2011; Fini et al., 2013). Leaf senescence and roots decay (Fig 5) were the key factors leading to increase in stem dry biomass ratio (63% higher than in the control) in *J. curcas* plants subjected to flooding stress, since this increase was not followed by an increase in stem dry biomass (SDB). Negative effects of flooding on the leaf (62% reduction) and root (38% reduction; in the stem, the reduction was 28%) biomass was also observed in *J. curcas* by Gimeno et al. (2012). According to these authors, this species can be considered sensitive to flooding for about 10 days, with a reduction of 30% of their total biomass. Such sensitivity has been confirmed in this study, in which *J. curcas* plants, to 16 DAIT, showed reductions of approximately 80%.
Table 2. Per plant (LA – cm²) and specific (SLA – cm²·g⁻¹) leaf area in plants of *Jatropha curcas* under different water conditions.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>LA</th>
<th>SLA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>1516.4 (87.3)a</td>
<td>217.4 (4.2)c</td>
</tr>
<tr>
<td>Drought</td>
<td>791.8 (35.0)b</td>
<td>271.0 (10.0)b</td>
</tr>
<tr>
<td>Flooding</td>
<td>244.7 (41.3)c</td>
<td>328.6 (4.0)a</td>
</tr>
</tbody>
</table>

Different letters indicate significant differences by Tukey test (P<0.05). Values are means (standard error) of 5 replicates.

Fig 2. Net photosynthetic rate (A) and transpiration (E) as a function of stomatal conductance (gs), in plants of *Jatropha curcas* under drought (a,c) and flooding (b,d). The points represent the average values of measurements (n=5).

Table 3. Mean values, maximum and minimum of photosynthetically active radiation (PAR – mol fotons m⁻²·day⁻¹), air temperature (Tair - °C) and relative humidity (RH - %), during the experimental period.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Maximum</th>
<th>Minimum</th>
<th>Mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>PAR</td>
<td>36.4</td>
<td>10.1</td>
<td>28.3</td>
</tr>
<tr>
<td>Tair</td>
<td>27</td>
<td>21</td>
<td>23.1</td>
</tr>
<tr>
<td>RH</td>
<td>94</td>
<td>58</td>
<td>73</td>
</tr>
</tbody>
</table>

Fig 3. Intrinsic water use efficiency (WUE) in plants of *Jatropha curcas* under different water conditions. Different letters indicate significant difference, by Tukey test (P≤0.05), between water regimes. The points represent the mean (n=5) and the bars indicate the standard error.
Fig 4. Instantaneous Carboxylation efficiency \( (A/Ci) \) in plants of *Jatropha curcas* under different water conditions. Different letters indicate significant difference, by Tukey test \((P \leq 0.05)\), between water regimes. The points represent the mean \((n=5)\) and the bars indicate the standard error.

Fig 5. A) Leaf, root, stem and total dry biomass; B) leaf, root, stem dry biomass ratio and root/shoot ratio \((R/S)\) in plants of *Jatropha curcas* under different water conditions. Columns are means \((n=5)\) and bars are the standard error.
From these results, it is possible to conclude that flooding has adversely affected leaf gas exchange and biomass production in J. curcas. Under water deficit, a high R/S associated with the lower responsiveness of the root system and the effectiveness of stomatal control as strategies for increasing water use efficiency under this stress and surviving to it. Hence, we conclude that these water stresses have exerted a negative effect on J. curcas through reductions in gas exchange and biomass production, with sharper decreases in plants subjected to flooding stress.

Materials and Methods

Plant material and experimental conditions

A greenhouse experiment was conducted at the campus of State University of Santa Cruz (UESC), in Ilhéus, Bahia, Brazil. J. curcas seeds, natural of Garanhuns-Pernambuco and donated by Brazilian Agricultural Research Corporation (EMBRAPA), Campina Grande, Paraíba, were germinated in 20 L pots (six seeds per pot) containing a mixture of soil:sand (2:1 v/v). The substrate obtained by the mixture soil:sand showed sandy loam texture and was previously fertilized according to chemical analysis of the substrate. After 15 days of germination, thinning was performed leaving only one plant per pot, constituting the experimental unit. Fifteen days after thinning, the plants were subjected to different water conditions for a period of 21 days. A set of 15 plants were used, five of which were subjected to water deficit, five to flooding and five remained irrigated to pot capacity. The drought treatment consisted of witholding water, whereas flooding condition was obtained by placing the pots into plastic bags, thus preventing drainage, and maintaining the water table above ground level. During the experiment, the photosynthetically active radiation (PAR) was monitored by means of quantum sensors S-LIA-M003, air temperature (Tair) and relative humidity (RH) were monitored using sensors microprocessor Hobo H8 Pro Series (Onset, USA). These variables were measured and stored permanently by data collectors Hobo Micro Station Data Logger (Onset, USA), where the values are shown in table 1. The soil moisture of control and drought plants was determined by the gravimetric method.

Leaf gas exchange

Leaf gas exchange was measured at 0, 2, 7, 9, 14 and 16 days after the imposition of treatments (DAIT) in one fully mature leaf per plant, using a portable photosynthesis system (LI-6400, Li-Cor, Lincoln, NE, USA). Measurements were performed from 8:30 a.m. to 11:30 a.m. under artificial saturating light of 1000 μmol photons m⁻² s⁻¹ and atmospheric CO₂ concentration (Ca) of 380 μmol mol⁻¹. Water use efficiency (WUE) was calculated as the ratio between net photosynthetic rate (A) and stomatal conductance (gs).

Chlorophyll content

Chloroplastidic pigment content was determined using a portable chlorophyll meter SPAD-502 (Soil Plant Analysis Development, Minolta, Japan), 16 DAIT, in fully mature leaves. From the SPAD index values, we have calculated the concentrations of chlorophyll a and b and carotenoids, according to Rigon et al. (2013) for J. curcas.

Biomass

At the end of the experiment, all the plants were harvested and sectioned in their leaves, stems and roots for dry biomass (DB) analysis. From these values, dry biomass ratio (DBR) of leaf, stem, and root, as well as the root:shoot ratio (R/S) have been calculated. Leaf area (LA) was measured using a leaf area meter (LI-3100, Li-Cor, Lincoln, NE, USA). Specific leaf area was calculated as SLA = LA / DB of leaf.

Experimental design and statistical analysis

A completely randomized experimental design was set, with three treatments (control, flooding and drought) and five replicates for each treatment, totaling 15 plants. The data were subjected to analysis of variance and the means were compared by Tukey test at 5% significance.

Conclusions

From these results, it is possible to conclude that flooding has a more pronounced negative effect than water deficit on leaf gas exchange and biomass production in J. curcas. Under water deficit, high R/S associated with lower responsiveness of the root system and the effectiveness of stomatal control, can be highlighted as strategies for increasing water use efficiency under this stress and surviving to it. Hence, we conclude that these water stresses have exerted a negative effect on J. curcas through reductions in gas exchange and biomass production, with sharper decreases in plants subjected to flooding stress.

Acknowledgments

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References

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