Responses of six *Brachiaria* spp. accessions to root zone flooding

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ABSTRACT - Some morphophysiological responses of *Brachiaria brizantha* cv. Marandu, cv. Piatã, cv. Arapoty, B163, B166 and *B. ruziziensis* R124 were compared in plants grown in pots, under flooded and well-drained conditions. Flooding reduced leaf elongation rate in all accessions. Leaf dry mass production was lower under flooding than in the control plants in all the studied accessions, except in cv. Piatã. Root dry mass production was reduced by flooding in cv. Marandu and in R124, while the proportion of biomass allocated to roots was reduced by flooding only in R124. It was not possible to detect significant differences between water regimes in the percentage of biomass allocated to culms and leaves for all accessions. Flooding decreased the number of tillers in cv. Marandu, in B163, B166 and in R124. Relative growth rate was decreased by flooding in cv. Marandu, in B163 and in R124. Flooding reduced gas exchange parameters in all accessions, but cv. Arapoty. The accessions tested differ in their relative tolerance to flooding. *B. brizantha* cv. Arapoty is considered the most tolerant.

Key Words: biomass allocation, *Brachiaria brizantha*, leaf elongation rate, photosynthesis, relative growth rate

Introduction

In tropical pastures periods of excess soil water can be relatively common events. Usually, excess water in pasture soils (waterlogging or flooding) is a short-term event and can be caused by the combination of intensive rains and low soil drainage (e.g., soils with a drainage impediment layer). In addition, management practices that lead to soil compaction (e.g., overgrazing) can disrupt the natural drainage ability of these soils, making them more prone to intermittent flooding or waterlogging periods.

Among the problems related to temporary excess soil water in pastures of tropical America, marandugrass (*Brachiaria brizantha* cv. Marandu) death syndrome is of particular importance (Dias-Filho, 2006). This problem was initially reported in Central America (Zúñiga et al., 1998) and for the last few years has become a major cause of pasture degradation in the North of Brazil, particularly in the states of Acre, Amazonas, Pará, Rondônia, Mato Grosso, Tocantins and Maranhão, where over 40% of the total pasture areas and 35% of the Brazilian beef cattle heard are present (Dias-Filho & Andrade, 2005). This syndrome is thought to be...
caused by the very low tolerance of marandugrass to excess soil water which adversely affects plant metabolism, resulting in predisposition to biotic stresses like fungal infection (Dias-Filho, 2005, 2006). Similar response mechanisms have also been reported in the literature for other plant species subjected to soil flooding, such as Eucalyptus marginata (Burgess et al., 1998, 1999), tomato (Chérif et al., 1997), and oak (Robin et al., 2001).

So far, the only economically viable alternative to cope with the death syndrome in areas prone to this problem is the replacement of marandugrass by grass species or cultivars with greater tolerance to temporary flooding or waterlogging (Andrade & Valentim, 2006; Dias-Filho, 2005, Dias-Filho & Andrade, 2005). Thus, information on the relative tolerance of forage grasses to excess soil water is essential to deal with this problem. For that reason, research programs to release new forage grass cultivars, such as the one currently being conducted for Brachiaria by the Brazilian Agricultural Research Corporation (Embrapa) (Valle et al., 2001), should prioritize the screening for flood tolerance in the evaluation process of these accessions.

Although knowledge about the tolerance of tropical forage grasses to excess soil water is key in determining the potential of use of these plants in habitats subjected to this stress, studies investigating this tolerance and the mechanisms involved in the responses of these plants to water stress have received little attention throughout the tropics (e.g., Baruch, 1994a; 1994b; Kibbler & Bahnisch, 1999; Medina & Motta, 1990; Ram, 2000), and particularly in Brazil (e.g., Dias-Filho, 2002; Dias-Filho & Carvalho, 2000; Haddade et al., 2002; Mattos et al., 2005).

We hypothesized that Brachiaria spp. accessions show differential ability to tolerate soil flooding. The objective of the present study was, therefore, to evaluate and compare selected physiological and morphological responses of five accessions of Brachiaria brizantha (including marandugrass) and one of B. ruziziensis under excess soil water.

**Material and Methods**

Seeds of Brachiaria brizantha cv. Marandu, cv. Piatã, cv. Arapoty, B163 and B166 and B. ruziziensis (R124), produced from germplasm originally introduced from east Africa in 1984 and 1985 (Valle, 1990) and supplied by Embrapa Beef Cattle, in Campo Grande, MS, Brazil, were germinated in sand and then planted individually in pots with 2 kg (dry mass) of soil (3:2; organic soil to cured manure). Seventeen days after planting, pots were fertilized with a solution of 40 mg of P (Ca(H2PO4)2.H2O) kg⁻¹ soil. One day after this fertilization, 50 mg N (urea) kg⁻¹ soil was applied. Plants were grown outdoors for the duration of the experiment under a shade net that intercepted ca. 60% of direct solar radiation. The trial was conducted at the Embrapa Eastern Amazonia headquarters, in Belém (1°28'S), PA, Brazil. Each pot was watered daily and fertilized every other day with 10 mL of a water soluble fertilizer solution (15:30:15; N:P2O5:K2O; 10 g L⁻¹). Flooding was imposed 20 days after planting by inundating the pots up to 3 cm above the soil level. Control pots were free-draining and watered daily. Flooding lasted five days for all accessions. Previous studies (Dias-Filho, 2002; M.B. Dias-Filho, unpublished data) have indicated that this period was ideal for detecting differences among forage grasses genotypes in their relative ability to tolerate flooding. All plants remained vegetative during the experimental period.

The length of a young expanding leaf blade (with the ligule not yet exposed) in a vegetative tiller of each plant was measured with a ruler at around the same time every day, starting immediately after flooding was imposed. Leaves being measured were marked with a plastic ring. Once the ligule was exposed, a new leaf, on the same plant, was marked and measured. Daily leaf elongation was calculated as the difference between leaf lengths of two consecutive days.

All plants were harvested at the end of the experiment; the number of tillers was counted for each plant and plant material was divided into leaf blades, culms (sheath and stem) and roots. Roots were washed free of soil using a manually manipulated jet spray of water. Plant dry mass was obtained by drying the plant material at 65°C for 48h. Leaf, culm and root mass ratios (respectively, leaf, culm and root dry mass per unit of dry mass of whole plant, LMR, CMR and RMR) were calculated according to Hunt (1990). Relative growth rate (change in total dry mass per total dry mass of plant per day, RGR) was also calculated.

Net photosynthesis, stomatal conductance and transpiration were measured with an open flow infrared gas analyzer with an attached red LED light source (LI-6400, Li-Cor, Inc., Lincoln, NE, EUA). Measurements were made between 9:00 a.m. and 11:00 a.m., local time, on the second and fourth days of flooding, on one young, fully expanded leaf blade of a vegetative tiller of each plant under a shade net that intercepted ca. 60% of direct solar radiation. The trial was conducted at the Embrapa Eastern Amazonia headquarters, in Belém (1°28’S), PA, Brazil. Each pot was watered daily and fertilized every other day with 10 mL of a water soluble fertilizer solution (15:30:15; N:P2O5:K2O; 10 g L⁻¹). Flooding was imposed 20 days after planting by inundating the pots up to 3 cm above the soil level. Control pots were free-draining and watered daily. Flooding lasted five days for all accessions. Previous studies (Dias-Filho, 2002; M.B. Dias-Filho, unpublished data) have indicated that this period was ideal for detecting differences among forage grasses genotypes in their relative ability to tolerate flooding. All plants remained vegetative during the experimental period.

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Pots were arranged in a factorial arrangement within a completely randomized design with four replicates. Differences in dry mass production and allocation, net
photosynthesis, stomatal conductance, transpiration and relative growth rate were assessed by two-way analyses of variance (ANOVA) with treatments (control and flooding) and accessions as main effects. Differences in leaf elongation rate throughout the experimental period were assessed by three-way ANOVA with treatments, accessions and evaluation dates as main effects. The assumption of homogeneity of variances and normality were tested for each ANOVA and, when necessary, data were log transformed. Transformed values were back transformed for presentation. Post hoc contrasts were calculated for assessing differences between treatments within the same accession whenever appropriate.

Factor analysis was used with the objective of grouping together (i.e., to classify) those accessions having the greatest similarities in their relative tolerance to flooding. Data for factor analysis were the mean difference between control and flooded plants in leaf elongation rate, leaf and root dry mass production, total dry mass production, relative growth rate, number of tillers, net photosynthesis, stomatal conductance and transpiration. The rationale being: the lower the difference between control and flooded plants, the higher the relative tolerance to flooding of that particular accession. All data were standardized to a mean of 0 and a standard deviation of 1 before conducting the analysis. The results of this analysis were presented as a two-dimensional plot of unrotated factor loadings.

The statistical package STATISTICA for Windows release 6.1 (StatSoft, Inc., Tulsa, USA) was used for all computations of the data.

Results and Discussion

Soil flooding reduced mean leaf elongation rate of all the studied accessions ($F_{5,211} = 3.92; P = 0.002$). The mean reduction in leaf elongation rate throughout the evaluation period was 48.6% for marandugrass, 56.3% for cv. Piatã, 35.6% for cv. Arapoty, 28.3% for B163, 48.5% for B166 and 40.9% for R124 (Figure 1). Similar results were reported by Dias-Filho (2002) and Mattos et al. (2005) studying the morphological responses of Brachiaria species under soil flooding.

According to Dias-Filho & Carvalho (2000), in Brachiaria spp. leaf elongation rate is related to the tolerance to flooding and could be used as an early detection mechanism of relative flood tolerance in Brachiaria spp. In the present study all accessions showed an immediate (day one) decrease in leaf elongation rate due to flooding.

Figure 1 - Leaf elongation rate of Brachiaria brizantha cultivar Piatã (A), Arapoty (B), B163 (C), B166 (D), cultivar Marandu (E) and B. ruziziensis R124 (F), under well-drained (open symbols, solid lines) and flooding (closed symbols, dotted lines) soil. Values are means±S.E.
indicating a low overall tolerance of these accessions to this stress. However, the reduction magnitude varied almost two-fold among all accessions, suggesting distinct degrees of relative flood tolerance.

A significant accession x treatment interaction was detected for relative growth rate (RGR) ($F_{5,25}=6.76; P<0.001$) (Figure 2). Post hoc contrasts of differences between treatments, within the same accession, showed that R124 ($F_{1,25}=59.1; P<0.0001$), marandugrass ($F_{1,25}=7.16; P=0.013$) and B163 ($F_{1,25}=9.73; P=0.0045$) suffered significant reductions in RGR (respectively 70.2%, 66.1% and 54.7%) due to flooding. No differences between treatments could be detected for cv. Piatã, Arapoty and B166. However, a clear tendency for lower RGR values were always observed in flooded plants.

Flooding reduced leaf biomass production for most of the accessions tested ($F_{5,25}=3.39; P=0.018$) (Figure 3). Only in cultivar Piatã it was not possible to detect statistical differences between treatments for leaf biomass production ($F_{1,25}=0.76; P=0.39$). Mean reduction in leaf production due to soil flooding was maximal in R124 (64.2%) and in marandugrass (62%).

Root dry mass was significantly reduced by flooding in R124 (82%) ($F_{1,25}=31.07; P<0.001$) and in marandugrass (74%) ($F_{1,25}=7.59; P<0.01$). For all the other accessions, although there was a tendency for lower root dry mass values under flooding, no statistical differences could be detected between flooded and control plants (Figure 4).

Flooding reduced the mean number of tillers in B166 (100%) ($F_{1,25}=16.4; P<0.001$), R124 (74.2%) ($F_{1,25}=177.5; P<0.0001$), marandugrass (55.5%) ($F_{1,25}=8.39; P<0.008$) and B163 (54.9%) ($F_{1,25}=11.6; P<0.001$). However, in cv. Arapoty and cv. Piatã it was not possible to detect significant differences between treatments in the mean number of tillers (data not shown). For most accessions tillering behavior of flooded plants was highly correlated to RGR and leaf biomass production (data not shown); the exception was B166 in which the decline in tillering did not affect those parameters.

For most of the accessions biomass allocation patterns were not significantly affected by flooding (Table 1). For R124, however, the amount of biomass allocated to roots (RMR) was significantly lower ($F_{1,25}=7.47; P=0.01$) in flooded plants. For the others accessions RMR generally showed a stronger tendency to be lower under flooding. An opposite tendency was observed for biomass allocated to leaves (LMR), while no clear pattern could be observed for biomass allocated to culms (CMR).

Miller & Zedler (2003) reported that the grasses *Phalaris arundinacea* and *Spartina pectinata* allocated more biomass to aboveground with flooding. Similarly, Rubio & Lavado (1999) observed a sharp decrease in the root:shoot ratio in the grasses *Paspalum dilatatum* and *Danthonia montevidensis* under flooding. Dias-Filho & Carvalho (2000) and Dias-Filho (2002) also found a decrease in RMR and an increase in LMR in flooded plants.
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Brachiaria spp. Also, a higher proportion of biomass allocated to leaves was observed in flooded *Leontodon taraxacoides* (Asteraceae) by Grimoldi et al. (1999). In contrast, Mattos et al. (2005) reported that flooding caused a lower LMR in *Brachiaria* spp.

As suggested by Rubio & Lavado (1999), the reduced allocation to roots in flooded plants could be a strategy to decrease the higher maintenance respiration of roots, when compared to that of shoots. Also, because roots cannot satisfy their own oxygen requirements, a smaller root system could be more easily ventilated by the shoot under waterlogging conditions.

There was a significant accession x treatment interaction on net photosynthesis ($F_{5,274} = 18.92; P<0.0001$), stomatal conductance ($F_{5,274} = 29.97; P<0.0001$) and transpiration ($F_{5,274} = 12.67; P<0.0001$). Post hoc contrasts of differences between treatments, within the same accession, revealed that only in cultivar Arapoty net photosynthesis was not significantly reduced by flooding (Figure 5). This reduction, however, was highest for R124 (63%), followed by marandugrass (43%) and by cv. Piatã (39%). A similar trend was observed for stomatal conductance (Figure 5) and transpiration (Figure 5). As with net photosynthesis, R124, marandugrass and cv. Piatã experienced the highest percentage decline due to flooding in leaf conductance (67, 51 and 44%, respectively) and transpiration (56, 34 and 32%, respectively), while no effect could be detected in cv. Arapoty. Previous studies have also shown a sharp decline of net photosynthesis and stomatal conductance of marandugrass subjected to root zone flooding (Dias-Filho, 2002; Dias-Filho & Carvalho, 2000). Interestingly, Mattos et al. (2005) found that net photosynthesis and stomatal conductance were not affected by flooding in marandugrass. The explanation could be the relatively low photosynthetic performance (9 $\mu$mol m$^{-2}$ s$^{-1}$) of control (field capacity) plants observed in that study, which could be viewed as an indication that those plants probably lacked the proper conditions (environmental or physiological) to express their full photosynthetic potential.

There was a significant and positive linear relationship between net photosynthesis and stomatal conductance for flooded plants from all accessions except for cultivar Arapoty, which showed a negative relationship between those parameters ($r^2 = 0.86; P<0.0001$). The positive relationship was strongest for cultivar Piatã ($r^2 = 0.99; P<0.0001$), but weaker for cultivar Marandu ($r^2 = 0.65; P<0.0001$). The strong positive relationship between net photosynthesis and stomatal conductance suggests that stomatal inhibition was a major factor affecting CO$_2$ assimilation. However, the
reduction in net photosynthesis observed in flooded plants was not always accompanied by a reduction in the internal CO₂ concentration (data not shown), which indicates that, in the present study, stomatal aperture, although important, was not the only limiting factor for the decrease in the photosynthetic capacity of flooded plants (Liao & Lin, 2001).

Conclusions

*B. brizantha* cv. Arapoty is the accession relatively most tolerant to flooding. *B. brizantha* cv. Piatã, B163 and B166 are intermediate in flooding tolerance. *B. ruziziensis* (R124), followed by marandugrass, are the most flood-sensitive accessions.

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