

*Journal of Experimental Agriculture International*

*44(11): 35-46, 2022; Article no.JEAI.89997 ISSN: 2457-0591 (Past name: American Journal of Experimental Agriculture, Past ISSN: 2231-0606)*

# **Gas Exchange of Oiticica Seedlings in Response to Potassium Fertilization and Water Turning**

**Rosilvam Ramos de Sousa a\* and Antonio Lucineudo de Oliveira Freire <sup>a</sup>**

*<sup>a</sup>Departamento de Ciências Florestais, Universidade Federal de Campina Grande, Brazil.*

### *Authors' contributions*

*This work was carried out in collaboration between both authors. Author RRS did the conceptualization, data curation, formal analysis, research, methodology, writing – first writing. Author ALOF did the conceptualization, data curation, obtaining funding, research, methodology, validation, writing – review and editing, supervision, project management and resources. Both authors read and approved the final manuscript.*

### *Article Information*

DOI: 10.9734/JEAI/2022/v44i1130906

#### **Open Peer Review History:**

This journal follows the Advanced Open Peer Review policy. Identity of the Reviewers, Editor(s) and additional Reviewers, peer review comments, different versions of the manuscript, comments of the editors, etc are available here: https://www.sdiarticle5.com/review-history/89997

*Original Research Article*

*Received 04 June 2022 Accepted 08 August 2022 Published 23 September 2022*

## **ABSTRACT**

**Aims:** This objective of this work was to evaluate the effects of water deficiency on physiological aspects of oiticica plants, as well as the capacity of potassium fertilization to attenuate such effects. **Study Design:** The treatments were distributed in an entirely randomized design (*DIC*), consisting of a 4x4 factorial, with four levels of potassium (K) and four water turning, with five repetitions. **Place and Duration of Study:** The experiment was conducted in a shaded environment, with 50% light interception, in the Forest Nursery of UAEF/UFCG, between January 2020 and July 2020. **Methodology:** The relative water content (*RWC*), transpiration (*E*) and photosynthesis (*A*) rates, stomatal conductance  $(gs)$ , and internal  $CO<sub>2</sub>$  concentration  $(Ci)$  were evaluated, and from the values of *A* and *E*, the intrinsic water use efficiency (*EUAi*) was calculated and, with the data of *A* and *Ci*, the carboxylation efficiency (*A*/*Ci*).

**Results:** There was significant interaction of treatments in the parameters *RWC*, *E*, *gs*, *A* and *EUAi* and isolated effect of water turning on *Ci* and *A*/*Ci*. Irrigation daily or every five days provided the maintenance of higher water status and better stomatal behavior, and the plants were sensitive to water stress provided by longer water turnings (every 10 and 15 days). The fertilization with 100 and 150 mg dm<sup>-3</sup> of K kept the values of *E* and A high, even under low water availability.

**Conclusion:** Potassium fertilization proved to be effective in attenuating the negative effects caused by water deficiency on gas exchange and *WUEi* of the seedlings, with highlights for 100 and 150 mg dm $^{-3}$  of K levels.

\_

*Keywords: Caatinga; forest essence; seedlings production; water stress.*

## **1. INTRODUCTION**

In recent decades, there has been growing concern about the effects of human activities on the increase in average global temperatures, especially in more sensitive areas, such as regions with arid and semi-arid climates. Regarding climate change, projections regarding the Brazilian semiarid region are worrying because, in addition to the increase in temperature, there should be an increase in the frequency, as well as in the duration, of dry periods [1,2], placing the Caatinga as one of the most vulnerable ecological regions on the planet [3].

In general, the species of the Caatinga present morphological modifications that guarantee their survival under the severe environmental conditions common to this biome, such as the presence of xylopodia in the roots, cut leaves, leathery leaves, etc. [4].

When subjected to water deficit conditions, plants promote stomatal closure, directly compromising processes such as stomatal conductance, transpiration and photosynthesis, affecting their growth [5,6,7]. However, the control on stomatal opening can be intensified by the use of potassium, which can increase stomatal conductance [8], thus attenuating the adverse effects of water stress.

The potassium, even though it is not part of any organic compound, nor does it play a structural role in plants, it is considered an essential element for plants [9]. One of the reasons for the high potassium requirements is probably the need to maintain high concentrations in the cytoplasm to ensure maximum enzymatic activity [10], in addition to functioning in the balance of negative charges and cellular pH [11] and in the maintenance of cell turgidity [12].

The oiticica (*Microdesmia rigida* (Benth.) Sothers & Prance), of the Crysobalanaceae family, is a large species, native and endemic to Brazil, found on the banks of rivers and streams in the agreste and northeastern sertão [13]. It is widely used in folk medicine, especially the leaves, used in the treatment of inflammation and diabetes. In the middle of the last century, the species was extensively exploited for the extraction of oil from its seeds as a raw material for the manufacture of soap and, more recently, for the production of biodiesel [14].

Despite the economic and ecological importance of oiticica, there are few studies that seek to understand the physiological strategies used by this species to survive in a challenging environment, such as the northeastern semi-arid region, which has only three months of rain, followed by nine months of drought, all of which the years. Given this fact, it is essential to understand its behavior under unfavorable water conditions, as well as to present strategies that can mitigate these effects, thus increasing the efficiency of plant water use, enabling the use of quality seedlings in forest stands, capable of surviving to adverse conditions after planting.

Such information can support management strategies for this species in degraded areas, allowing, in addition to revegetation, the infiltration of water into the soil, due to its deep root system, breaking the naturally compacted upper layer, characteristic of Brazilian semiarid soils, facilitating the establishment of other species in the area.

Thus, this study aimed to evaluate the effects of water deficit on physiological aspects of oiticica plants, as well as the ability of potassium fertilization to attenuate these effects.

## **2. MATERIALS AND METHODS**

The experiment was carried out in a screened environment, with 50% light interception, in the Forestry Nursery of the Forest Engineering Academic Unit, belonging to the Rural Health and Technology Center of the Universidade Federal de Campina Grande, located in the municipality of Patos - PB (07º03' 39'' S and 37º16'38'' W), with an average altitude of 250 m. The climate of the region is classified, according to Köppen, as BSh, semi-arid, with average annual temperatures above 25°C and annual average rainfall around 750 mm, with irregular rainfall distribution and average relative humidity of 65% [15].

The oiticica seeds, from 5 matrices present in the community of the District of Socorro, in the rural area of the municipality of Olho d'Água - PB, were placed in a seedbed containing sand, and, five days after emergence (*DAE*), the seeds were seedlings were transferred to containers (black plastic bags) containing 5 kg of a mixture of subsoil soil and cattle manure, proportion 2:1. Before filling the containers, the substrate was fertilized with N, P and K. Nitrogen and phosphorus were added as recommended by Furtini Neto et al. [16], and fertilization with potassium was carried out according to the treatments tested.

The soil used in the experiment was collected at Fazenda Nupeárido, belonging to UFCG, crushed and sieved and subjected to physical and fertility analysis at the Laboratory of Soil and Water Analysis of UFCG (Table 1).

During the experimental period, the irrigation of the pots was done trying to keep the humidity close to 80% of the substrate retention capacity, obtained through weighing. To determine this retention capacity, five pots were submerged in a bucket with water until the substrate was completely soaked. Then, they were removed and, after complete leaching, weighed. This value corresponded to their weight with 100% of the vessel capacity (cv) and, based on it, the weight it should have when reaching 80% of the holding capacity was calculated.

The treatments were distributed in a completely randomized design (DIC), consisting of a 4 x 4 factorial, with four levels of potassium (K) (0, 50,100 and 150 mg dm<sup>-3</sup> K) and four water turning (ID - daily irrigation (control), and every 5 (5D), 10 (10D) and 15 days (15D)), with five replications and two plants per sampling unit, totaling 160 plants.

Water regimes were started at 25 days after seedlings were transferred to pots (plants at 30 days of age). On that day, the height of the plants (initial height) was measured. This measurement was also performed at the end of the experiment, which occurred 90 days after the start of water treatments [120 days after emergence (DAE)].

### **2.1 Evaluated Parameters**

a) Relative water content (RWC): at the end of the experiment, two leaves per plant were collected, taken to the Laboratory of Plant Physiology (CSTR), from which 5 disks of 1  $cm<sup>2</sup>$ per leaf were removed and subjected to weighing, obtaining if the fresh mass (FM). Then, they were placed under two sheets of hydrated germination paper, in Petri dishes, and placed in a refrigerator (5°C) until complete hydration. Upon reaching this point, they were removed, lightly dried and weighed to determine the turgid mass (TM). Subsequently, they were placed in an oven for drying (±65 oC), until reaching constant mass, to determine the dry mass (DM). The relative water content was calculated using the Weatherley [17] formula:

$$
RWC = \left(\frac{FM - DM}{TM - DM}\right) \times 100\tag{1}
$$

b) Stomatal evaluations: at the end of the experiment, readings of transpiration (*E*) and photosynthesis (*A*), stomatal conductance (*gs*) and internal concentration of CO<sub>2</sub> (Ci) were performed with the aid of a portable LCpro-SD photosynthesis (ADC BioScientific Ltd.). These readings were taken on fully expanded leaves, two per plant, inserted at the third and fourth nodes from the apex of the plants, between 9 am and 10 am. From the values of A and gs, the intrinsic efficiency in water use (*EUAi*) was calculated and, with the data of *A* and *Ci*, the carboxylation efficiency (*A*/*Ci*).

The data were submitted to analysis of variance by the F test, and the means were compared by the Tukey test, at 5% of significance, using the statistical program *Assistet*® [18].





#### **3. RESULTS AND DISCUSSION**

There was a significant interaction of the treatments in the parameters *RWC*, *gs*, *E*, *A* and *EUAi* and isolated effect in the water turning in *Ci* and *A/Ci*.

Comparing the K doses in each water turning treatment (Table 2), it was verified that there was no significant effect of potassium fertilization on the *RWC* when the plants received water daily (ID) and every 5 days (5D). However, despite the water turning increasing to 10 (10D) and 15 days (15D), the use of 100 and 150 mg dm<sup>-3</sup> of K enabled higher *RWC*, compared to 0 and 50 mg  $dm^{-3}$  of K.

In the analysis of the *RWC* in the water regimes at each dose of K, there was statistical equality between treatments ID and 5D, which were superior to 10D and 15D, both at 0 and 50 mg  $dm^{-3}$  of K. 100 and 150 mg dm<sup>-3</sup> of K provided statistical equality between water turning.

It can be seen from the data contained in Table 2 that, even under irrigation every five days, the oiticica plants were able to maintain a high *RWC*, regardless of potassium supply, evidencing the species' resistance to water deficit. However, the positive effect of provisioning with this nutrient, specifically 100 and 150 mg dm $^3$  of K, became evident when the plants were irrigated at longer intervals (10D and 15D), which allowed the maintenance of high *RWC*. According to Prado (2010), *RWC* values below 50% are considered critical and can lead to tissue death, with the exception of some xerophilous species, which can withstand this value without tissue loss.

Reduction in *RWC* due to water deficit has been reported in other studies [19,20,21,22]. However, in aroeira (*Myracrodruon urundeuva* Fr. Allem) and jucá (*Libidibia ferrea* (Mart. ex Tul) under moderate and severe water deficits, Almeida et al. [22] found an increase in *RWC* when they supplied potassium to the plants. Presented by plants, when supplied with K, demonstrates its ability to provide drought tolerance, helping to<br>carry out photosynthesis and transport out photosynthesis and transport carbohydrates from leaves to roots [23], and the osmotic effect exerted on plants. Roots, contributing to water absorption and maintenance of high *RWC* [24].

Low *RWC* causes numerous damages to various physiological processes, from the arrest of cell growth, the closing of stomata and the reduction in photosynthesis, in addition to impairing basic metabolic processes in plants, such as the production and degradation of proteins and carbohydrates and the accumulation of solutes, which may compromise plant growth [25,26].

*RWC* value below the critical (46.5%) mentioned above was verified only in the plants of the 15D treatment, which did not receive the K supply, evidencing once again the role of this nutrient in the absorption and maintenance of tissue water in the species here studied. Potassium is the ion found in the highest concentration in the vacuole of cells, being closely related to osmotic balance, thus acting in the maintenance of cell turgidity [27,12]. The accumulation of osmotically active solutes in the cell is an important strategy developed by plants that tolerate drought, promoting a reduction in water potential [28].

As the dose of K supplied increased, there was a progressive decrease in stomatal conductance (*gs*) in all water turning evaluated (Table 3). Analyzing the water turning within the K doses provided, it was found that, in unfertilized plants  $(0 \text{ mg dm}^3$  of K), the highest gs values were recorded in plants under 5D and 10D. However, in plants fertilized with 50 mg dm<sup>-3</sup> of K, those maintained in water turning 10D and 15D stood out, and at 100 mg dm $^{-3}$  of K there was no significant difference between water turning. Providing 150 mg dm $^{-3}$  of K, plants under daily irrigation have lower stomatal conductance, statistically different from other water turning.

**Table 2. Relative water content (***RWC***) (%) of** *M. rigida* **plants, as a function of potassium fertilization and water turning**

<b>Water turning</b>	K (mg dm <sup>-3</sup> )				
		50	100	150	
ID	77.1 aA	73.3 aA	74.6 aA	78.9 aA	
5D	82.1 aA	74.2 aA	76.1 aA	75.5 aA	
10D	54.8 bB	65.4 bB	79.6 aA	74.8 aA	
15D	46.5 bC	54.8 bB	77.0 aA	77.1 aA	

*\*Means followed by equal letters, uppercase in the lines and lowercase in the columns, do not differ statistically from each other (P ≤0.05)*

<b>Water turning</b>	K (mg dm <sup>-3</sup> )				
		50	100	150	
ID	0.79 <sub>bA</sub>	0.37 <sub>bB</sub>	$0.26$ aB	$0.12$ bB	
5D	$1.08$ aA	0.43 <sub>bB</sub>	$0.18$ aC	$0.29$ aC	
10D	0.95aA	$0.57$ aB	$0.19$ aC	$0.24$ aC	
15D	0.67 <sub>bA</sub>	$0.53$ aA	$0.21$ aB	$0.25$ aB	

**Table 3. Stomatal conductance (***gs***) (mmol H2O dm-2 s -1 ) of** *M. rigida* **plants, as a function of potassium fertilization and water turning**

*\*Means followed by equal letters, uppercase in the lines and lowercase in the columns, do not differ statistically from each other (P ≤0.05)*

Potassium fertilization did not positively influence *gs*, regardless of the water turning, with an average reduction of 74% between the treatment without K and the one that received the highest dose (150 mg dm $^{-3}$ ). Similar results were obtained in *Mimosa tenuiflora*, when submitted to 100% and 25% of vessel capacity (cv), in which the gs reduced with the increase of the amount of K [22]. In contrast, Costa, Carvalho and Pinto [29] reported that the highest dose of potassium contributed to greater stomatal conductance, adding that it provided a greater balance between the turgor of the guard cells and the control of the osmotic potential of the plant.

Reduction in stomatal conductance is probably associated with the adaptation mechanism of the plant when exposed to water deficit conditions, since, under these circumstances, plants tend to close their stomata, causing a decrease in leaf transpiration and, consequently, in water loss [12]. The reduction of *gs* is a mechanism for evaluating the degree of water stress to which the seedling is exposed due to the sensitivity of the stomata to changes in water availability [30]. According to Pita et al. [31], this aspect can be used in genetic improvement, aiming at increasing productivity, even in drought conditions.

Regarding the transpiration rate (*E*) (Table 4), comparing the K treatments in each water turning, there was an increase in the values as the supplied dose increased, except in plants kept under daily irrigation (ID), which showed lower E when they received 150 mg dm<sup>-3</sup> of K. Comparing the water turning, at each dose of K, the highest transpiration rate occurred in plants irrigated every five days (5D), when they received 0. 50 and 100 mg dm $3$  of K, higher than water turning ID, 10D and 15D.

In a parallel between treatments 5D and 10D, reductions in  $E$  of 29% (0 mg dm<sup>-3</sup> of K), 47% (50 mg dm $^{-3}$  of K) and 24% (100 mg dm $^{-3}$  of K) were

observed. While in the 15D treatment, the reductions were, respectively, 32%, 46% and 25%. In plants fertilized with 150 mg dm<sup>-3</sup> of K, statistical equality was verified between the 5D, 10D and 15D water turning, which were statistically superior to the ID treatment. There is, therefore, a positive effect of potassium fertilization in the attenuation of water deficit on plant transpiration.

The reduction in water availability that occurred in plants irrigated at longer water turning (10D and 15D) had direct consequences on the *RWC* (Table 1) and on plant transpiration (Table 3). It can be seen that plants from treatments 10D and 15D, fertilized with K at doses of 0 and 50 mg dm-3 , showed lower values of *RWC* and, therefore, greater stomatal closure and lower *E*. According to Chaves, Flexas and Pinheiro [32], this stomatal closure resulting from water deficit is a strategy to reduce water loss through transpiration. With this, the plant guarantees the maintenance of an adequate state of hydration and, consequently, of activities vital to its growth. Stomatal closure due to water deficit was also reported by Albuquerque et al. [33], França et al. [34], Campelo et al. [35] and Almeida et al. [22], among others. The increase in water turning and consequent decrease in soil moisture content may have resulted in the production of abscisic acid (ABA) in the roots and its transport to the leaves, resulting in the closing of the stomata [36,37]. This, in turn, limits the loss of water by the plant, preventing tissue dehydration and consequent death of the plant [38,39].

The increase in plant transpiration with the increase in the dose of K supplied (100 and 150 mg dm<sup>-3</sup>) evidences the beneficial effect of potassium for plants, especially under water deficit conditions, through a possible cellular osmotic adjustment, ensuring greater water absorption and perspiration. Silva et al. [11] reinforced the role of potassium as an osmotic agent, ensuring the maintenance of water

<b>Water turning</b>	K (mg dm <sup>-3</sup> )				
		50	100	150	
ID	3.82 bA	3.92 bA	4.85 bA	3.27 bB	
5D	4.53 aB	5.88 aA	6.36 aA	5.42 aA	
10D	3.23 <sub>bB</sub>	3.11 <sub>bB</sub>	4.82 bA	5.00 aA	
15D	3.08 <sub>bB</sub>	3.16 <sub>bB</sub>	4.77 bA	5.75 aA	

**Table 4. Transpiration rate (***E***) (µmol H2O dm-2 s -1 ) of** *M. rigida* **plants, as a function of potassium fertilization and water turning**

*\*Means followed by equal letters, uppercase in the lines and lowercase in the columns, do not differ statistically from each other (P ≤0.05)*

absorption by plants. Subjecting acerola plants to saline stress, Dias et al. [40] reported that low transpiration was a direct reflection of stomatal behavior and that the use of potassium in plants promoted osmotic adjustment, ensuring greater water uptake and increases in transpiration and  $CO<sub>2</sub>$  assimilation rate.

Regarding the photosynthesis rate (*A*) (Table 5), there was a gradual reduction as the water turning increased, regardless of the amount of K supplied. This effect was more severe in plants that did not receive potassium, with *A* values reducing from 31.26  $\mu$ mol dm<sup>-2</sup> s<sup>-1</sup> (ID) to 5.08  $\mu$ mol dm<sup>-2</sup> s<sup>-1</sup> (15D), equivalent to 84% decrease, while in the other K treatments, there was an average decrease of 60%.

Comparing K doses, in each water turning, in plants irrigated daily (ID), no significant difference was detected between treatments. However, in the 5D water turning, there was an elevation in A when they received 150 mg dm<sup>-3</sup> of K and, in the 10D and 15D water turning, in which there was an elevation in *A* from 100 mg dm<sup>-3</sup> of K, with the most pronounced effect in plants irrigated every 15 days (15D), with a 66% increase in this variable when comparing plants that did not receive K (0 mg dm $^{-3}$ ) with those that received 150 mg dm $^{-3}$  of K.

The increase in the water turning imposed a water deficit condition on the plants, causing the stomata to close, with consequent decreases in transpiration (Table 4) and photosynthesis (Table 5) rates. Such a response is a common fact to

plants, and its intensity varies with the species and with the degree of water condition imposed.

For some authors, photosynthesis is directly impaired by the limitation of gas exchange that occurs in the stomata, since it reduces the availability of  $CO<sub>2</sub>$  to the enzyme ribulose-1,5bisphosphate carboxylase/oxygenase (Rubisco) [41,31,42,32,43,44], which consequently affects their activity [45]. However, this process may originate from other non-stomatal factors, such as the impairment of the photosynthetic system, through the disruption of thylakoid membranes [46], decrease in ATP synthesis, electron transport in photosystem II (PSII), the capacity and speed of regeneration of Rubisco [47,41,32] and the regeneration of RuBP [48].

Silveira et al. [49] point out that resistances related to water flow in the soil-plant-atmosphere system cause disharmony between water absorption and transpiration. These authors also emphasize that the partial closure of the stomata is a maneuver to prevent excessive dehydration or a leaf water imbalance, which, consequently, would lead to a reduction in the photosynthetic rate. Several studies relate the decrease in stomatal conductance and in transpiration and photosynthesis rates with soil water deficit in arboreal species of the caatinga, such as aroeira (*Myracrondruon urundeuva* Allemão) [50], ipêroxo (*Handroanthus impetiginosus* (Mart. ex DC.) Mattos) [51] and faveleira (*Cnidoscolus quercifolius* Pohl.) [52].

**Table 5. Photosynthesis (***A***) (µmol CO<sup>2</sup> dm-2 s -1 ) of** *M. rigida* **plants, as a function of potassium fertilization and water turning**

<b>Water turning</b>	K (mg dm <sup>-3</sup> )			
		50	100	150
ID	31.26 aA	25.68 aA	35.92 aA	37.99 aA
5D	17.74 bB	17.82 bB	20.91 bB	26.82 bA
10D	12.12 bB	10.88 <sub>bB</sub>	13.50 bA	14.62 cA
15D	$5.08 \text{ cC}$	10.80 bB	13.72 bA	14.75 cA

*\*Means followed by equal letters, uppercase in the lines and lowercase in the columns, do not differ statistically from each other (P ≤0.05)*

Despite the reduction in the photosynthetic rate. due to the increase in the water turning, it can be seen that this effect is attenuated by the use of potassium, especially at doses of 100 and 150 mg dm $3$  of K (Table 5). A positive effect of potassium in maintaining a high *RWC* and increasing stomatal conductance, transpiration and photosynthesis was also obtained by Almeida et al. [22] in *Myracrondruon urundeuva*, *Libidibia ferrea* and *Mimosa tenuiflora* under conditions of low water availability. These authors attributed these results to the osmotic action of this nutrient, providing cellular osmoregulation and greater water absorption by plants, corroborating the statements by Mendes et al. [53], Battie-Laclau et al. [8], Zahoor et al. [54] and Dias et al. [40].

Due to the decrease in *gs* (Table 3) and the increase in *A* (Table 5), the intrinsic water use efficiency (*EUAi*) (Table 6) decreased as the interval between water turning, at all levels of K evaluated. Comparing the ID and 15D treatments, the decreases were 81%, 71%, 53% and 81%, respectively, at the levels of 0, 50, 100 and 150 mg dm $^3$  of K.

In each water turning, there was an increase in *EUAi*, with the increase in the amount of K supplied to the plants (Table 5), and between treatments 0 and 150 mg dm<sup>-3</sup> of K, the values obtained in this variable were about 8, 5.6, 4.8 and 7.8 times, respectively, in the ID, 5D, 10D and 15D water turning.

Therefore, *EUAi* was strongly affected by water stress caused by water turning, regardless of potassium dose. Furthermore, it was found that the supply of K (100 and 150 mg dm<sup>-3</sup>) enabled the attenuation of the effects of eventual water deficit in the plants under the water turning evaluated. The values presented in Table 5 are mainly due to the positive effect exerted by K on photosynthesis *A* (Table 4) and negative effect on stomatal conductance (Table 2). This statement corroborates the studies by Tsonev et al. [55], who reported that the reduction in *gs* and not in *A* can explain the higher *EUAi* values of plants when submitted to water deficit. This is

**Table 6. Intrinsic efficiency in water use (***EUAi***) of** *M. rigida* **plants, as a function of potassium fertilization and water turning**

<b>Water turning</b>	K (mg dm <sup>-3</sup> )			
		50	100	150
ID	39.6 aD	69.7 aC	138.1 aB	316.6 aA
5D	16.4 bD	41.4 bC	116.2 bA	92.5 <sub>bB</sub>
10D	12.7 <sub>b</sub> C	$19.1 \text{ cB}$	71.0 cA	60.9 cA
15D	7.58 cC	$20.4 \text{ cB}$	65.3 dA	59.0 cA

*\*Means followed by equal letters, uppercase in the lines and lowercase in the columns, do not differ statistically from each other (P ≤0.05)*



**Fig. 1. Internal CO<sup>2</sup> (***Ci***) concentration of** *M. rigida* **leaves at 120 DAE, as a function of water turning**

*\*Means followed by equal letters do not differ statistically from each other (P ≤ 0.05)*



#### **Fig. 2. Carboxylation efficiency (***A***/***Ci***) of** *M. rigida* **leaves at 120 DAE, as a function of water turning**

*\*Means followed by equal letters do not differ statistically from each other (P ≤ 0.05)*

because, when under water stress, a small reduction in *gs* can have a protective effect against water stress, allowing plants to reduce water consumption and thus improve water use efficiency [32]. A negative effect of water deficit on water use efficiency and photosynthesis rate was also reported by Hommel et al. [56], when<br>evaluating the stomatal and mesophilic evaluating the stomatal and mesophilic conductances of forest species under conditions of low water availability.

According to Dias et al. [40], potassium is essential for enzyme activation, carbohydrate translocation and water regulation. Taiz et al. [12] emphasize that, as a result of gas exchange, during the absorption of carbon dioxide from the external environment, the plant loses water to the atmosphere and, in order to reduce this loss, it reduces the entry of  $CO<sub>2</sub>$ .

Figs. 1 and 2 show the data for *Ci* and *A*/*Ci*, respectively.

Regarding the internal concentration of  $CO<sub>2</sub>$ (*Ci*), the highest value was observed in plants irrigated every 5 days. However, there was statistical equality between this and treatments 10D and 15D (Fig. 2). The same behavior was verified in the carboxylation efficiency (*A*/*Ci*). Fernandes et al. [57,58] explain that, during gas exchange, as the stomata are responsible for regulating the substomatic concentration of CO2, the *Ci* remains slightly constant, a similar behavior observed in this research.

## **4. CONCLUSION**

Water stress impaired gas exchange in *M. rigida* seedlings. Daily irrigation or every five days provided the maintenance of higher water status and better stomatal behavior, and they were sensitive to water stress caused by longer water turning (10D and 15D). Potassium fertilization proved to be effective in mitigating the negative effects caused by water deficiency on gas exchange and *EUAi* of plants, especially at doses of 100 and 150 mg dm $^{-3}$  of K.

## **ACKNOWLEDGEMENTS**

We thank the Coordenação de Aperfeiçoamento de Pessoal de Nível Superior (CAPES) for the scholarship during the entire research period.

## **COMPETING INTERESTS**

Authors have declared that no competing interests exist.

#### **REFERENCES**

- 1. Lacerda FF, Nobre P, Sobral MdCM, Lopes GMB, Assad ED. Trends in the semiarid climate against the perspectives of global climate change; the case of Araripina, Pernambuco. Rev Dep Gegr. 2016;31:132-41. DOI: 10.11606/rdg.v31i0.114843
- 2. Marengo JA, Torres RR, Alves LM. Drought in Northeast Brazil: Past, present

and future. Theor Appl Climatol. 2017;129(3-4):1189-200. DOI: 10.1007/s00704-016-1840-8

- 3. Cavalcante AdMB, Fernandes PHC, Silva EMd. *Opuntia ficus-indica* (L.) Mill. and Climate Change: An Analysis in Light of Species Distribution Modeling in the Caatinga Biome. Rev Bras Meteorol. 2020;35(3):375-85 DOI: 10.1590/0102-7786353001.
- 4. Larcher W. Ecofisiologia vegetal. São Carlos: Rima. 2000;531.
- 5. Chaves MM, MAROCO JP, PEREIRA JS. Understanding plant responses to drought—from genes to the whole plant. Funct Plant Biol. 2003;30(3):239-64. DOI: 10.1071/FP02076, PMID 32689007.
- 6. Chaves MM, Oliveira MM. Mechanisms underlying plant resilience to water deficits: prospects for water-saving agriculture. J Exp Bot. 2004;55(407):2365-84. DOI: 10.1093/jxb/erh269, PMID 15475377.
- 7. Souza BCd, Oliveira RS, Araújo FSd, Lima ALAd, Rodal MJN. Functional divergences and drought resistance strategies among tropical deciduous and evergreen species. Rodríguesia. 2015;66(1):21-32 DOI: 10.1590/2175-7860201566102
- 8. Battie-Laclau P, Laclau JP, Beri C, Mietton L, Muniz MR, Arenque BC, et al. Photosynthetic and anatomical responses of Eucalyptus grandis leaves to potassium and sodium supply in a field experiment. Plant Cell Environ. 2014;37(1):70-81. DOI: 10.1111/pce.12131, PMID 23663049.
- 9. Malavolta E. Plant mineral nutrition manual. São Paulo: Agronomic Ceres. 2006;638.
- 10. Malavolta E, Vitti GC, Oliveira SA. Assessment of the nutritional status of plants: principles and applications. 2nd ed. Piracicaba: POTAFOS. 1997;105.
- 11. Silva PMC, Uchôa SCP, Barbosa JBF, Bastos VJ, Alves JMA, Farias LC. Effect of potassium and limestone on the quality of sweet cedar (Bombacopsis quinata) seedlings. AGRO@ENVIRONMENT. 2013;7(1):63-9.

DOI: 10.18227/1982-8470ragro.v7i1.842

- 12. Taiz L, et al. Plant physiology and development. 6. ed. Porto Alegre: Artmed. 2017;858.
- 13. Diniz NETO MA, Silva IdFd, Cavalcante LF, Diniz BLMT, Silva JCAd, Silva ECd. Mudas de oiticica irrigadas com águas salinas no solo com biofertilizante bovino e

potássio. Rev Bras Eng Agric Ambient. 2014;18(1):10-18.

DOI: 10.1590/S1415-43662014000100002

- 14. Souza F. Oiticica: Licania rigida Benth., fauna and flora of RN; 2018. Available:http://faunaefloradorn.blogspot.c om/2018/01/oiticica-licania-rigidabenth.html Acesso em: 12 jun. 2019.
- 15. Alvares CA, Stape JL, Sentelhas PC, de Moraes Gonçalves JL, Sparovek G. Köppen's climate classification map for Brazil. Meteorol Z. 2013;22(6):711-28. DOI: 10.1127/0941-2948/2013/0507
- 16. Furtini Neto AE, et al. Soil acidity, growth and mineral nutrition of some tree species in the seedling phase. Rev Cerne. 1999;5:1-12.
- 17. Weatherley PE. Studies in the water relations of the cotton plant I: The field measurements of water deficits in leaves. New Phytol. 1950;49(1):81-97. DOI: 10.1111/j.1469-8137.1950.tb05146.x
- 18. Silva FAS, Azevedo CAV. Version of the computer program Assistet for the Windows operating system. Rev Bras Product Agroind. 2002;4(1):71-8. DOI: 10.15871/1517-8595/rbpa.v4n1p71- 78.
- 19. Wang LF. Physiological and molecular responses to drought stress in rubber tree (*Hevea brasiliensis* Muell. Arg.). Plant Physiol Biochem. 2014;83:243-9. DOI: 10.1016/j.plaphy.2014.08.012, PMID 25194774.
- 20. Ataíde W, Neto C, Silva K, Brito A, Cardoso K, Nogueira G, et al. Growth and gas exchange of Tachigali vulgaris submitted to water deficiency. Rev Ciênc. Agr. 2018;41(3):771-82. DOI: 10.19084/RCA17231
- 21. Leite RdS, Nascimento MNd, Tanan TT, Ramos CAdS, Neto LPG, Guimarães DS. Physiological responses of Physalis angulata plants to water deficit. J Agric Sci. 2018;10(10):287-97. DOI: 10.5539/jas.v10n10p287
- 22. Almeida EP, Freire ALdO, Bakke IA, Ferreira CD, França GM, Batista AW. Effects of potassium in *Myracrodruon urundeuva*, *Libidibia ferrea* and *Mimosa tenuiflora* seedlings under a short-term water deficit. Res Soc Dev. 2020;9(5): 1-19.

DOI: 10.33448/rsd-v9i5.3269.

23. Martineau E, et al. The role of potassium on maize leaf carbon exportation under drought condition. Acta Physiol Plant. 2017;39:1-13.

- 24. Premachandra GS, Saneoka H, Fujita K, Ogata S. Leaf water relations, osmotic adjustment, cell membrane stability, epicuticular wax load and growth as affected by increasing water deficits in sorghum. J Exp Bot. 1992;43(12):1569-76. DOI: 10.1093/jxb/43.12.1569.
- 25. Kumar A, Singh DP. Use of physiological indices as a screening technique for drought to tolerance in oilseed Brassica species. Ann Bot. 1998;81:413-20.
- 26. Franco AC, Bustamante M, Caldas LS, Goldstein G, Meinzer FC, Kozovits AR, et al. Leaf functional traits of Neotropical savanna trees in relation to seasonal water deficit. Trees. 2005;19(3):326-35. DOI: 10.1007/s00468-004-0394-z
- 27. Kerbauy GB. Plant physiology. 2nd ed. Rio de Janeiro: Guanabara Koogan. 2013;431.
- 28. Chaves Filho JT, Stacciarini-Seraphin E. Change in osmotic potential and soluble carbohydrate content in young plants of Lobeira (*Solanum lycocarpum* St. Hil.) in response to water stress. Rev Bras Bot. 2001;24(2):199-204.

DOI: 10.1590/S0100-84042001000200010

- 29. Costa WRS, Carvalho PHMdS, Pinto PAdC. Potassium fertilization in protected pepper cultivation under organic system.<br>Rev Verde Agroecol Sustainable Rev Verde Agroecol Sustainable Development. 2019;14(2):224-8 DOI: 10.18378/rvads.v14i2.6134
- 30. Eksteen AB, Grzeskowiak V, Jones NB, Pammenter NW. Stomatal characteristics of *Eucaliptus grandis* clonal hybrids in response to water stress. South Forests J Forest Sci. 2013;75(3):105-11. DOI: 10.2989/20702620.2013.804310
- 31. Pita P, Cañas I, Soria F, Ruiz F, Toval G. Use of physiological traits in tree breeding for improved yield in drought-prone environments. The case of Eucalyptus globulus. Invest Agrar.: Sist Recur For. 2005;14(3):383-93. DOI: 10.5424/srf/2005143-00931
- 32. Chaves MM, Flexas J, Pinheiro C. Photosynthesis under drought and salt stress: regulation mechanisms from whole plant to cell. Ann Bot. 2009;103(4): 551-60.

DOI: 10.1093/aob/mcn125, PMID 18662937.

- 33. Albuquerque MPFd, Moraes FKC, Santos RIN, Castro GLSd, Ramos EMLS, Pinheiro HA. Ecofisiologia de plantas jovens de mogno-africano submetidas a déficit hídrico e reidratação. Pesq agropec bras. 2013;48(1):9-16. DOI: 10.1590/S0100-204X2013000100002
- 34. França PHTd, Silva ECAd, Silva TC, Brasil NA, Nogueira RJMC. Physiological analysis in guanandi seedlings (*Calophyllum brasiliense* Cambess.) submitted to water deficit. Agriculture In the Semi-ARID. 2017;13(4):264-269. DOI: 10.30969/acsa.v13i4.886
- 35. Campelo DdH, Lacerda CFd, Sousa JAd, Bezerra AME, Araújo JDM, Neves ALR, et al. Leaf morphophysiological characteristics and nutritional status of six woody species as a function of soil water availability. Science Forest. 2018; 28(3):924-36
- DOI: 10.5902/1980509832581 36. inácio MC, et al. Secondary messengers related to the action of plant hormones. Rev Bras Agroscience. 2011;17:
- 438-46. 37. Valadares J, Paula NFd, Paula RCd. Physiological changes in eucalyptus hybrids under different irrigation regimes. Rev Ciênc. Agron. 2014;  $45(4)$ :805-14

DOI: 10.1590/S1806-66902014000400019

- 38. Chaves MM, Pereira JS, Maroco J, Rodrigues ML, Ricardo CP, Osório ML et al. How plants cope with water stress in the field? Photosynthesis and growth. Ann Bot. 2002;89;Spec No:907-16. DOI: 10.1093/aob/mcf105, PMID 12102516
- 39. Chaves MM, Costa JM, Zarrouk O, Pinheiro C, Lopes CM, Pereira JS. Controlling stomatal aperture in semi-arid regions – the dilemma of saving water or being cool? Plant Sci. 2016; 251:54-64. DOI: 10.1016/j.plantsci.2016.06.015, PMID 27593463.
- 40. Dias AS, Lima GSD, Pinheiro FWA, Gheyi HR, Soares LADA. Gas exchange, quantum yield and photosynthetic pigments of acerola tree under saline stress and potassium fertilization. Rev Caatinga. 2019;32(2):429-39 DOI: 10.1590/1983-21252019v32n216rc
- 41. Flexas J, Bota J, Loreto F, Cornic G, Sharkey TD. Diffusive and metabolic

limitations to photosynthesis under drought and salinity in C3 plants. Plant Biol (Stuttg). 2004;6(3):269-279.

DOI: 10.1055/s-2004-820867, PMID 15143435.

42. Šircelj H, Tausz M, Grill D, Batič F. Detecting different levels of drought stress in apple trees (Malus *domestica* Borkh.) with selected biochemical and physiological parameters. Sci Hortic. 2007; 113(4):362-9.

DOI: 10.1016/j.scienta.2007.04.012

43. Pinheiro C, Chaves MM. Photosynthesis and drought: Can we make metabolic connections from available data? J Exp Bot. 2011;62(3):869-82.

DOI: 10.1093/jxb/erq340, PMID 21172816.

44. Jacinto Júnior SG, Moraes JGL, Silva FDBd, Silva BdN, Sousa GGd, Oliveira LLBd, et al. Physiological responses of fava bean (*Phaseolus lunatus* L.) genotypes subjected to water stress cultivated in the State of Ceará. Rev Bras Meteorol. 2019;34(3):413-22.

DOI: 10.1590/0102-7786343047.

45. Parry MA, Andralojc PJ, Khan S, Lea PJ, Keys AJ. RuBisCO activity: Effects of drought stress. Ann Bot. 2002;89;Spec No:833-9.

> DOI: 10.1093/aob/mcf103, PMID 12102509.

46. Dias MC, Brüggemann W. Limitations of photosynthesis in Phaseolus vulgaris under drought stress: gas exchange, chlorophyll fluorescence and Calvin cycle enzyms. Photosynthetica. 2010;48(1):96- 102.

DOI: 10.1007/s11099-010-0013-8

47. Flexas J, MEDRANO H. Drought-inhibition of photosynthesis in C3 plants: Stomatal and non-stomatal limitations revisited. Ann Bot. 2002;89(2):183-189.

> DOI: 10.1093/aob/mcf027, PMID 12099349.

48. Boussadia O, Mariem FB, Mechri B, Boussetta W, Braham M, Hadj SBE. Response to drought of two olive tree cultivars (cv Koroneki and Meski). Sci Hortic. 2008;116(4):388-93.

DOI: 10.1016/j.scienta.2008.02.016

49. Silveira NM, Ribeiro RV, de Morais SFN, de Souza SCR, da Silva SF, Seabra AB, et al. Leaf arginine spraying improves leaf

gas exchange under water deficit and root antioxidant responses during the recovery period. Plant Physiol Biochem. 2021; 162:315-26.

DOI: 10.1016/j.plaphy.2021.02.036, PMID 33714146.

50. Costa AS, Freire ALO, Bakke IA, Pereira FHF. Physiological and biochemical responses of mastic plants (*Myracrodruon urundeuva* Allemão) to water deficit and subsequent recovery. IRRIGATE. 2015;20(4):705-17

DOI: 10.15809/irriga.2015v20n4p705

51. Pessoa JL, Freire ALO, Costa AS. Gas exchange of Handroanthus impetiginosus (Mart. ex DC) Mattos plants submitted to water deficit and subsequent rehydration. Rev Cienc Agrovet. 2017; 16(3):269-76

DOI: 10.5965/223811711632017269

52. Ramos FR, Freire ALdO. Growth and gas exchange of Cnidoscolus quercifolius fertilized with potassium and under water deficit. J Agric Sci. 2019;11(11): 23-9.

DOI: 10.5539/jas.v11n11p23

53. Mendes HSJ, Paula NFd, Scarpinatti EA, Paula RCd. Physiological responses of Eucalyptus grandis x E. urophylla genotypes to water availability and potassium fertilization. Rev Cerne. 2013; 19(4):603-11

DOI:10.1590/S0104-77602013000400010.

54. Zahoor R, Dong H, Abid M, Zhao W, Wang Y, Zhou Z. Potassium fertilizer improves drought stress alleviation potential in cotton by enhancing photosynthesis and carbohydrate metabolism. Environ Exp Bot. 2017;137:73-83.

DOI: 10.1016/j.envexpbot.2017.02.002

55. Tsonev T, Velikova V, Yildiz-Aktas L, Gürel A, Edreva A. Effect of water deficit and potassium fertilization on photosynthetic activity in cotton plants. Plant Biosyst. 2011;145(4):841-7.

DOI: 10.1080/11263504.2011.560199

56. Hommel R, Siegwolf R, Saurer M, Farquhar GD, Kayler Z, Ferrio JP, et al. Drought response of mesophyll conductance in forest understory species – impacts on water-use efficiency and interactions with leaf water movement. Physiol Plant. 2014;152(1):98-114.

## DOI: 10.1111/ppl.12160, PMID 24483818.

- 57. Fernandes PD, et al. Growth of citrus hybrids and rootstock varieties under salinity. Acta Sci Agron. 2011;33: 259-267.
- 58. Pardo JM. Biotechnology of water and salinity stress tolerance. Curr Opin Biotechnol. 2010;21(2):185-96. DOI: 10.1016/j.copbio.2010.02.005, PMID 20189794.

\_ *© 2022 Sousa and Freire; This is an Open Access article distributed under the terms of the Creative Commons Attribution License [\(http://creativecommons.org/licenses/by/4.0\)](http://creativecommons.org/licenses/by/4.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.*

> *Peer-review history: The peer review history for this paper can be accessed here: https://www.sdiarticle5.com/review-history/89997*